Obstetric Evidence Based Guidelines
Third Edition
Edited by Vincenzo Berghella
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Edited by
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Philadelphia, Pennsylvania, USA
To Paola, Andrea, Pietro, mamma, and papá,
for giving me the serenity, love, and strength at home now, then, and in the future
to fulfill my dreams and spend my talents as best as possible.
To all those who loved the 1st edition
To the health of mothers and babies
And, as I often toast: To the next generation!
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Introduction

Welcome to the third edition of our evidence-based books on obstetrics and maternal-fetal medicine! I am indebted for your support! I can’t believe how much praise we have gotten for these companion volumes. Your words of encouragement have kept me and all the collaborators past and present going now for well over a decade (we are indebted to contributors to previous editions of this text for their work). It has been extremely worthwhile, and fulfilling. You are making me happy! In return, I hope we are helping you and your patients toward ever better evidence-based care of pregnant women and their babies, and therefore better outcomes. Indeed, maternal and perinatal morbidities and mortalities throughout the world are improving.

To me, pregnancy has always been the most fascinating and exciting area of interest, as care involves not one, but at least two persons—the mother and the fetus—and leads to the miracle of a new life. I was a third-year medical student when, during a lecture, a resident said: "I went into obstetrics because this is the easiest medical field. Pregnancy is a physiologic process, and there isn’t much to know. It is simple.” I knew from my “classical” background that “obstetrics” means to “stand by, stay near,” and that indeed pregnancy used to receive no medical support at all. After more than 25 years of practicing obstetrics, I now know that although physiologic and at times simple, obstetrics and maternal-fetal medicine can be the most complex of the medical fields: pregnancy is based on a different physiology than for nonpregnant women, can include any medical disease, require surgery, etc. It is not so simple. In fact, ignorance can kill, in this case with the health of the woman and her baby both at risk. Too often, I have gone to a lecture, journal club, rounds, or other didactic event to hear presented only one or a few articles regarding the subject, without the presenter reviewing the pertinent best review of the total literature and data. It is increasingly difficult to read and acquire knowledge of all that is published, even just in obstetrics, with about 3,000 scientific manuscripts published monthly on this subject. Some residents or even authorities would state at times that “there is no evidence” on a topic. We indeed used to be the field with the worst use of randomized trials [1]. As the best way to find something is to look for it, my coauthors and I searched for the best evidence. On careful investigation, indeed there are data on almost everything we do in obstetrics, especially on our interventions. Indeed, our field is now the pioneer for numbers of meta-analysis and extension of work for evidence-based reviews [2]. Obstetricians are now blessed with lots of data, and should make the best use of it.

The goals of this book are to summarize the best evidence available in the obstetrics and maternal-fetal medicine literature, and make the results of randomized controlled trials (RCTs) and meta-analyses of RCTs easily accessible to guide clinical care. The intent is to bridge the gap between knowledge (the evidence) and its easy application. To reach these goals, we reviewed all trials on effectiveness of interventions in obstetrics. Millions of pregnant women have participated in thousands of properly conducted RCTs. The efforts and sacrifice of mothers and their fetuses for science should be recognized at least by the physicians’ awareness and understanding of these studies. Some of the trials have been summarized in over 600 Cochrane reviews, with hundreds of other meta-analyses also published in obstetrical topics (Table 1). All of the Cochrane reviews, as well as other meta-analyses and trials in obstetrics and maternal-fetal medicine, were reviewed and referenced. The material presented in single trials or meta-analyses is too detailed to be readily translated to advice for the busy clinician who needs to make dozens of clinical decisions a day. Even the Cochrane Library, the undisputed leader for evidence-based medicine efforts, has been criticized for its lack of flexibility and relevance in failing to be more easily understandable and clinically readily usable [3]. It is the gap between research and clinicians that needed to be filled, making sure that proven interventions are clearly highlighted, and are included in today’s care. Just as all pilots fly planes under similar rules to maximize safety, all obstetricians should manage all aspects
of pregnancy with similar, evidenced-based rules. Indeed, only interventions that have been proven to provide benefit should be used routinely. On the other hand, *primum non nocere*: interventions that have clearly been shown to be not helpful or indeed harmful to mother and/or baby should be avoided. Another aim of the book is to make sure the pregnant woman and her unborn child are not marginalized by the medical community. In most circumstances, medical disorders of pregnant women can be treated as in nonpregnant adults. Moreover, there are several effective interventions for preventing or treating specific pregnancy disorders.

Evidence-based medicine is the concept of treating patients according to the best available evidence. While George Bernard Shaw said: “I have my own opinion, do not confuse me with the facts,” this can be a deadly approach, especially in medicine, and compromise two or more lives at the same time in obstetrics and maternal-fetal medicine. What should be the basis for our interventions in medicine? Meta-analyses provide a comprehensive summary of the best research data available. As such, they provide the best guidance for “effective’ clinical care [4]. It is unscientific and unethical to practice medicine, teach, or conduct research without first knowing all that has already been proven [4]. In the absence of trials or meta-analyses, lower level evidence is reviewed. This book aims at providing a current systematic review of all the best evidence, so that current practice and education, as well as future research can be based on the full story from the best-conducted research, not just the latest data or someone’s opinion (Table 2).

These evidence-based guidelines cannot be used as a “cookbook,” or a document dictating the best care. The knowledge from the best evidence presented in the guidelines needs to be integrated with other knowledge gained from clinical judgment, individual patient circumstances, and patient preferences, to lead to best medical practice. These are guidelines, not rules. Even the best scientific studies are not always perfectly related to any given individual, and clinical judgment must still be applied to allow the best “particularization” of the best knowledge for the individual, unique patient. Evidence-based medicine informs clinical judgment, but does not substitute it. It is important to understand though that greater clinical experience by the physician actually correlates with inferior quality of care if not integrated with knowledge of the best evidence [5]. The appropriate treatment is given in only 50% of visits to general physicians [5]. At times, limitations in resources may also limit the applicability of the guidelines, but should not limit the physician’s knowledge. Guidelines and clinical pathways based on evidence not only point to the right management, but also can decrease medicolegal risk [6]. We aimed for brevity and clarity. Suggested management of the healthy or sick mother and child is stated as straightforwardly as possible, for everyone to easily understand and implement (Table 3). If you find the Cochrane reviews, scientific manuscripts, and other publications difficult to “translate” into care of your patients, this book is for you. We wanted to prevent information overload.

Table 1 Obstetrical Evidence

<table>
<thead>
<tr>
<th>Evidence Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Over 600 current Cochrane reviews</td>
</tr>
<tr>
<td>Hundreds of other current meta-analyses</td>
</tr>
<tr>
<td>More than 1000 RCTs</td>
</tr>
<tr>
<td>Millions of pregnant women randomized</td>
</tr>
</tbody>
</table>

Table 2 Goals of This Book

<table>
<thead>
<tr>
<th>Goal</th>
</tr>
</thead>
<tbody>
<tr>
<td>Improve the health of women and their children</td>
</tr>
<tr>
<td>“Make it easy to do it right”</td>
</tr>
<tr>
<td>Implement the best clinical care based on science (evidence), not opinion</td>
</tr>
<tr>
<td>Education</td>
</tr>
<tr>
<td>Develop lectures</td>
</tr>
<tr>
<td>Decrease disease, use of detrimental interventions, and therefore costs</td>
</tr>
<tr>
<td>Reduce medicolegal risks</td>
</tr>
</tbody>
</table>
Table 3  This Book Is For

- Obstetricians
- Midwives
- Family medicine and others (practicing obstetrics)
- Residents
- Nurses
- Medical students
- Maternal-fetal medicine attendings
- Maternal-fetal medicine fellows
- Other consultants on pregnancy
- Lay persons who want to know “the evidence”
- Politicians responsible for health care

On the other hand, “everything should be made as simple as possible, but not simpler” (A. Einstein). Key management points are highlighted at the beginning of each guideline, and in bold in the text. The chapters are divided in two volumes, one on obstetrics and one on maternal-fetal medicine; cross-references to chapters in Maternal-Fetal Evidence Based Guidelines have been noted in the text where applicable. Please contact us (vincenzo.berghella@jefferson.edu) for any comments, criticisms, corrections, missing evidence, etc.

I have the most fun discovering the best ways to alleviate discomfort and disease. The search for the best evidence for these guidelines has been a wonderful, stimulating journey. Keeping up with evidence-based medicine is exciting. The most rewarding part, as a teacher, is the dissemination of knowledge. I hope, truly, that this effort will be helpful to you, too.

REFERENCES

5. Arky RA. The family business—To educate. NEJM. 2006;354:1922–1926. [Review]
How to “Read” This Book

The knowledge from randomized controlled trials (RCTs) and meta-analyses of RCTs is summarized and easily available for clinical implementation. Key management points are highlighted at the beginning of each guideline, and in bold in the text. Relative risks and 95% confidence intervals from studies are quoted sparingly. Instead, the straight recommendation for care is made if one intervention is superior to the other, with the percent improvement often quoted to assess degree of benefit. If there is insufficient evidence to compare to interventions or management, this is clearly stated.

References: Cochrane reviews with 0 RCT are not referenced, and, instead of referencing a meta-analysis with only one RCT, the actual RCT is usually referenced. RCTs that are already included in meta-analyses are not referenced, for brevity and because they can be easily accessed by reviewing the meta-analysis. If new RCTs are not included in meta-analysis, they are obviously referenced. Each reference was reviewed and evaluated for quality according to a modified method as outlined by the U.S. Preventive Services Task Force (http://www.ahrq.gov):

I  Evidence obtained from at least one properly designed randomized controlled trial.
II-1 Evidence obtained from well-designed controlled trials without randomization.
II-2 Evidence obtained from well-designed cohort or case-control analytic studies, preferably from more than one center or research group.
II-3 Evidence obtained from multiple time series with or without the intervention. Dramatic results in uncontrolled experiments could also be regarded as this type of evidence.
III (Review) Opinions of respected authorities, based on clinical experience, descriptive studies, or reports of expert committees.

These levels are quoted after each reference. For RCTs and meta-analyses, the number of subjects studied is stated and, sometimes, more details are provided to aid the reader to understand the study better.
<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ab</td>
<td>antibody</td>
</tr>
<tr>
<td>AC</td>
<td>abdominal circumference</td>
</tr>
<tr>
<td>ACA</td>
<td>anticardiolipin antibody</td>
</tr>
<tr>
<td>ACOG</td>
<td>American College of Obstetricians and Gynecologists</td>
</tr>
<tr>
<td>ACS</td>
<td>acute chest syndrome</td>
</tr>
<tr>
<td>ADR</td>
<td>autosomic dysreflexia</td>
</tr>
<tr>
<td>AF</td>
<td>amniotic fluid</td>
</tr>
<tr>
<td>AFI</td>
<td>amniotic fluid index</td>
</tr>
<tr>
<td>AFP</td>
<td>alpha-fetoprotein</td>
</tr>
<tr>
<td>AFV</td>
<td>amniotic fluid volume</td>
</tr>
<tr>
<td>Ag</td>
<td>antigen</td>
</tr>
<tr>
<td>AIDS</td>
<td>acquired immune deficiency syndrome</td>
</tr>
<tr>
<td>ALT</td>
<td>alanine aminotransferase</td>
</tr>
<tr>
<td>ANA</td>
<td>antinuclear antibodies</td>
</tr>
<tr>
<td>aPT</td>
<td>activated prothrombin time</td>
</tr>
<tr>
<td>APS</td>
<td>antiphospholipid syndrome</td>
</tr>
<tr>
<td>aPTT</td>
<td>activated partial thromboplastin time</td>
</tr>
<tr>
<td>AROM</td>
<td>artificial rupture of membranes</td>
</tr>
<tr>
<td>ART</td>
<td>assisted reproductive technologies</td>
</tr>
<tr>
<td>ARV</td>
<td>antiretroviral therapy</td>
</tr>
<tr>
<td>ASA</td>
<td>aspirin</td>
</tr>
<tr>
<td>ASD</td>
<td>atrial septal defect</td>
</tr>
<tr>
<td>AST</td>
<td>aspartate aminotransferase</td>
</tr>
<tr>
<td>AT III</td>
<td>antithrombin III</td>
</tr>
<tr>
<td>AZT</td>
<td>ziduvudine</td>
</tr>
<tr>
<td>bid</td>
<td>“bis in die,” i.e., twice per day</td>
</tr>
<tr>
<td>BPD</td>
<td>biparietal diameter</td>
</tr>
<tr>
<td>BPD</td>
<td>bronchopulmonary dysplasia</td>
</tr>
<tr>
<td>BPP</td>
<td>biophysical profile</td>
</tr>
<tr>
<td>BMI</td>
<td>body mass index</td>
</tr>
<tr>
<td>BP</td>
<td>blood pressure</td>
</tr>
<tr>
<td>CAP</td>
<td>community-acquired pneumonia</td>
</tr>
<tr>
<td>CBC</td>
<td>complete blood count</td>
</tr>
<tr>
<td>CCB</td>
<td>calcium channel blocker</td>
</tr>
<tr>
<td>CDC</td>
<td>Centers for Disease Control</td>
</tr>
<tr>
<td>CF</td>
<td>cystic fibrosis</td>
</tr>
<tr>
<td>CHD</td>
<td>congenital heart defect</td>
</tr>
<tr>
<td>CL</td>
<td>cervical length</td>
</tr>
<tr>
<td>CMV</td>
<td>cytomegalovirus</td>
</tr>
<tr>
<td>CNS</td>
<td>central nervous system</td>
</tr>
<tr>
<td>COX</td>
<td>cyclooxygenase</td>
</tr>
<tr>
<td>CRL</td>
<td>crown-rump length</td>
</tr>
<tr>
<td>CSE</td>
<td>combined spinal epidural</td>
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<tr>
<td>CSF</td>
<td>cerebrospinal fluid</td>
</tr>
<tr>
<td>CT</td>
<td>computerized tomography</td>
</tr>
<tr>
<td>CVS</td>
<td>chorionic villus sampling</td>
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<tr>
<td>DES</td>
<td>diethylstilbestrol</td>
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<tr>
<td>DIC</td>
<td>disseminated intravascular coagulation</td>
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<td>dilute Russell’s viper venom time</td>
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<td>deepest vertical pocket</td>
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<td>deep vein thrombosis</td>
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<tr>
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<td>external cephalic version</td>
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<tr>
<td>EDC</td>
<td>estimated date of confinement</td>
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<tr>
<td>EDD</td>
<td>estimated date of delivery</td>
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<td>electrocardiogram</td>
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<td>FOB</td>
<td>father of baby</td>
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<td>grams</td>
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<td>hemoglobin</td>
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<td>LR</td>
<td>likelihood ratio</td>
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<td>meconium aspiration syndrome</td>
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<td>middle cerebral artery</td>
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<td>MCH</td>
<td>methemoglobin</td>
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<tr>
<td>Abbreviation</td>
<td>Description</td>
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</tr>
<tr>
<td>MCV</td>
<td>mean corpuscular volume</td>
</tr>
<tr>
<td>MOM</td>
<td>multiple of the median</td>
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<tr>
<td>MRI</td>
<td>magnetic resonance imaging</td>
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<td>methylenetetrahydrofolate reductase</td>
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<td>MVP</td>
<td>maximum vertical pocket</td>
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<tr>
<td>NA</td>
<td>not available</td>
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<td>nonstress test</td>
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<td>n/v</td>
<td>nausea and/or vomiting</td>
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<td>OR</td>
<td>operating room</td>
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<td>ORA</td>
<td>oxytocin receptor agonist</td>
</tr>
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<td>PC</td>
<td>protein C</td>
</tr>
<tr>
<td>PCR</td>
<td>polymerase chain reaction</td>
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<td>pulmonary embolus</td>
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<td>pulmonary function tests</td>
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<td>prothrombin gene mutation</td>
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<td>pelvic inflammatory disease</td>
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<td>PL</td>
<td>pregnancy loss</td>
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<td>PNC</td>
<td>prenatal care</td>
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<td>po</td>
<td>“per os,” i.e., by mouth</td>
</tr>
<tr>
<td>PPH</td>
<td>postpartum hemorrhage</td>
</tr>
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<td>PRCD</td>
<td>planned repeat cesarean delivery</td>
</tr>
<tr>
<td>PS</td>
<td>protein S</td>
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<td>prothrombin time</td>
</tr>
<tr>
<td>PTB</td>
<td>preterm birth</td>
</tr>
<tr>
<td>PTT</td>
<td>partial thromboplastin time</td>
</tr>
<tr>
<td>PPROM</td>
<td>preterm premature rupture of membranes</td>
</tr>
<tr>
<td>pRBC</td>
<td>packed red blood cells</td>
</tr>
<tr>
<td>PROM</td>
<td>preterm rupture of membranes</td>
</tr>
<tr>
<td>PSV</td>
<td>peak systolic velocity</td>
</tr>
<tr>
<td>PTL</td>
<td>preterm labor</td>
</tr>
<tr>
<td>PTU</td>
<td>propylthiouracil</td>
</tr>
<tr>
<td>PUBS</td>
<td>percutaneous umbilical blood sampling</td>
</tr>
<tr>
<td>PWH</td>
<td>periventricular hemorrhage</td>
</tr>
<tr>
<td>qd</td>
<td>once a day</td>
</tr>
<tr>
<td>qid</td>
<td>four times per day</td>
</tr>
<tr>
<td>qhs</td>
<td>before bedtime</td>
</tr>
<tr>
<td>QS</td>
<td>quadruple screen</td>
</tr>
<tr>
<td>RBC</td>
<td>red blood cell</td>
</tr>
<tr>
<td>RCT</td>
<td>randomized controlled study</td>
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<td>RDS</td>
<td>respiratory distress syndrome</td>
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<tr>
<td>RNA</td>
<td>ribonucleic acid</td>
</tr>
<tr>
<td>ROM</td>
<td>rupture of membranes</td>
</tr>
<tr>
<td>RPR</td>
<td>rapid plasma reagin</td>
</tr>
<tr>
<td>RR</td>
<td>respiratory rate</td>
</tr>
<tr>
<td>Rx</td>
<td>treatment</td>
</tr>
<tr>
<td>SAB</td>
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<tr>
<td>SC</td>
<td>subcutaneous</td>
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<td>SCI</td>
<td>spinal cord injury</td>
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<tr>
<td>SDP</td>
<td>single deepest pocket</td>
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<td>SIDS</td>
<td>sudden infant death syndrome</td>
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<tr>
<td>SLE</td>
<td>systemic lupus erythematosus</td>
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<tr>
<td>SPTB</td>
<td>spontaneous preterm birth</td>
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<tr>
<td>STD</td>
<td>sexually transmitted diseases (synonym of STI)</td>
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<tr>
<td>TB</td>
<td>tuberculosis</td>
</tr>
<tr>
<td>TG</td>
<td><em>Toxoplasma gondii</em></td>
</tr>
<tr>
<td>tid</td>
<td>three times per day</td>
</tr>
<tr>
<td>TOL</td>
<td>trial of labor</td>
</tr>
<tr>
<td>TRAP</td>
<td>twin reversal arterial perfusion</td>
</tr>
<tr>
<td>TRH</td>
<td>thyrotropin-releasing hormone</td>
</tr>
<tr>
<td>TSH</td>
<td>thyroid-stimulating hormone</td>
</tr>
<tr>
<td>TSI</td>
<td>thyroid-stimulating immune globulins</td>
</tr>
<tr>
<td>TTTS</td>
<td>twin-twin transfusion syndrome</td>
</tr>
<tr>
<td>TVU</td>
<td>transvaginal ultrasound</td>
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<tr>
<td>UA</td>
<td>umbilical artery</td>
</tr>
<tr>
<td>UFH</td>
<td>unfractionated heparin</td>
</tr>
<tr>
<td>U/S (or u/s)</td>
<td>ultrasound</td>
</tr>
<tr>
<td>VBAC</td>
<td>vaginal birth after cesarean</td>
</tr>
<tr>
<td>VDRL</td>
<td>venereal disease research laboratory</td>
</tr>
<tr>
<td>VSD</td>
<td>ventricular septal defect</td>
</tr>
<tr>
<td>VTE</td>
<td>venous thromboembolism</td>
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<tr>
<td>WHO</td>
<td>World Health Organization</td>
</tr>
</tbody>
</table>
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Preconception care

Johanna Quist-Nelson

KEY POINTS

- Preconception care is a set of interventions that aim to identify and modify biomedical, behavioral, and social risks to a woman's health or pregnancy outcome through prevention and management. The foundation of preconception care is prevention.
- Preconception care should occur any time if any health-care provider sees a reproductive-age woman (e.g., 15–44 years old).
- Personal and family history, physical exam, laboratory screening, reproductive plan, nutrition, supplements, weight, exercise, vaccinations, and injury prevention should be reviewed in all reproductive-age women.
- Folic acid 400 µg/day, as well as proper diet and exercise, should be encouraged.
- Regarding vaccinations, women should receive the influenza vaccine if planning pregnancy during flu season; the rubella and varicella vaccines if there is no evidence of immunity to these viruses; and tetanus/diphtheria/pertussis if lacking adult vaccination.
- Specific interventions to reduce morbidity and mortality for both the woman and her baby should be offered to those identified with chronic diseases or exposed to teratogens or illicit substances.

HISTORY

Preconception care has ancient origins. Plutarch (46–120 CE) wrote that the ancient Spartans “[...] ordered the maidens to exercise [...] to the end that the fruit they conceived might [...] take firmer root and find better growth” [1].

DEFINITION

Preconception care is a set of interventions that aim to identify and modify biomedical, behavioral, and social risks to a woman's health or pregnancy outcome through prevention and management [2,3]. This care has also been called prepregnancy, interpregnancy care, or periconceptional medicine [4].

AIM AND EFFECTIVENESS

The foundation of preconception care is prevention. Prevention of disease is the most effective form of medicine, and health care should shift from the delivery of procedure-based acute care to the provision of counseling-based preventative care [5,6]. For example, the two leading causes of death in the first year of life—birth defects and disorders caused by preterm birth (PTB)—can both be significantly reduced by preconception care. Randomized controlled trials have corroborated that women are likely to incorporate change in modifiable health behaviors in response to preconception counseling [7]. General practitioner-initiated preconception counseling not only decreases adverse pregnancy outcomes but also reduces anxiety in reproductive-age women [8].

TIMING AND TARGET POPULATION

The time that people should start caring for a pregnancy is not after, but before, conception. Preconception care should occur any time any health-care provider sees a reproductive-age woman. A reproductive-age woman is usually defined as between 15 and 44 years of age, but occasionally even younger or older women contemplate, or at least are at risk of, pregnancy. The first prenatal visit is “months too late!” [9]. It often happens after first-trimester exposure to a potential teratogen has already occurred. There are about 1 billion reproductive-age women worldwide. In the United States, as an example, only about half of pregnancies are planned. As women get pregnant later in life, disease prevalence and medication exposures increase. Approximately 80% of reproductive-age U.S. women have dental disease, 66% are obese or overweight, 55% drink alcohol, 11% continue to smoke during pregnancy, 9% have diabetes, 6% asthma, 3% hypertension, and 3% cardiac disease [2]. The incidences of many of these conditions, even among pregnant women, are on the rise.

While some beneficial interventions could be started as soon as a pregnancy is diagnosed, this is unrealistic. Many of the preventive measures take time, often months, such as quitting smoking, losing weight, folic acid supplementation, and stabilization of medical conditions with effective and safe medications.

OPPORTUNITIES FOR PRECONCEPTION CARE

By age 25, about 50% of U.S. women have had at least one birth. The highest fertility rate occurs in 25- to 30-year-old women. By age 44, >85% have given birth at least once. About 84% of reproductive-age women, when asked, answer that they had a health-care visit within the prior year [6]. Therefore, universal preconception care can be achieved if health-care providers make it a priority and plan for it at every opportunity (Table 1.1). The approach should be “every reproductive-age woman, every time” [6]. Every reproductive-age woman should be asked at every health-care encounter: “Are you considering pregnancy?” and “Could you possibly become pregnant?” Increased awareness of preconception care can be accomplished through improving health resources, public outreach, and advertising. Despite its great effectiveness, not all health-care plans cover preconception care. A preconception visit (or often more than one) should be standard primary care, as stated by the Center for Disease Control [2]. It should be as routine, if not more so, as prenatal care, as should the screening and interventions associated with it. A clear political will to drive the funding and insurance coverage for preconception care is required.

Therefore, providers of all specialties should be aware of the evidence-based recommendations (Tables 1.1–1.8).
Organizations representing family and internal medicine, obstetrics and gynecology, nurse midwifery, nursing, public health, diabetes, neurology, cardiology, and many other associations have supported recommendations for preconception care. Unfortunately, practitioners seldom implement them [10], even though it is an opportunity to optimize the health of the woman independent of whether she is planning pregnancy [6]. Only one out of six obstetrician-gynecologists (ob-gyns) or family physicians provides preconception care to the majority of women for whom they provide prenatal care [11].

Preconception care may often need to be multidisciplinary care. Prior to pregnancy, a woman can have numerous different medical problems affecting different specialties, and her care should occur in close collaboration among the different fields involved. Maternal physiology is different than nonpregnant adult physiology. An entire field, maternal-fetal medicine, is dedicated to the care of pregnancies with maternal or fetal problems, and these specialists are particularly adept at directing best practices for preconception counseling. Preconception care occurs best if all practitioners, including primary and specialty care, either directly implement or appropriately refer for implementation of effective preconception screening and intervention. The worse scenario is the belief that a positive pregnancy test is a good reason to “stop all medicines” thereby stopping disease treatment. Prevent panic; get women ready for a healthy pregnancy before contraception is stopped.

CONTENT OF PRECONCEPTION CARE
Topics pertinent to optimizing preconception health and therefore future maternal and perinatal outcome should be discussed. Topics to be discussed in preconception care are listed in Table 1.2 [2,12]. Further research is needed to determine the best content of preconception care and the most effective way to implement it [13,14].

UNIVERSAL SCREENING AND RELATED INTERVENTIONS
History, Exam, and Laboratory Screen
Suggested preconception screening assessment is shown in Table 1.3 [12,15]. A questionnaire should be completed ahead of time, either on paper or online, to review this extensive list. A standardized form improves the completeness of preconception screening, which necessitates time and commitment [16]. This standardized preconception form should be integrated into the permanent record of all reproductive-age woman. In a randomized trial, women assigned to be screened with a preconception risk survey were found to have an average of nine risk factors, supporting the facts that even low-risk women may benefit from preconception screening [13].

History should be detailed, especially when pertinent positives are detected. Prior inpatient and outpatient medical records should be reviewed. Women should be empowered with easy access to their records (best if electronic), to facilitate multispecialty care coordination. Personal prenatal medical record access has been associated with increased maternal control, satisfaction during pregnancy, and increased availability of antenatal records during hospital attendance [17].

Prior obstetrical and gynecological history, including prior pregnancy complications, should be reviewed. Other reproductive issues should also be assessed: fertility, including the possibility of assisted reproductive technology needs, sexuality (in particular high-risk behaviors), contraception, partner selection, and sexual function. Several social issues need to be reviewed as well (Table 1.3).

All couples should have a basic screen for family history of heritable genetic disorders, with a pedigree to at least the second prior generation. Women belonging to an ethnic group at increased risk for a recessive condition (Table 1.4) should be offered appropriate screening. All couples should be made aware of the option for cystic fibrosis (CF) screening, especially those who have a family history of CF, are in a high-risk group, or are reproductive partners of individuals with CF [18]. Women with a specific indication for genetic testing should be referred for formal genetic counseling (see Chapters 5 and 6).

Physical exam details are shown in Table 1.3. Pelvic exam may include cytologic and sexually transmitted infection screening for women with certain risk factors. Laboratory tests are done routinely (Table 1.3) and depend on risk factors (Table 1.4) [12].

Reproductive Health Plan
Asking a reproductive-age woman, and therefore inducing her to think about her reproductive health plan should be a priority of any medical visit [19]. Such a plan should address the desire (or not) for children; the optimal number, spacing, and timing of pregnancies; contraception to achieve this plan; opportunities to improve her health and therefore a successful reproductive life; and age-related changes in fertility [19]. Having a reproductive health plan reduces unintended pregnancies, age-related infertility, and fetal exposure to teratogens [2]. Very few women know that a short interpregnancy interval (i.e., <6 months from the end of last pregnancy to
the next conception) is associated with increased incidence of both small-for-gestational-age and low-birth-weight neonates [20]. Folic acid depletion may be the etiology for these increased risks [21]. Education and contraception advice are necessary to aim for the wished reproductive plan, avoiding unplanned pregnancies, and optimizing the 18- to 24-month interpregnancy interval goal. In a nonrandomized study, preconception care decreased the number of unintended pregnancies [22].

Table 1.3 Preconception Screening Assessment for all Reproductive-Age Women (15–44 Years Old)

<table>
<thead>
<tr>
<th>History</th>
<th>Reason for visit</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>Health status: obstetrical, gynecological, medical, surgical, and family history</td>
</tr>
<tr>
<td></td>
<td>Use of prescription, over-the-counter, complementary, and alternative medicines</td>
</tr>
<tr>
<td></td>
<td>Allergies (to medications or other)</td>
</tr>
<tr>
<td></td>
<td>Tobacco, alcohol, other drug use</td>
</tr>
<tr>
<td></td>
<td>Work-related exposures</td>
</tr>
<tr>
<td></td>
<td>Dietary/nutrition assessment</td>
</tr>
<tr>
<td></td>
<td>Physical activity</td>
</tr>
<tr>
<td></td>
<td>Urinary and fecal incontinence</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Physical examination</th>
<th>Height, weight, body mass index (BMI)</th>
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<tbody>
<tr>
<td></td>
<td>Blood pressure</td>
</tr>
<tr>
<td></td>
<td>Head</td>
</tr>
<tr>
<td></td>
<td>Neck: adenopathy and thyroid</td>
</tr>
<tr>
<td></td>
<td>Breasts</td>
</tr>
<tr>
<td></td>
<td>Heart, lungs</td>
</tr>
<tr>
<td></td>
<td>Abdomen</td>
</tr>
<tr>
<td></td>
<td>Pelvic examination</td>
</tr>
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<td></td>
<td>Skin</td>
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<table>
<thead>
<tr>
<th>Laboratory testing</th>
<th>Rubella titer*</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Varicella titer*</td>
</tr>
<tr>
<td></td>
<td>Human immunodeficiency virus (HIV) testing*</td>
</tr>
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<td></td>
<td>Cervical cytology*</td>
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<tr>
<td></td>
<td>Chlamydia testing (if aged 25 years or younger and sexually active)</td>
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</table>

<table>
<thead>
<tr>
<th>Evaluation and counseling</th>
<th>Sexuality and reproductive planning</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>High-risk behaviors</td>
</tr>
<tr>
<td></td>
<td>Discussion of a reproductive health plan</td>
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<tr>
<td></td>
<td>Contraceptive options for prevention of unwanted pregnancy, including emergency contraception</td>
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<tr>
<td></td>
<td>Genetic counseling</td>
</tr>
<tr>
<td></td>
<td>Sexually transmitted diseases</td>
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<tr>
<td></td>
<td>Partner selection</td>
</tr>
<tr>
<td></td>
<td>Barrier protection</td>
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</table>

<table>
<thead>
<tr>
<th>Sexual function</th>
<th>Partner selection</th>
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</thead>
<tbody>
<tr>
<td>Fitness and nutrition</td>
<td>Barrier protection</td>
</tr>
<tr>
<td>Dietary/Nutrition assessment</td>
<td></td>
</tr>
<tr>
<td>Exercise program</td>
<td></td>
</tr>
<tr>
<td>Folic acid supplementation (0.4 mg/day)</td>
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<tr>
<td>Calcium intake</td>
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<table>
<thead>
<tr>
<th>Psychosocial evaluation</th>
<th>Abuse/neglect/violence (physical, sexual, and emotional)</th>
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<tbody>
<tr>
<td></td>
<td>Sexual practices</td>
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<tr>
<td></td>
<td>Lifestyle/stress</td>
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<td>Sleep disorders</td>
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<td>Home and work (including satisfaction, and environmental hazards)</td>
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<td></td>
<td>Interpersonal/family relationships; social support</td>
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<tr>
<td></td>
<td>Depression (suicide)</td>
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<td>Criminality</td>
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<td>Education</td>
</tr>
<tr>
<td></td>
<td>Language and culture</td>
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<td>Health insurance status; coverage; access; public programs</td>
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<tr>
<th>Cardiovascular risk factors</th>
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<tr>
<td></td>
<td>Hypertension</td>
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<tr>
<td></td>
<td>Dyslipidemia</td>
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<tr>
<td></td>
<td>Obesity</td>
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<td></td>
<td>Diabetes mellitus</td>
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<thead>
<tr>
<th>Health/risk behaviors</th>
<th>Hygiene (including dental)</th>
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<tr>
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<td>Injury prevention</td>
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<td>Safety belts and helmets</td>
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<td>Recreational hazards</td>
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<td>Hearing</td>
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<td>Exercise and sports involvement</td>
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<tr>
<th>Breast self-examination</th>
<th>Vaccinations</th>
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<tr>
<td></td>
<td>See Table 1.5</td>
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</table>


All women should be counseled that 2%-3% of babies are born with minor (usually) or major anomalies. Screening and diagnostic options to detect aneuploidy and birth defects should be reviewed so that women may consider their options in relation to their personal values.

**Nutrition, Weight, and Exercise**

Lifelong habits of healthy diet and regular exercise should be established preconceptionally [23]. Proper diet and exercise can prevent several complications of pregnancy, including gestational diabetes and hypertensive complications [24]. Some studies suggest a correlation between a diet high in fruits, vegetables, nuts, and legumes, less than two servings of meat weekly and at least two servings of fish weekly (the “Mediterranean Diet”) with decreased rates of infertility and PTB [25–27].

In addition to following a healthy diet, issues of food safety are important to review. All meat, seafood, and shellfish should be thoroughly cooked. Eating at least 12 oz. of fish weekly is associated with several benefits, including a lower rate of PTB (see Chapter 17), but women must avoid >2 serving/week of shark, swordfish, king mackerel, some tuna, or tilefish, all of which may contain high concentrations of methyl mercury. Albacore (white) tuna has more mercury than canned, light tuna [28]. Other recommendations include eating only pasteurized eggs and dairy products and washing raw fruits and vegetables before eating. Women should try to obtain a minimum daily iodine intake of 150 mg/day. Education about proper hand, food, and cooking utensil hygiene is important, especially in developing countries.

Body mass index (BMI) should be calculated at least annually for reproductive-age women [29]. For women with a BMI that falls outside the normal range [28–34], preconception...
Table 1.4  Preconception Laboratory Screening Depending on Risk Factors

**Personal history:**

**Age:**
- >35: fasting glucose

**Race:**
- African-American: fasting glucose; hemoglobin electrophoresis (for sickle cell disease)
- Hispanic, Native American, Pacific Islander: fasting glucose
- Mediterranean: mean corpuscular volume (MCV) screening (for thalassemia)
- Ethnic testing: Ashkenazi—familial dysautonomia; Tay-Sachs; Canavan; Fanconi anemia type C; Niemann-Pick disease type A; Bloom’s syndrome; Gaucher disease; glycogen storage 1a; maple syrup urine disease; mucolipidosis type IV

**Prior obstetrical history:**
- Prior birth of a newborn weighting more than 9 lb or >4500 g (macrosomia): fasting glucose
- History of gestational diabetes mellitus: fasting glucose
- Prior unexplained fetal death: check autopsy and karyotype of fetal death; antiphospholipid antibody testing; fasting glucose
- Prior infant with congenital anomaly (if not screened in that pregnancy): fasting glucose
- Prior recurrent unexplained early pregnancy loss: antiphospholipid antibody testing; study of uterine anatomy; parental karyotype

**Prior medical history:**
- Metabolic syndrome/obesity; family history of lipid or coronary disorders: cholesterol/lipid profile
- Diabetes: lipid profile; hemoglobin A1c; cardiac and renal baseline function assessment; ophthalmologic exam
- Hypertension: fasting glucose; baseline cardiac, renal, and liver functions
- Multiple coronary heart disease risk factors (e.g., tobacco use, hypertension): lipid profile
- High-density lipoprotein cholesterol level ≤35 mL/dL: fasting glucose
- Triglyceride level ≥250 mg/dL: fasting glucose
- History of impaired glucose tolerance or impaired fasting glucose: fasting glucose
- Chronic use of steroids: fasting glucose
- Polycystic ovary syndrome: fasting glucose
- History of vascular disease: fasting glucose
- Marfan syndrome: echocardiogram for assessment of aortic root; eye exam for lens
- History of STD, drug abuse, etc.: HIV, Hep C
- Recipients of blood from donors who later tested positive for HCV infection: Hep C
- Recipients of blood or blood-component transfusion or organ transplant before July 1992: Hep C
- Recipients of clotting factor concentrates before 1987: Hep C
- Chronic (long-term) hemodialysis: Hep C
- History of transfusion from 1978 to 1985: HIV
- Invasive cervical cancer: HIV
- HIV infection: STD screening; PPD
- Medical risk factors known to increase risk of TB if infected: PPD
- Not sure whether patient had varicella infection or vaccination in past: varicella titer

**Social history:**
- HIV or TB contact, IV drug use, etc.: TB testing
- History of injecting illegal drugs: Hep C; HIV; STD screening (chlamydia, gonorrhea, syphilis, etc.); PPD
- Occupational percutaneous or mucosal exposure to HCV-positive blood: Hep C
- More than one sexual partner since most recent HIV test or a sex partner with more than one sexual partner since most recent HIV test: HIV
- Seeking treatment for STDs: HIV
- History of prostitution: STD screening; HIV
- Past or present sexual partner who is HIV positive or bisexual or injects drugs: HIV
- Long-term residence or birth in an area with high prevalence of HIV infection: HIV
- Adolescents who are or ever have been sexually active: HIV
- Adolescents entering detention facilities: HIV; STD screening
- Offer to women seeking preconception evaluation: HIV (all women should be screened)
- History of multiple sexual partners or a sexual partner with multiple contacts: STD screening
- Sexual contact with individuals with culture-proven STD: STD screening
- History of repeated episodes of STDs: STD screening
- Attendance at clinics for STDs: STD screening
- All sexually active women aged 25 years or younger: chlamydia
- All sexually active adolescents: gonorrhea
- Close contact with individuals known or suspected to have TB: PPD
- Born in country with high TB prevalence: PPD
- Medically underserved: PPD
- Low income: PPD
- Alcoholism: PPD
- Resident of long-term care facility (e.g., correctional institutions, mental institutions, and nursing homes and facilities): PPD
- Health professional working in high-risk health-care facilities: PPD

**Family history:**
- Family history of diabetes mellitus: fasting glucose
- Family history of diabetes; history of gestational diabetes, overweight/obese, hypertension, high-risk ethnic group (African-American, Hispanic, Native American): fasting glucose every 3 years
counseling regarding the woman’s increased risk of complications in pregnancy is extremely important. **Formal nutritional counseling should be offered and goals set to avoid pregnancy until optimal weight is achieved.** Women with low BMI should be screened for eating disorders. In overweight and obese women, **calorie and portion-size control** may be the most effective methods of sustained preconception weight loss. Unfortunately, there are no current evidence-based guidelines as to the most effective method of weight loss in the preconception period for obese and overweight patients [34]. Postpartum individual counseling on diet and physical activity increased the proportion of women returning to prepregnancy weight from 30% to 50% in one randomized trial [30].

An exercise routine that can be started preconceptionally and safely continued in pregnancy may include yoga; brisk walking (including hiking and backpacking); jogging; swimming; biking; cross-country skiing; and using fitness equipment such as an elliptical trainer, treadmill, or stationary bike. Women should be given standard advice for engaging in regular physical activity for 30–60 min/day for 5 or more days per week.

**Supplements**

The preconception intervention with the most evidence-based data to support its efficacy is **folic acid supplementation**. Folic acid supplementation is recommended, with a **minimum of 400 µg/day** for all women (93% decrease in neural tube defects [NTDs]), and **4 mg/day for women with prior children with NTDs** (69% decrease in recurrent NTDs) [32].

Supplementation should start at least 1 month before conception and continue until at least 28 days after conception (time of neural tube closure). Given the unpredictability of planned conception, all reproductive-age women should be on folic acid supplementation from menarche to menopause. Women taking oral contraceptives, all reproductive-age women should be on folic acid of neural tube closure). Given the unpredictability of planned conception and continue until at least 28 days after conception (time of neural tube closure). Folic acid supplementation has also been associated with a decrease in the risk of congenital anomalies other than NTDs (e.g., cardiac, facial clefts) [34,35].

The overall benefits or risks of fortifying basic foods such as grains with added folate have been associated with a 140–200 µg/day increase in supplementation and a 20%–50% decrease in incidence of NTD [33,36]. Education with provision of printed material [32,37], computerized counseling [38], and learner-centered nutrition education [39] all increase the awareness of the folate/NTDs association and the use of the folate supplements. These interventions may be effective in increasing the prophylactic use of additional preconception care activities.

There is insufficient evidence to justify the routine use of other supplements in reproductive-age women, especially in the developed world, unless a nutritional deficiency has been identified. It is important to obtain a **minimum daily iodine intake of 150 mg/day** and 10,000 IU daily of vitamin A (as beta-carotene) if deficiencies in these nutrients are identified. The use of certain supplements may be detrimental, especially if excessive amounts of lipid-soluble vitamins such as vitamin A (>10,000 IU/day) are taken, since they can be teratogenic. All supplements, including alternative and complementary medicines, should be reviewed (see also Chapter 2) [40,41].

**Vaccines**

Preconception vaccination for the prevention of fetal and maternal disease is an important preconception intervention (Table 1.5) (see also Chapter 38 in Maternal-Fetal Evidence Based Guidelines). Maternal immunity to infections such as rubella and varicella should be assessed for potential vaccination of nonimmune women, thus eliminating their risk for congenital syndromes associated with these viruses. Vaccination with live attenuated viruses should occur at least 4 weeks prior to conception due to theoretical risk of live virus affecting the fetus.

**Annual influenza vaccination** for women and their partners contemplating pregnancy will reduce the chance of maternal prenatal infection, a time during which higher morbidity has been documented. Influenza vaccination for new mothers and other close contacts of the newborn will reduce risk of infection for the child who is unable to receive vaccination until 6 months of age. Through this process of “cocooning,” the newborn is protected from the high morbidity and mortality rates associated with influenza in the first year of life [42].

### Table 1.4  Preconception Laboratory Screening Depending on Risk Factors (Continued)

<table>
<thead>
<tr>
<th>Risk Factor</th>
<th>Laboratory Screening</th>
</tr>
</thead>
<tbody>
<tr>
<td>Family history suggestive of familial hyperlipidemia: lipid profile</td>
<td>Family history of premature (age &lt; 50 years for men, age &lt; 60 years for women) cardiovascular disease: lipid profile</td>
</tr>
<tr>
<td>Family history of premature (age &lt; 50 years for men, age &lt; 60 years for women) cardiovascular disease: lipid profile</td>
<td>Family history of premature (age &lt; 50 years for men, age &lt; 60 years for women) cardiovascular disease: lipid profile</td>
</tr>
<tr>
<td>Colorectal cancer or adenomatous polyps in first-degree relative younger than 60 years or in two or more first-degree relatives of any ages; family history of familial adenomatous polyposis or hereditary nonpolyposis colon cancer: colonoscopy</td>
<td>First-degree relative (i.e., mother, sister, or daughter) or multiple other relatives who have a history of premenopausal breast or breast or ovarian cancer: mammography</td>
</tr>
<tr>
<td>Family history of Marfan syndrome: echocardiogram for assessment of aortic root; eye exam for lens</td>
<td>Family history of breast cancer: mammography</td>
</tr>
<tr>
<td>Family history of thyroid disease: TSH</td>
<td>Family history of Marfan syndrome: echocardiogram for assessment of aortic root; eye exam for lens</td>
</tr>
<tr>
<td>Physical examination:</td>
<td>Physical examination:</td>
</tr>
<tr>
<td>• Overweight (BMI ≥ 25): fasting glucose</td>
<td>• Overweight (BMI ≥ 25): fasting glucose</td>
</tr>
<tr>
<td>• Hypertension: fasting glucose</td>
<td>• Hypertension: fasting glucose</td>
</tr>
<tr>
<td>Laboratory screening:</td>
<td>Laboratory screening:</td>
</tr>
<tr>
<td>• Persistently abnormal alanine aminotransferase levels: Hep C</td>
<td>• Persistently abnormal alanine aminotransferase levels: Hep C</td>
</tr>
<tr>
<td>• Glycosuria: fasting glucose</td>
<td>• Glycosuria: fasting glucose</td>
</tr>
</tbody>
</table>

Source: Modified from Henderson JT et al., Women Health Issue, 12, 138–149, 2002. 
Abbreviations: STDs, sexually transmitted diseases; HIV, human immunodeficiency virus; Hep C, hepatitis C; HCV, hepatitis C virus; PPD, purified protein derivative; TB, tuberculosis; IV, intravenous; TSH, thyroid-stimulating hormone; BMI, body mass index.
### Table 1.5  Recommended Preconception Vaccinations

**All reproductive-age women**

- During flu season: Influenza
- No evidence of immunity to rubella: MMR
- No evidence of immunity to varicella: Varicella
- No adult Td vaccination in last 2 years: Tetanus/Diphtheria/Pertussis (Tdap)
- Hepatitis B nonimmune: Hepatitis B vaccine

**Age**

- All girls and women 9–26 years old: HPV
- All persons 18 years old and younger without immunity to hepatitis B infection: Hepatitis B

**Occupational**

- Health-care workers: Hepatitis B, Influenza, MMR, Varicella
- Public safety workers who have exposure to blood in the workplace: Hepatitis B
- Students in schools of medicine, dentistry, nursing, laboratory technology, and other allied health professions: Hepatitis B
- Staff of institutions for the developmentally disabled: Hepatitis B
- Individuals who work with HAV-infected nonhuman primates or with HAV in a research laboratory setting: Hepatitis A
- Military recruits: Meningococcus
- Microbiologists routinely exposed to *Neisseria meningitidis* isolates: Meningococcus

**Social history/living situation**

- Individuals with more than one sexual partner in the previous 6 months: Hepatitis B
- Households with child who is allergic to eggs and children who are on the autism spectrum: Hepatitis B
- Clients of institutions for the developmentally disabled: Hepatitis B
- Illegal injected drug users: Hepatitis B
- Illegal drug users (injected and noninjected): Hepatitis A
- Exposure to environment where pneumococcal outbreaks have occurred: Pneumococcus
- Native Alaskan/Native American: Pneumococcus
- Alcohol abuse: Pneumococcus
- Tobacco smoking: Pneumococcus
- Residents of long-term care facilities: Influenza, Pneumococcus
- First-year college students living in dormitories: Meningococcus

**Travel/immigration**

- Individuals traveling to or working in countries that have high or intermediate endemicity of hepatitis A: Hepatitis A
- International travelers who will be in countries with high or intermediate prevalence of chronic hepatitis B infection for more than 6 months: Hepatitis B
- Travel to areas hyperendemic or epidemic for *Neisseria meningitides*: Meningococcus

**Pulmonary conditions**

- Chronic pulmonary disorders, including asthma: Pneumococcus

**Cardiac conditions**

- Chronic cardiovascular disorders (e.g., CHF, cardiomyopathies): Influenza, Pneumococcus

**Renal conditions**

- Chronic metabolic diseases, including renal dysfunction: Influenza, Pneumococcus
- Nephrotic syndrome: Pneumococcus
- End-stage renal disease including those on dialysis: Hepatitis B

**Endocrine conditions**

- Diabetes mellitus: Influenza, Pneumococcus

**Hematologic/Immunologic conditions**

- Prior transfusions: Hepatitis A, Hepatitis B
- Patients with clotting factor disorders (those who receive clotting factor concentrates): Hepatitis A
- Chronic illness, such as functional asplenia (e.g., sickle cell disease) or splenectomy: Pneumococcus
- Immunocompromised patients (e.g., HIV infection, hematologic or solid malignancies, chemotherapy, steroid therapy): Pneumococcus
- Adults with anatomic or functional asplenia: Pneumococcus, Meningococcus
- Terminal complement component deficiencies: Meningococcus

**Infectious conditions**

- Individuals with a recently acquired or recent evaluation for STI: Hepatitis B
- All clients in STD clinics: Hepatitis B
- HIV: Hepatitis B, Influenza, Pneumococcus, consider Meningococcus
- Individuals with Hepatitis C: Hepatitis A, Hepatitis B

**GI/Hepatic conditions**

- Chronic liver disease: Hepatitis A, Hepatitis B, Pneumococcus

**Neurologic conditions**

- Cerebrospinal fluid leaks: Pneumococcus

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*Source:* Modified from Henderson JT et al., *Women Health Issue*, 12, 138–149, 2002 (see also Chapter 38 in Maternal-Fetal Evidence Based Guidelines).

*Abbreviations:* MMR, measles, mumps, and rubella; HPV, human papillomavirus; HAV, hepatitis A virus; CHF, congestive heart failure; HIV, human immunodeficiency virus; STI, sexually transmitted infection; STD, sexually transmitted disease; Td, tetanus diphtheria.
**Hepatitis B vaccination** should be offered to all susceptible women of reproductive age in regions with intermediate and high rates of endemcity (where ≥2% of the population is hepatitis B surface antigen [HBsAg] positive). Perinatal transmission of hepatitis B results in 90% chance of chronic infection in the newborn, which places the child at risk for cirrhosis and hepatocellular carcinoma. In regions of low prevalence, vaccination should be targeted to high-risk groups (Table 1.5).

**Tetanus vaccination** should remain up-to-date in reproductive-age women, particularly in regions of the world where maternal and neonatal tetanus is prevalent [43]. This has been shown to markedly reduce the incidence of tetanus related to parturition. Due to increasing prevalence and the high morbidity and mortality rates of neonatal pertussis, vaccination (in combination with tetanus and diphtheria) is recommended for all women and their partners of reproductive age who have not been immunized in their adult lives (since age 11 years) [44]. It is well documented that 75% of cases of neonatal pertussis have a family member as the index case [45]. Again, through the concept of cocooning, the incidence of neonatal pertussis can be reduced.

Other vaccination recommendations based on medical, occupational, or social risks are described in Table 1.5.

**Injury Prevention**
The second leading cause of death in reproductive-age women is accidents. Use of **seat belts** and **helmets** should be reviewed and strongly encouraged where appropriate. Inquiry should be made regarding occupational and recreational hazards. Possession and use of firearms should be evaluated. Possession and use of firearms should be evaluated, especially in individuals with a history of significant mental health diagnoses.

**Universal Recommendations**
Preconception recommendations for all women are listed in Table 1.6. Reproductive-age women should be aware of these evidence-based recommendations, both through their doctors and through public awareness campaigns. Several online resources are available [46–49]. Women and their partners should take more responsibility for their care and the future health of their offspring, and implement the health and lifestyle changes recommended.

**SPECIFIC INDIVIDUAL ISSUES**

**History of PTB**
There are currently no preconception recommendations for a woman with a history of PTB outside of the general recommendations for women trying to conceive. Randomized controlled trials examining preconception initiation of low-dose aspirin did not demonstrate an increased live birth rate or decrease in PTB [50,51]. Interval antibiotic treatment with azithromycin and metronidazole between pregnancies in women with a prior spontaneous PTB <34 weeks has not been associated with decreased risk of preterm delivery [52,53].

**Advanced Maternal Age**
In recent years, there has been a trend to delay childbearing. This trend is especially prevalent in developed countries, for example, in the United States where the birth rate in women age 40–44 has increased from 5.2 births per 1000 in 1990 to 10.4 births per 1000 women in 2013 [54]. It is well established that women of advanced maternal age (AMA) are at increased risks of poor obstetric outcomes, stillbirth, and fetal death [55–57]. Women of extreme AMA (>45 years old) have been found to increase the prevalence of preexisting chronic disease [58]. Although no Level I evidence exists for preconception testing in this population, it is reasonable to screen patients of extreme AMA for chronic hypertension, diabetes, hyperlipidemia, or heart disease with a cardiac echocardiogram.

**Chronic Diseases**
The incidences of several medical disorders such as obesity, diabetes mellitus, and hypertension are high and on the rise in reproductive-age women. There is literature for evidence-based recommendations on each disease or condition that can involve the reproductive-age woman and affect her reproductive health [3,4,15]. Full review of each is behind the scope of this chapter (see individual chapters in Maternal-Fetal Evidence Based Guidelines). Some common conditions are discussed for brief preconception management review (Table 1.7).

**Diabetes**
Diabetes (see Chapters 4 and 5 in Maternal-Fetal Evidence Based Guidelines) is associated with an increased risk of congenital anomalies, in particular cardiac defects and NTDs, if poorly controlled in the first weeks of pregnancy. The risk of congenital anomalies is related to long-term diabetic control, reflected in the level of glycosylated hemoglobin (HgB A1c): <7% = no increased risk (2%–3% baseline); 7%–9% = 15%; 9%–11% = 23%; >11% = 25% [32]. It has been estimated that euglycemia (with normal HgB A1c) during the first trimester, which can only be achieved through attentive preconception counseling, could prevent >100,000 U.S. pregnancy losses or birth defects per year [2]. Another cost analysis reported that universal preconception care could lead to averted lifetime costs for the affected cohort of children as high as $4.3 billion [59,60]. The benefits of preconception diabetes care have been previously demonstrated [61,62], even in teenagers [63]. Preconception care is also essential for counseling of the woman with conditions severe enough to make a successful pregnancy extremely challenging.

**Table 1.6** Preconception Interventions for All Women

<table>
<thead>
<tr>
<th>Intervention</th>
<th>Prevention of</th>
</tr>
</thead>
<tbody>
<tr>
<td>Folic acid 400 μg/day</td>
<td>NTDs, and also probably cardiac defects, facial clefts</td>
</tr>
<tr>
<td>Vaccinations</td>
<td>Maternal/perinatal infection (Table 1.5)</td>
</tr>
<tr>
<td>Proper diet and exercise</td>
<td>Obesity, diabetes, hypertensive diseases, and their consequences</td>
</tr>
<tr>
<td>Injury prevention (e.g., seat belts, helmets)</td>
<td>Physical trauma</td>
</tr>
<tr>
<td>Screen for specific risk factors</td>
<td>See Table 1.7</td>
</tr>
</tbody>
</table>

*Consider higher dose, especially for women taking antiseizure medications, other drugs that might interfere with folic acid metabolism, those with homozygous MTHFR enzyme mutations, or those who are obese.*

*By decreasing perinatal transmission, also decrease congenital defects caused by infection.*

**Abbreviations:** NTDs, neural tube defects; STD, sexually transmitted disease; MTHFR, methylenetetrahydrofolate reductase.
<table>
<thead>
<tr>
<th>Disorder</th>
<th>Chapter in MFM Evidence-Based Guidelines</th>
<th>Brief preconception recommendations</th>
<th>Prevention of</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hypertensive disorders</td>
<td>1</td>
<td>- Discontinue ACE inhibitors and ARB; transition to another antihypertensive</td>
<td>Congenital anomalies, HTN complications, CD, IUGR, placental abruption, PTB, perinatal death</td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Investigation into other etiologies</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Baseline creatinine</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Any necessary or possible cardiac interventions undergone prior to pregnancy</td>
<td>Worsening maternal cardiac condition, PTB, HTN</td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Patients with group III lesions or dilated cardiomypathy are advised not to conceive</td>
<td></td>
</tr>
<tr>
<td>Cardiac disease</td>
<td>2</td>
<td>- Counseling, diet and exercise to return to normal BMI</td>
<td>Infertility, fetal NTDs, PTB, CD, HTN disorders, diabetes, VTE</td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Evaluation of fasting lipids, fasting blood sugar</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Screening for thyroid disease, OSA, HTN</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>- If coexisting HTN or DM, obtain EKG and ECHO</td>
<td></td>
</tr>
<tr>
<td>Obesity</td>
<td>3</td>
<td>- Motivational interviewing</td>
<td></td>
</tr>
<tr>
<td>Pregestational diabetes</td>
<td>4</td>
<td>- Optimize glycemic control with goal HgbA1c &lt;7%</td>
<td>Congenital anomalies, length of NICU admission, perinatal mortality and long-term health consequences in infant; miscarriage; maternal hospitalizations, maternal renal disease</td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Screen for asymptomatic bacteriuria</td>
<td></td>
</tr>
<tr>
<td>Hypothyroidism</td>
<td>6</td>
<td>- Monitor TSH and FT4 to assure euthyroid state</td>
<td>Infertility, maternal HTN, miscarriage, preeclampsia, abortion, anemia, PTB, LBW, fetal death, possibly neurological problems in infant</td>
</tr>
<tr>
<td>Hyperthyroidism</td>
<td>7</td>
<td>- If radioiodine is required, should be completed 6–12 months before attempting conception</td>
<td>Spontaneous pregnancy loss, PTB, preeclampsia, fetal death, FGR, maternal congestive heart failure, and thyroid storm; neonatal Graves' disease</td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Emphasize minimum of 150 µg iodine daily (recommendation for all preconception women)</td>
<td></td>
</tr>
<tr>
<td>Prolactinoma</td>
<td>8</td>
<td>- Treat with dopamine agonist until decreasing size of adenoma to at least &lt;1 cm, and normal prolactin</td>
<td>Risk of increasing size of maternal prolactinoma, possibly causing optic nerve impairment</td>
</tr>
<tr>
<td>History of hyperemesis</td>
<td>9</td>
<td>- Start prenatal vitamins at 3 month prior to conception</td>
<td>Decreases risk of recurrence of hyperemesis</td>
</tr>
<tr>
<td>gravidarum</td>
<td></td>
<td>- Plan conception when disease is in remission &gt;6 months</td>
<td>Birth defects</td>
</tr>
<tr>
<td>Inflammatory bowel disease</td>
<td>11</td>
<td>- Discontinue MTX 3–6 months prior to conception</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Screen for B12, vitamin D and iron deficiency</td>
<td></td>
</tr>
<tr>
<td>Liver transplantation</td>
<td>13</td>
<td>- Plan pregnancy when stable on immunosuppressive regimen &gt;1 year</td>
<td>PTB, HTN, preeclampsia, IUGR, GDM, graft rejection</td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Assess baseline kidney/liver function, 24-hour urine</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>- If patient is on mycophenolic acid products, assess fetal risks and consider switching to alternative immunosuppressant</td>
<td></td>
</tr>
<tr>
<td>Anemia</td>
<td>14</td>
<td>- Evaluation of etiology, assessment for iron, B12, and folate deficiency</td>
<td>SGA, PTB, maternal CV compromise, need for transfusion</td>
</tr>
<tr>
<td>Sickle cell disease</td>
<td>15</td>
<td>- Start on 4 mg folic acid daily to optimize hemoglobin status</td>
<td>Birth defects, crises</td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Vaccinate with pneumococcal and influenza</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Discontinue teratogenic medications (ACE inhibitors, iron chelators)</td>
<td></td>
</tr>
<tr>
<td>von Willebrand disease</td>
<td>16</td>
<td>- Consult hematology, genetics; administer Hepatitis B vaccine</td>
<td>Postpartum hemorrhage</td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Baseline labs (von Willebrand factor antigen, ristocetin cofactor activity, factor VIII, low-dose ristocetin-induced platelet aggregation, multimer assay)</td>
<td></td>
</tr>
<tr>
<td>Disorder</td>
<td>Chapter in MFM Evidence-Based Guidelines</td>
<td>Brief preconception recommendations</td>
<td>Prevention of</td>
</tr>
<tr>
<td>----------</td>
<td>------------------------------------------</td>
<td>-------------------------------------</td>
<td>--------------</td>
</tr>
<tr>
<td>Renal disease/ transplant</td>
<td>17</td>
<td>- Assess baseline creatinine, 24-hour proteinuria, intravenous pyelogram&lt;br&gt;- Plan pregnancy when stable on immunosuppressive regimen, with drug therapies at maintenance levels if possible&lt;br&gt;- Post transplant, await &gt;1 year before conception&lt;br&gt;- Recommend deferring conception until seizure-free on minimal medication, preferably monotherapy&lt;br&gt;- Consult neurology to consider weaning medication if &gt;2 years seizure-free&lt;br&gt;- Start on folic acid 2–4 mg daily</td>
<td>Preeclampsia</td>
</tr>
<tr>
<td>Seizures</td>
<td>19</td>
<td></td>
<td>Congenital anomalies</td>
</tr>
<tr>
<td>Spinal cord injury</td>
<td>20</td>
<td>- If cause is congenital, start on folic acid 4 mg daily and genetic counseling</td>
<td>Congenital anomalies</td>
</tr>
<tr>
<td>Mood disorders</td>
<td>21</td>
<td>- Counsel on the risks of discontinuing antidepressants in pregnancy&lt;br&gt;- Stabilize mood on lowest effective dose prior to pregnancy&lt;br&gt;- Avoid Paroxetine given risks of cardiac malformations</td>
<td>Cardiac malformations</td>
</tr>
<tr>
<td>Smoking</td>
<td>22</td>
<td>- Counsel regarding preventable pregnancy outcomes in patient who smoke&lt;br&gt;- Encourage cessation with behavioral and educational interventions</td>
<td>PTB, LBW</td>
</tr>
<tr>
<td>Drug abuse</td>
<td>23</td>
<td>- Encourage patients to postpone conception until after completing detox</td>
<td>PTB, IUGR, neonatal withdrawal, etc. (Effect depends on drug of abuse)</td>
</tr>
<tr>
<td>Asthma</td>
<td>24</td>
<td>- Control of asthma with appropriate regimen through multidisciplinary care, set expectations to continue management throughout pregnancy</td>
<td>PTB, LBW, preeclampsia, perinatal mortality</td>
</tr>
<tr>
<td>Tuberculosis</td>
<td>24</td>
<td>- Screen high risk patients (hx of incarceration, TB exposure, international travel or immigration) with PPD or interferon gamma-release assay and treat accordingly</td>
<td>Active TB</td>
</tr>
<tr>
<td>Lupus</td>
<td>25</td>
<td>- Recommend conception when disease is in remission for &gt;6 months&lt;br&gt;- Screen for HTN, renal, heart, lung, or brain disease as well as antiphospholipid and SSA/SSB antibodies&lt;br&gt;- Decrease meds to lowest possible effective dose&lt;br&gt;- Replace mycophenolate mofetil and with other medications</td>
<td>HTN, preeclampsia, PTB, fetal death, IUGR, neonatal lupus</td>
</tr>
<tr>
<td>Venous thromboembolism and mechanical heart valves</td>
<td>28</td>
<td>- Screen all patients with history of VTE for thrombophilia&lt;br&gt;- Perform any necessary valve replacements before pregnancy&lt;br&gt;- If mechanical heart valve, consider continuing warfarin after full counseling of risks of warfarin embryopathy and under direction of cardiologist</td>
<td>Recurrence of venous thromboembolism</td>
</tr>
<tr>
<td>Hepatitis A</td>
<td>29</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hepatitis B</td>
<td>30</td>
<td>- Administer HBV vaccine to any woman who is susceptible before pregnancy&lt;br&gt;- If chronically infected, screen for Hepatitis A and vaccinate prior to pregnancy</td>
<td>Perinatal HBV transmission</td>
</tr>
<tr>
<td>Hepatitis C</td>
<td>31</td>
<td>- Screen high risk populations prior to pregnancy&lt;br&gt;- Vaccinate against hepatitis A and B if non-immune&lt;br&gt;- Consider treatment preconception</td>
<td>Cirrhosis, HCC, HCV infant transmission</td>
</tr>
<tr>
<td>HIV</td>
<td>32</td>
<td>- Initiate or modify antiretroviral therapy avoiding teratogenic agents (e.g., efavirenz)&lt;br&gt;- CD4 count and indicated prophylaxis based on level&lt;br&gt;- Screen for STIs&lt;br&gt;- Advise how to optimize conception, yet minimizing risk of transmission</td>
<td>Perinatal HIV infection</td>
</tr>
</tbody>
</table>

(Continued)
Table 1.7 Preconception Care for Specific Maternal Medical Disorders (Continued)

<table>
<thead>
<tr>
<th>Disorder</th>
<th>Chapter in MFM Evidence-Based Guidelines</th>
<th>Brief preconception recommendations</th>
<th>Prevention of</th>
</tr>
</thead>
<tbody>
<tr>
<td>STI testing</td>
<td>33, 34, 35, 36</td>
<td>- Screen and treat for gonorrhea, chlamydia, syphilis and trichomonas in high risk patients (e.g., &lt;25 prior STI, multiple sexual partners, inconsistent condom use, sex work, or drug use)</td>
<td>Ectopic pregnancy</td>
</tr>
<tr>
<td>PKU</td>
<td></td>
<td>- Low-phenylalanine diet</td>
<td>PKU-related mental retardation</td>
</tr>
<tr>
<td>Social issues (e.g., abuse)</td>
<td></td>
<td>- Counseling; Referral to appropriate agency</td>
<td>Physical and emotional trauma and their consequences</td>
</tr>
<tr>
<td>Alcohol</td>
<td></td>
<td>- Avoid all alcohol intake</td>
<td>Congenital anomalies, mental retardation</td>
</tr>
<tr>
<td>Supplements and over-the-counter medications</td>
<td></td>
<td>- Review and counsel: Avoid excess of recommended daily allowance (RDA) (see also Chapter 2)</td>
<td>Congenital anomalies</td>
</tr>
</tbody>
</table>

Abbreviations: ACE, angiotensin converting enzyme; ARB, angiotensin receptor blocker; HTN, hypertension; IUGR, intrauterine growth restriction; CD, cesarean delivery; PTB, preterm birth; LBW, low birth weight; NICU, neonatal intensive care unit; BMI, body mass index; DM, diabetes mellitus; EKG, electrocardiogram; ECHO, echocardiogram; OSA, obstructive sleep apnea; NTD, neural tube defects; VTE, venous thromboembolism; FGR, fetal growth restriction; TSH, thyroid-stimulating hormone; FT4, free thyroxine; CV, cardiovascular; MTX, methotrexate; TB, tuberculosis; PPD, purified protein derivative; SSA/SSB, Sjogren syndrome related antigen A and B; HIV, human immunodeficiency virus; STI, sexually transmitted infections; PKU, phenylketonuria.

unlikely. The diabetic woman with either ischemic heart disease, untreated proliferative retinopathy, creatinine clearance <50 mL/min, proteinuria >2 g/24 hours, creatinine >2 mg/dL, uncontrolled hypertension, or gastropathy should be told not to get pregnant before the above conditions can be improved, and counseled regarding adoption if the conditions cannot be improved [64]. The frequency of fetal/infant and maternal morbidity and mortality is reduced in diabetic women seeking consultation in preparation for pregnancy, but unfortunately only about one-third of these women receive such consultation [65]. The preconception consultation affords the opportunity to screen for vascular consequences of the diabetes, with ophthalmologic, electrocardiogram (EKG), and renal evaluation via a 24-hour urine collection for total protein and creatinine clearance, and determine ancillary pregnancy risks. Proliferative retinopathy should be treated with laser before pregnancy. A thyroid-stimulating hormone (TSH) level should be checked, as 40% of young women with type 1 diabetes have subclinical hypothyroidism. Of note, there is insufficient evidence to treat subclinical hypothyroidism [66].

Diabetes evaluation should emphasize the importance of tight glycemic control, with normalization of the HgbA1c to at least <7%. To achieve euglycemia, diet, glucose monitoring, and exercise are always stressed. If euglycemia is not achieved with these means, oral hypoglycemic agents or insulins are utilized, and their regimens should be optimized preconceptionally. Of the oral hypoglycemic agents, glyburide and glipizide can be used, and probably continued during pregnancy. The original safety data available for glyburide showed that it did not cross the placenta in appreciable amounts [67], but recent data have shown a 70% level in umbilical blood compared with maternal blood [68]. The other oral hypoglycemic agents should not be used for preconception glycemic control, as there is no sufficient evidence for their safety and efficacy in pregnancy. A common insulin regimen currently used by diabetologists is long-acting (e.g., glargine) and short-acting (e.g., lispro). This is a safe and effective regimen in pregnancy, too. Women compliant with insulin pumps should continue this regimen.

If a woman has a history of gestational diabetes, appropriate postpartum diabetes screening should be performed. Interconception counseling and lifestyle modifications may be beneficial for future pregnancies [69].

**Hypertension** (see Chapter 1 in *Maternal-Fetal Evidence Based Guidelines*) is associated with several maternal [worsening hypertension; superimposed preeclampsia; severe preeclampsia; eclampsia; hemolysis, elevated liver enzyme levels, and a low platelet count (HELLP) syndrome; cesarean delivery] and fetal (growth restriction; oligohydramnios; placental abruption; PTB; perinatal death) risks in pregnancy. Serum creatinine, 24-hour urine for total protein and creatinine clearance, EKG, and ophthalmologic exam are suggested, especially in women with long-standing or severe hypertension. It is important to identify cardiovascular risk factors and any reversible cause of hypertension, as well as assess for target organ damage or cardiovascular disease. If hypertension is newly diagnosed and has not been evaluated previously, a medical consult may be indicated to assess for any of these factors. Secondary hypertension, target organ damage (left ventricular dysfunction, retinopathy, dyslipidemia, microvascular disease, and prior stroke), maternal age >40, previous pregnancy loss, systolic blood pressure ≥180 mmHg, or diastolic blood pressure ≥110 mmHg are associated with higher risks in pregnancy. Abnormalities should be addressed and managed appropriately. If, for example, serum creatinine is >1.4 mg/dL, the woman should be aware of increased risks in pregnancy (pregnancy loss, reduced birth weight, PTB, and accelerated deterioration of maternal renal disease). Even mild renal disease (creatinine 1.1–1.4 mg/dL) with uncontrolled hypertension is associated with tenfold higher risk of fetal loss. Preconception prevention can be enormously effective. Thirty minutes of exercise five times per week in all women with hypertension and weight reduction if overweight are recommended. Restriction of sodium intake to the same <2.4 g sodium daily intake recommended for essential hypertension is beneficial in nonpregnant adults. If antihypertensive medical therapy is necessary, angiotensin-converting enzyme (ACE) inhibitors and angiotensin II (AII) receptor antagonists should be discontinued as they are associated with birth defects, fetal growth restriction,
oligohydramnios, neonatal renal failure, and neonatal death in pregnancy. All other antihypertensive agents should be used at the lowest effective dose and are probably safe if started preconceptionally and continued in pregnancy.

Seizure Disorders
Conception should be deferred until seizures are well controlled on the minimum effective dose of medication (see Chapter 19 in Maternal-Fetal Evidence Based Guidelines). Monotherapy is preferable. Lamotrigine has been reported to be the first-line therapy for nonpregnant adults for partial seizures [70–72] and is associated with a low incidence of major malformations [73], but not in all studies [74]. The best choice is the antiepileptic drug (AED) that best controls the seizures. The AEDs are usually U.S. Food and Drug Administration (FDA) category C (human risk unknown, but none proven yet) except for the following AEDs that are known potential teratogens: carbamazepine, primidone, phenytoin, and valproate (Table 1.8). These four AEDs should therefore be avoided if possible, by using a different therapy beginning in the preconception period. Women who have been seizure-free for ≥2 years with a normal electroencephalogram (EEG) may be eligible to stop anticonvulsant therapy after consulting with a neurologist [75].

Medications/Teratogens
Detailed discussion regarding prescribed and over-the-counter medications should occur at the preconception visit. The indication, safety, effectiveness, and necessity of each drug need to be reviewed. Often, women and their doctors stop efficacious and necessary medications as soon as the woman finds out she is pregnant, compromising the health of both the woman and her baby. The vast majority of prescribed medications are safe in pregnancy, even in the first trimester. Only a few drugs, chemicals, infections, or reactions are known teratogens (Table 1.8) [76,77]. These should be avoided, except in rare circumstances (e.g., the woman with mechanical cardiac valves who accepts the teratogenic risk of warfarin). This medication counseling is often a crucial part of preconception care and can save and ameliorate significantly the health of a future offspring. Great resources exist on the Web for up-to-date teratologic information [78–80].

### Table 1.8 Teratogens

<table>
<thead>
<tr>
<th>Prescribed drugs</th>
<th>Chemicals</th>
<th>Drugs of abuse</th>
<th>Infections</th>
<th>Radiation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Androgens and testosterone derivatives (e.g., danazol)</td>
<td>Lithium</td>
<td>Alcohol</td>
<td>Cytomegalovirus</td>
<td>Radiation</td>
</tr>
<tr>
<td>Angiotensin-converting enzyme (ACE) inhibitors (e.g., enalapril, captopril) and angiotensin II receptor blockers</td>
<td>Phenyltoin</td>
<td>Cocaine</td>
<td>Rubella</td>
<td>Ovarian cyst</td>
</tr>
<tr>
<td>Coumadin derivatives (e.g., warfarin)</td>
<td>Primidone</td>
<td>Infections</td>
<td>Syphilis</td>
<td>Radiation</td>
</tr>
<tr>
<td>Carbamazepine</td>
<td>Streptomycin and kanamycin</td>
<td>Infections</td>
<td>Toxoplasmosis</td>
<td>Varicella</td>
</tr>
<tr>
<td>Diethylstilbestrol</td>
<td>Tetracycline</td>
<td>Infections</td>
<td></td>
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</tr>
<tr>
<td>Folic acid antagonists (methotrexate and aminopterin)</td>
<td>Thalidomide and leflunomide</td>
<td>Infections</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HMG-CoA reductase inhibitors (statins)</td>
<td>Trimethadione and paramethadione</td>
<td>Infections</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lithium</td>
<td>Valproic acid</td>
<td>Chemicals</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Phenyltoin</td>
<td>Vitamin A above RDA, and its derivatives (e.g., isotretinoin, etretinate, and retinoids)</td>
<td>Drugs of abuse</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Primidone</td>
<td></td>
<td>Infections</td>
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<tr>
<td>Streptomyacin and kanamycin</td>
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<tr>
<td>Tetracycline</td>
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<tr>
<td>Thalidomide and leflunomide</td>
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<tr>
<td>Trimethadione and paramethadione</td>
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<tr>
<td>Valproic acid</td>
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<tr>
<td>Vitamin A above RDA, and its derivatives (e.g., isotretinoin, etretinate, and retinoids)</td>
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</tbody>
</table>


### Substance Abuse/Environmental Hazards/Toxins
Tobacco smoking during pregnancy is associated with increased risks of several complications (see Chapter 22 in Maternal-Fetal Evidence Based Guidelines). The benefits of smoking cessation are tremendous: prevention of 10% of perinatal deaths, 35% of low-birth weight births, and 15% of preterm deliveries [81]. Smoking only one to five cigarettes per day is associated with a 55% higher incidence of low birth weight compared with nonsmokers. Reproductive-age women should be informed of other smoking-related diseases, such as ischemic heart disease, cancer, lung diseases, pneumonia, stroke, and congestive heart failure. Women at greatest risk for smoking are those <25 years old with less than a high school education. Smoking makes a major contribution to disparities in mortality [82]. Smoking cessation programs are associated with a 6% increase in smoking cessation, and decreases in incidences of low birth weight (by 19%) and PTB (by 16%) [83]. Support and reward techniques to help quit smoking are one of the best form of evidence-based medicine, supported by over 20 high-quality randomized trials. The “5 As” for screening and interventions to prevent smoking in pregnancy are Ask, Advise, Assess, Assist, and Arrange [67]. Counseling with behavioral and educational interventions is associated with highest cessation rates. If necessary, most pharmacotherapies are effective preconception, but contraindicated or with uncertain safety and efficacy during pregnancy. Nicotine replacement therapy (e.g., patch, gum, and bupropion) is safe and effective in reproductive-age women, but there is insufficient evidence for recommending them in pregnant smokers. Nicotine replacement therapy is associated with known adverse fetal effects, and nicotine is detected in breast milk. Possibly the best prevention of the adverse effects of smoking on pregnancy is achieved by avoiding sale of tobacco to young people, prohibition of smoking in public places, increase in tobacco taxation, workplace smoking cessation programs, and banning of tobacco sponsorship of sporting and cultural events.

Numerous recreational drug exposures have adverse pregnancy effects (see Chapter 23 in Maternal-Fetal Evidence Based Guidelines). This list is extensive and includes, but not limited to, common recreational drugs such as alcohol, cannabinoids, cocaine, heroin, and methamphetamines. Working to ensure that women with substance abuse issues engage in safe sex practices and family planning is a constant challenge, and these women are disproportionately overrepresented among women with unplanned pregnancies.
REFERENCES


KEY POINTS

• Prenatal care is of benefit to pregnant women, especially those with modifiable risk factors.
• Most low-risk women can be offered midwife-led models of care, and women should be encouraged to ask for this option. Continuity of care by midwives has been associated with improved patient satisfaction. Caution should be exercised in applying this advice to women with substantial medical or obstetric complications.
• Group prenatal care should be promoted as it has been associated reduction in preterm birth (PTB), greater satisfaction with care, and higher breastfeeding initiation. In the developing world, participatory intervention with women's groups is associated with decreased maternal and neonatal mortality.
• Women should be allowed to carry their record.
• Prenatal care usually consists of 7–12 visits per pregnancy, with a first prenatal visit soon after the pregnancy test is positive, and in time to establish location and number of embryo(s), usually at around 6–8 weeks, then at 11–14 weeks for aneuploidy screening, followed by visits about every 4 weeks approximately at 16, 20, 24, and 28 weeks; about every 2 weeks from 34 to 36 weeks, then weekly until delivery. In settings with limited resources where the number of visits is already low, reduced visits programs of antenatal care (<5) are associated with an increase in perinatal mortality compared with standard care.
• See Table 2.1 for screening and interventions at different times in pregnancy.
• Early ultrasonography should be used to determine the estimated date of confinement (EDC) if there is any uncertainty regarding last menstrual period (LMP).
• Content issues that should be included in prenatal care are lifestyle, nutrition, supplements, vaccinations, drugs, environment, prenatal education, and others.
• Regular aerobic exercise for 35–90 minutes 3–4 times per week during pregnancy is beneficial to overall maternal fitness and sense of well-being, as well as associated with prevention of excessive weight gain and higher chance of vaginal delivery.
• Most studies report that sexual activity is associated with better pregnancy outcomes, probably because women who are sexually active are healthier to begin with compared with women with less sexual activity.
• Balanced nutrition and protein supplementation is associated with modest increases in maternal weight gain and in mean birth weight, and reduction in risk of small-for-gestational-age (SGA), stillbirth, and neonatal death. High-protein and isocaloric protein supplementation should be avoided as they are associated with increased risk of SGA.
• Suggested weight gain in pregnancy is shown in Table 2.4. Women who are underweight are at increased risk for low birth weight (LBW) and PTB and have better outcomes with a higher total weight gain. Excessive weight gain in women with normal body mass index (BMI) can be prevented with dietary and lifestyle counseling.
• Folic acid supplementation is recommended for neural tube defect (NTD) prevention, with 400 μg/day for all women, and 4 mg/day for women with prior children with NTD. All reproductive-age women should be on folic acid (FA) supplementation.
• Immunity to rubella, varicella, hepatitis B, influenza, tetanus, and pertussis should be assessed at the first prenatal visit. Ideally needed vaccinations should be provided preconception. Influenza vaccine is recommended for pregnant women during flu season. Tetanus, diphtheria, and acellular pertussis vaccine, also known as TDAP vaccine, is recommended for all pregnant women after 28 weeks. Partners and family members should be encouraged to be vaccinated as well.
• Prenatal education directed at specific objectives has been demonstrated to be effective.
• Implementation of community-based interventional care packages is associated with a trend for reduction in maternal mortality, and with significant reductions in maternal morbidity, neonatal mortality, stillbirths, and perinatal mortality.
• Perineal massage with sweet almond oil for 5–10 minutes daily from 34 weeks until delivery is associated with a significantly higher chance of intact perineum in nulliparous women.
• Antenatal classes with training to prepare for labor and delivery are associated with arriving to labor and delivery (L & D) ward more often in active labor, and less use of epidural analgesia.
• Identifying mothers at risk for postpartum depression assists in prevention compared with intervening on the general population.
• Breastfeeding is the best feeding method for most infants and should be strongly encouraged. Continued counseling and education facilitate breastfeeding success.
• Unsensitized RhD-negative women should be offered anti-D immunoglobulin prophylaxis.
• Sweeping or “stripping” of membranes during cervical exam at ≥38 weeks reduces the rate of postterm delivery.
• Magnesium lactate or citrate chewable tablets 5 mmol in the morning and 10 mmol in the evening for 3 weeks for women with leg cramps are associated with significant improvement in persistent leg cramps.
• Water gymnastics for 1 hour weekly starting at <19 weeks reduces back pain in pregnancy and allows more women to continue to work, with no adverse effects. Both
Table 2.1  Suggested Prenatal Care Counseling, Screening, and Intervention

<table>
<thead>
<tr>
<th>Initial visit ≤14 weeks</th>
<th>Visits at: 14–24 weeks</th>
<th>24–28 weeks</th>
<th>28–34 weeks</th>
<th>34–41 weeks</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Assessments/procedures</strong></td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>• Complete history and risk identification</td>
<td>• Fetal heart tones</td>
<td>• Fetal heart tones</td>
<td>• Fetal heart tones</td>
<td>• Fetal heart tones</td>
</tr>
<tr>
<td>• Assessment of EDB by LMP and sizing; ultrasound if indicated</td>
<td>• Fundal height</td>
<td>• Fundal height</td>
<td>• Fundal height</td>
<td>• Fundal height/ EFW</td>
</tr>
<tr>
<td>• Baseline BP screening</td>
<td>• Fetal movement</td>
<td>• Fetal movement</td>
<td>• Fetal movement</td>
<td>• Fetal movement</td>
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<tr>
<td>• Weight and BMI</td>
<td>• BP</td>
<td>• BP</td>
<td>• BP</td>
<td>• BP</td>
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<tr>
<td>• Screening for domestic abuse</td>
<td>• Weight</td>
<td>• Weight</td>
<td>• Weight</td>
<td>• Weight</td>
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<tr>
<td>• Vaccines according to risk status and season</td>
<td>• Screening ultrasound for anatomy</td>
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<td>• Referral for specialist care according to history</td>
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<tr>
<td>• Offer 11–13 6/7 weeks aneuploidy screening ultrasound</td>
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<tr>
<td>• Multiple-marker aneuploidy screen</td>
<td>• Fetal heart tones</td>
<td>• Gestational diabetes screen; repeat CBC and antibody screen</td>
<td>• Urine dipstick for protein</td>
<td>• Group B Strep</td>
</tr>
<tr>
<td></td>
<td>• CBC; blood type, Rh, antibody screen; Rubella IgG; RPR; HBsAg; HIV</td>
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<td>• Rh immunoglobulin if indicated</td>
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<td></td>
<td>• Urine dipstick for protein if indicated</td>
<td>• Antibody screen if indicated</td>
<td>• Urine dipstick for protein</td>
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<td></td>
<td>• Urinalysis and urine culture</td>
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<td></td>
<td>• Gonorrhea/chlamydia</td>
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<td>• Papa</td>
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<td></td>
<td>• Additional testing as directed by history and PEa</td>
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<td><strong>Laboratory tests</strong></td>
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<tr>
<td>• Multiple-marker aneuploidy screen</td>
<td>• Review and discuss results of testing</td>
<td>• Preterm labor s/sx</td>
<td>• Preterm labor s/sx</td>
<td>• Labor symptoms/when to call</td>
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<td></td>
<td>• Urine dipstick for protein if indicated</td>
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<tr>
<td></td>
<td>• Gestational diabetes screen</td>
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<td></td>
<td>• Antibody screen if indicated</td>
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<td>• Urine dipstick for protein</td>
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<tr>
<td><strong>Education/counseling</strong></td>
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<tr>
<td>• Cessation of harmful substances</td>
<td>• Preterm labor s/sx</td>
<td>• Preterm labor s/sx</td>
<td>• Post-dates management</td>
<td>• Breastfeeding</td>
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<tr>
<td>• Exercise/activity</td>
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<td>• Nutrition</td>
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<td>• Weight gain</td>
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<td>• Supplements</td>
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<td></td>
<td>• Food safety</td>
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<tr>
<td></td>
<td>• Breastfeeding</td>
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</tbody>
</table>

**Education/counseling not limited to specific weeks gestation**

| | | | | |
| • Danger signs | | | | |
| • Dental care | | | | |
| • Family planning | | | | |
| • Labor preparation, options, s/sx to report | | | | |
| • Travel | | | | |
| • TOLAC | | | | |

**Assessments/procedures**

| | | | | |
| • Complete history and risk identification | • Fetal heart tones | • Fetal heart tones | • Fetal heart tones | • Fetal heart tones |
| • Assessment of EDB by LMP and sizing; ultrasound if indicated | • Fundal height | • Fundal height | • Fundal height | • Fundal height/ EFW |
| • Baseline BP screening | • Fetal movement | • Fetal movement | • Fetal movement | • Fetal movement |
| • Weight and BMI | • BP | • BP | • BP | • BP |
| • Screening for domestic abuse | • Weight | • Weight | • Weight | • Weight |
| • Vaccines according to risk status and season | • Screening ultrasound for anatomy | • Screening ultrasound for anatomy | • Screening for domestic abuses | • Screening for domestic abuses |
| • Referral for specialist care according to history | | | | |
| • Offer 11–13 6/7 weeks aneuploidy screening ultrasound | | | | |

**Sources:** Adapted from a review of current prenatal care guidelines from four major groups: U.S. Veterans Health Administration, Department of Veteran Affairs, and Health Affairs, Department of Defense; Institute for Clinical Systems Improvement; the American Academy of Pediatrics and the American College of Obstetricians and Gynecologists; and the American Academy of Family Physicians; Hanson L et al., J Midwifery Women's Health, 54(6), 458–468, 2009.

*aSee text, only in certain circumstances.*

**Abbreviations:** EDB, expected date of birth; LMP, last menstrual period; BP, blood pressure; BMI, body mass index; EFW, estimation of fetal weight; RPR, rapid plasma regain; HIV, human immunodeficiency virus; CBC, complete blood count; PE, physical exam; TOLAC, trial of labor after cesarean; s/sx, signs and symptoms.
physiotherapy and acupuncture starting <32 weeks for 10 sessions might reduce back and pelvic pain.

- Exercise, increase in water intake, dietary counseling, and certain foods (e.g., prunes) have shown relief in constipation. If these self-help measures are inadequate, the pregnant woman should then try daily bran or wheat fiber supplements. Docusate sodium is an effective stimulant laxative.

DEFINITION

Prenatal care is the care provided to pregnant women with the aim to prevent complications and decrease the incidence of perinatal and maternal morbidity and mortality [1]. This care consists of health promotion, risk assessment, and intervention linked to the risks and conditions uncovered. These activities require the cooperative and coordinated efforts of the woman, her family, her prenatal care providers, and other specialized providers. Prenatal care begins when conception is first considered and continues until labor begins. The objectives of prenatal care for the mother, infant, and family relate to outcomes through the first year following birth [1].

PURPOSE

Prenatal care developed, historically, to reduce the incidence of LBW and preterm infants [2]. It has evolved to encompass a broader purpose; to identify pregnancies with maternal or fetal conditions associated with morbidity/mortality, to provide interventions to prevent or treat such complications, and to provide education support and health promotion that can have lasting effects on the health of an entire family [3]. Care should be systematic, evidence-based, and should result in informed shared decision making between the patient and the provider.

EFFECTIVENESS

Prenatal care is of benefit to pregnant women. Nonetheless, the value of prenatal care is controversial, as there is no definite evidence that prenatal care improves birth outcomes. There are no randomized control trials (RCTs) of prenatal care versus no prenatal care. Most studies are observational. Selection bias (women who self-select to prenatal care usually are more inclined to have better outcomes) leads to confounding bias (e.g., risk factors associated with LBW and neonatal death are also risk factors for inadequate prenatal care).

There are several RCTs on the number of prenatal care visits, which indirectly demonstrate the beneficial effects of prenatal care. There is a higher incidence of perinatal mortality (relative risk [RR] 1.14, 95% confidence interval [CI] 1.00–1.31) in programs with significantly less (<5) numbers of prenatal visits, compared with the usual 8–12. This is particularly significant for low- and middle-income countries [4]. Also, studies demonstrate a reduction in poor outcomes in high-risk pregnancies with enhanced prenatal care at no added cost [4] (see Section “Number and Timing of Visits”). In addition, women are dissatisfied with a reduced schedule of prenatal visits indicating a perceived benefit by women [4]. Specific interventions for specific risks may reduce morbidity and mortality. Prenatal care is probably of most benefit to medically high-risk women [2].

ORGANIZATIONAL ISSUES

Health-Care Provider

There is no evidence that physicians need to be involved in the prenatal care of every woman experiencing an uncomplicated pregnancy. The effect of midwife-led care compared with physician-led care or to other provider-led care has been evaluated mostly for the whole pregnancy, including together both antepartum care and care during labor and delivery (see also Chapter 7). Therefore it is difficult to assess the effect of midwife-led care just on antepartum care. From the evidence from both antepartum and L and D care, most women can be offered midwife-led models of care and women should be encouraged to ask for this option. Caution should be exercised in applying this advice to women with substantial medical or obstetric complications. In a meta-analysis, women, the vast majority low risk, who had midwife-led models of care, were less likely to experience antenatal hospitalization, and less likely to experience fetal loss before 24 weeks’ gestation (RR 0.79, 95% CI 0.65–0.97), although there were no statistically significant differences in fetal loss/neonatal death of at least 24 weeks (RR 1.01, 95% CI 0.67–1.53) or in fetal/neonatal death overall (RR 0.83, 95% CI 0.70–1.00) [5] (see also Chapter 7). It is not clear whether these associations are due to greater continuity of care or to midwifery care [5].

Group Prenatal Care

In a meta-analysis, educational interventions were the focus of group prenatal care, and no consistent results were found. Sample sizes were very small to moderate. No data were reported concerning anxiety, breastfeeding success, or general social support. Knowledge acquisition, sense of control, factors related to infant-care competencies, and some labor and birth outcomes were measured. The largest of the included studies (n = 1275) examined an educational and social support intervention to increase vaginal birth after cesarean delivery. This high-quality study showed similar rates of vaginal birth after cesarean delivery in “verbal” and “document” groups (RR 1.08, 95% CI 0.97–1.21) [6]. One large RCT demonstrated significant reduction in PTB, greater satisfaction with care, and higher breastfeeding initiation at no added cost for group prenatal care over standard care in a group of medically low-risk (but socially at-risk) women in an urban clinic [7]. In this study, group care included, among other interventions, continuity of care from a single provider, patient keeping copies of their records, no waiting time at visits, about 20 hours of provider/patient time, with 8–10 women in each group session. In the developing world, participatory intervention with women’s groups is associated with decreased maternal and neonatal mortality in several large cluster-randomized trials [8–10]. In one of these studies, participatory care involved a female facilitator convening nine women’s group meetings every month. The facilitator supported groups through an action–learning cycle in which they identified local perinatal problems and formulated strategies to address them [8]. This strategy holds great promise in decreasing maternal and perinatal deaths among the most vulnerable in our world.

Group prenatal care may even be utilized in a higher risk population. In a non-RCT study, group prenatal care for women with gestational diabetes (GDM) is associated with decreased progression to A2 gestational diabetes and improved postpartum follow-up for appropriate diabetes screening without significantly affecting obstetrical or neonatal outcomes [11].
Group prenatal care should be promoted and further studied among more diverse populations.

Prenatal Record
A formal, structured record should be used for documenting care during the pregnancy. Structured records with reminder aids help ensure that providers incorporate evidence-based guidelines into clinical practice. There is no trial comparing different records. Women should be allowed to carry their record. A meta-analysis of three trials showed that carrying the record is associated with increased maternal control and satisfaction during pregnancy, increased availability of antenatal records during hospital attendance, but also with more operative deliveries. Importantly, all of the three trials included in the meta-analysis report that more women in the case notes group would prefer to hold their antenatal records in another pregnancy [12].

Number and Timing of Visits
There is insufficient evidence to recommend an ideal schedule of prenatal visits for all pregnant women. The most important visit to optimize pregnancy outcomes is the preconception visit (see Chapter 1). A visit early, soon after the pregnancy test is positive, and in time to establish location and number of embryo(s), usually around 6–8 weeks, is also desirable. At this early visit, each woman should be assessed for risk factors (see Tables 1.3 and 1.4 in Chapter 1). The frequency of subsequent visits can be determined based on risk factors.

In developed countries, prenatal care usually consists of 7–12 visits per pregnancy, with a prenatal visit ideally at 10–14 weeks for aneuploidy screening (see Chapters 5 and 6), followed by visits about every 4 weeks approximately at 16, 20, 24, and 28 weeks; about every 2 weeks from 32 to 36 weeks, then weekly until delivery (Table 2.1) [13]. Uncomplicated multiparous women may need fewer visits than uncomplicated nulliparous ones. Individual patient needs and risk factors should be assessed at the first prenatal visit and reassessed at each appointment thereafter.

A small reduction in the traditional number of prenatal visits in both developed and developing countries has not been associated with adverse biological maternal or perinatal outcomes, but women may feel less satisfied with fewer visits [4]. But, in settings with limited resources where the number of visits is already low, reduced antenatal visits (<5) are associated with an increase in perinatal mortality compared with standard care, although admission to neonatal intensive care may be reduced [4]. Women prefer the standard visits schedule. Where the standard number of visits is low, visits should not be reduced without close monitoring of fetal and neonatal outcome [4]. In addition, women in high-resource settings were more often dissatisfied with a reduced schedule of visits (defined as eight). The schedule of visits should be determined by the purpose of the appointment. A minimum of four prenatal care visits is recommended even for low-risk women [4].

### STRUCTURE

#### Initial Visit

Ideally, this visit should occur prior to 12 weeks of gestation. Women should receive written information regarding their pregnancy care services, the proposed schedule of visits, screening tests that will be offered, and lifestyle issues, such as nutrition and exercise. Major parts of the visit include history, risk identification, physical examination, laboratory testing, education for health promotion, and a detailed plan of care for any risks identified (see Table 2.1) (see also Chapter 1, Tables 1.2–1.5).

#### History

A comprehensive history should be performed, preferably using standardized record forms (e.g., www.acog.org). Risk assessment should be performed with detailed review of systems. In particular, the woman who may require additional care or referral should be identified. Early ultrasonography should be used to determine the EDC if there is any uncertainty regarding LMP [14]. Accuracy of EDC is critical for timing of screening tests and appropriate interventions, managing complications, and consideration of delivery timing. It also provides early identification and chorionicity of multiple pregnancies (see Section “Ultrasonography” and Chapter 4). Content issues such as lifestyle, nutrition, supplements, drugs, environment, vaccinations, prenatal education, and others should be discussed (see Section “Content of Prenatal Care”). Prenatal diagnosis and screening for aneuploidy (Chapter 5) and genetic screening (Chapter 6) should be reviewed.

#### Physical Exam

The physical exam should be both general (Table 2.1) and directed by any risks identified in the history (see Chapter 1).

**Weight and height** should be determined at the initial prenatal visit in order to determine BMI (BMI = weight (kg)/height squared [m²]). BMI should be based on weight at time of conception or the earliest known weight in pregnancy. Categories of BMI are in Table 2.2. Women with obesity are at increased risk for diabetes, shoulder dystocia, cesarean section, and other complications, and have better outcomes with a lower (or no) total weight gain. Women who are underweight (<50 kg or <120 lb.) also are at increased risk for LBW and PTB, and have better outcomes with a higher total weight gain (see Section “Nutrition”).

**Blood pressure** is recommended at each prenatal visit. Initial blood pressure evaluation may help to identify women with chronic hypertension, while subsequent blood pressure readings aid in preeclampsia screening. A diastolic blood pressure of >80 at booking is associated with later risks of preeclampsia [15]. There are significant risks associated with both hypertension and preeclampsia in pregnancy. This simple, inexpensive, and widely accepted screening tool may help to identify abnormal trends in blood pressure over time. Blood pressure should be taken in the sitting position using an appropriately sized cuff and correct technique (see Chapter 1 in Maternal-Fetal Evidence Based Guidelines).

#### Pelvic Examination

Routine pelvic examination early in pregnancy is not as accurate for assessment of gestational age compared with ultrasound (see Chapter 4) and not a reliable predictive test of PTB.

#### Table 2.2 Body Mass Index (BMI) Categories

<table>
<thead>
<tr>
<th>Weight category</th>
<th>BMI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Underweight</td>
<td>&lt;18.5</td>
</tr>
<tr>
<td>Normal weight</td>
<td>18.5–24.9</td>
</tr>
<tr>
<td>Overweight</td>
<td>25–29.9</td>
</tr>
<tr>
<td>Obesity (class I)</td>
<td>30–34.9</td>
</tr>
<tr>
<td>Obesity (class II)</td>
<td>35–39.9</td>
</tr>
<tr>
<td>Extreme obesity (class III)</td>
<td>&gt;40</td>
</tr>
</tbody>
</table>
or cephalopelvic disproportion later in pregnancy (see also Chapters 7 and 17), and so it is not recommended for these assessments. Abdominal and pelvic examination to detect gynecologic pathology can be included in the initial examination, with no level 1 evidence for effectiveness of this screening test.

**Laboratory Screening**

Recommended initial universal laboratory screening is listed in Table 2.1. Other lab testing may be ordered if other risks/conditions are present.

**ABO/Rh (D) type and antibody screen.** Testing for blood group, Rh status, and atypical red cell antibodies at the initial visit is recommended. Unsensitized RhD-negative women should be offered anti-D immunoglobulin at 28 weeks (see Chapter 53 in *Maternal-Fetal Evidence Based Guidelines*). Anti-D immunoglobulin should also be offered for any invasive procedure (e.g., amniocentesis, chorionic villus sampling (CVS), percutaneous umbilical blood sampling [PUBS]), second- or third-trimester bleeding, partial molar pregnancies, spontaneous abortion, elective termination, and any condition that might be associated with fetal-maternal hemorrhage, such as abdominal trauma, external cephalic version, or placental abruption. It may also be offered for any first-trimester threatened abortion and ectopic pregnancy, although the evidence is not as strong, and it is probably not cost-effective or necessary unless the bleeding is significant. For the RhD-negative woman in a known RhD-negative father of the pregnancy, anti-D immunoglobulin can be deferred. Du-positive women do not need anti-D immunoglobulin (see Chapter 53 in *Maternal-Fetal Evidence Based Guidelines*).

**Complete blood count.** Recommended at the first prenatal visit to identify anemia (hemoglobin and hematocrit) and to screen for thalassemia (mean corpuscular volume [MCV]). Pregnant women identified with anemia (Hgb < 11.0 g/dL in first trimester) should be treated as per Chapter 14 in *Maternal-Fetal Evidence Based Guidelines*. Initial determination of platelet count (optimally also before pregnancy) may help identify chronic thrombocytopenias and aid in diagnosis of gestational thrombocytopenia or HELLP (hemolysis, elevated liver enzyme levels, and a low platelet count) syndrome later in pregnancy.

**Rubella antibody.** Screen all women at first encounter. Nonimmune pregnant women should be counseled to avoid exposure and seek immunization postpartum (see Chapter 38 in *Maternal-Fetal Evidence Based Guidelines*).

**Syphilis screening.** All pregnant women should be screened with a serologic test for syphilis at the first prenatal visit. Women who are at high risk, live in areas of high syphilis morbidity, or are previously untested should be screened at 28 weeks and again at delivery (see Chapter 35 in *Maternal-Fetal Evidence Based Guidelines*).

**HBsAg.** Screen at initial encounter, and rescreen high-risk populations in third trimester. Postnatal intervention is recommended in all HBsAg-positive women to reduce the risk of viral transmission to the neonate. Pregnancy and breastfeeding are not contraindications to immunization in women who are at risk for acquisition of the hepatitis B virus (see Chapter 30 in *Maternal-Fetal Evidence Based Guidelines*).

**HIV serology.** Screening is recommended for all pregnant women. The "opt-out" approach is recommended. It should be emphasized that testing not only provides the opportunity to maintain maternal health, but interventions can be offered to dramatically reduce the risk of viral transmission to the fetus (see Chapter 32 in *Maternal-Fetal Evidence Based Guidelines*).

**Urine dipstick for protein.** Screening for proteinuria should occur at the initial visit and routinely after 20 weeks in women at risk for preeclampsia. Urine dipsticks for protein do not reliably detect the variable elevations in albumin that may occur in preeclampsia and may not be indicated at each visit in low-risk women [16]. In women at high risk for preeclampsia, the 24-hour collection is a reasonable screen for proteinuria as a baseline at the first prenatal visit, and when other signs/symptoms of preeclampsia are present. The proteinuria/creatinine (P/C) ratio may be used as a screening test as a good predictor for remarkable proteinuria since it seems to be highly predictive for diagnosis to detect proteinuria over one gram but inadequate in detecting lower levels [17] (see Chapter 1 in *Maternal-Fetal Evidence Based Guidelines*).

**Urine dipstick for glucose.** Glicosuria ≥250 mg/dL (equivalent to 1+) on urine dipstick in the first or second trimester is associated with abnormal GDM screening later in pregnancy. Presence of significant glycosuria before 24–28 weeks is an indicator for earlier gestational glucose screening (see Chapters 4 and 5 in *Maternal-Fetal Evidence Based Guidelines*).

**Urine culture for asymptomatic bacteriuria.** Screening for bacteriuria is recommended at the first prenatal visit for all women. Pregnant women with asymptomatic bacteriuria are at increased risk for symptomatic infection and pyelonephritis. There is also a positive relationship between untreated bacteriuria and LBW/PTB. Treatment of asymptomatic bacteriuria prevents these complications (see Chapter 17 in *Maternal-Fetal Evidence Based Guidelines*).

**Cervical cancer screening.** Cervical cancer screening should be obtained if not current according to guidelines. Pap smear screening should be initiated at age 21, regardless of onset of sexual activity. Routine screening intervals have also been extended to every 3 years for women in their 20s without human papillomavirus (HPV) co-testing and every 5 years in women over 30 with the addition of HPV co-testing. Colposcopy can be performed during pregnancy and a plan can be made for treatment postpartum (see Chapter 31).

**Selective (Only Women with Risk Factors) Laboratory Screening**

**Hepatitis C serology.** A test for hepatitis C antibodies should be performed in pregnant women at increased risk for exposure, such as those with a history of IV drug abuse, exposure to blood products or transfusion, organ transplants, kidney dialysis, etc. (see Chapter 31 in *Maternal-Fetal Evidence Based Guidelines*).

**Chlamydia screening.** All women of age <25 years (strongest risk factor), multiple sex partners, new partner within past 3 months, single marital status, inconsistent use of barrier contraception, previous or concurrent sexually transmitted infection (STI), vaginal discharge, mucopurulent cervicitis, friable cervix, or signs of cervicitis on physical examination should be screened. Some agencies advocate universal chlamydia screening. Rescreen in the third trimester if at increased risk for infection. Screening using polymerase chain reaction (PCR) technology is most accurate (see Chapter 34 in *Maternal-Fetal Evidence-Based Guidelines*).

**Gonorrhea screening.** All women of age <25 years, prior STI, multiple sexual partners, having a partner with a past history of any sexually transmitted disease (STD), sex work, drug use, and inconsistent condom use should be screened for gonorrhea. Some agencies advocate universal gonorrhea screening. Rescreen in the third trimester if at increased risk for infection. Screening using PCR technology is most accurate (see Chapter 33 in *Maternal-Fetal Evidence Based Guidelines*).
Bacterial vaginosis. There is no benefit to routine screening and treatment for asymptomatic bacterial vaginosis. Consideration can be given to screening and treating women with a prior PTB, but given the inconclusive evidence we do not recommend it as routine. However, those women who are symptomatic should be screened (see Chapter 17 in Maternal-Fetal Evidence Based Guidelines).

Genital herpes. Routine serologic or other screening for herpes simplex virus (HSV) in asymptomatic pregnant women is not recommended. In the absence of lesions during the third trimester, routine serial cultures are not indicated for women with a history of recurrent genital herpes (see Chapter 50 in Maternal-Fetal Evidence Based Guidelines).

Varicella. Screening is indicated if a woman has had neither past infection nor vaccination. Varicella vaccine (live attenuated) is not recommended during pregnancy, but seronegative women should be advised to take appropriate precautions (see Chapters 38 and 51 in Maternal-Fetal Evidence Based Guidelines).

Tuberculosis. Quantiferon gold or purified protein derivative (PPD) can be offered to high-risk women at any gestational age in pregnancy to screen for tuberculosis, and follow-up chest x-ray is recommended for recent converters. High-risk factors include human immunodeficiency virus (HIV) disease, homeless or impoverished women, prisoners, recent immigrants from areas where tuberculosis is prevalent, and others (see Chapter 24 in Maternal-Fetal Evidence Based Guidelines).

Cytomegalovirus (CMV). Routine testing is not recommended. Good hand washing and practicing universal precautions are recommended to prevent transmission [18] (see Chapter 47 in Maternal-Fetal Evidence Based Guidelines).

Parvovirus. Routine screening is not recommended, but can be considered for high-risk groups (see Chapter 48 in Maternal-Fetal Evidence Based Guidelines).

Toxoplasmosis. Universal screening is not recommended. Education regarding prevention of disease should be addressed (Table 2.3) (see Chapter 49 in Maternal-Fetal Evidence Based Guidelines).

Follow-Up Visits
Follow-up visits should provide for the following:
• Follow-up physical exam, laboratory screening, and testing as indicated
• Ongoing assessment of risk factors and plan for intervention as indicated

<table>
<thead>
<tr>
<th>Table 2.3</th>
<th>Prevention of Food-Borne Illnesses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Food-borne illness to avoid</td>
<td>Preventive strategy</td>
</tr>
<tr>
<td>Listeriosis</td>
<td>Cook meat thoroughly including luncheon meats; avoid raw or smoked meats or fish, pates, unpasteurized cheese, and raw milk.</td>
</tr>
<tr>
<td>Toxoplasmosis</td>
<td>Cook meat and wash fruits and vegetables thoroughly; avoid cat litter; wear gloves when gardening outdoors.</td>
</tr>
<tr>
<td>Escherichia coli and Salmonella Methylmercury</td>
<td>Follow food-handling guidelines above.</td>
</tr>
</tbody>
</table>

Table 2.4 | Institute of Medicine Recommended Total Weight Gain in Pregnancy by Prepregnancy BMI (kg [lb.]) |
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>BMI</td>
<td>Singleton</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>--------------------------------------------------</td>
</tr>
<tr>
<td>&lt;18.5</td>
<td>12.5–18 (27–40)</td>
</tr>
<tr>
<td>18.5–24.9</td>
<td>11.5–16 (25–35)</td>
</tr>
<tr>
<td>25.0–29.9</td>
<td>7–11.5 (15–25)</td>
</tr>
<tr>
<td>≥30*</td>
<td>5–9 (11–20)</td>
</tr>
</tbody>
</table>

*See Table 2.7 for our recommendations.

• Education and health promotion directed to individual plan of care
• Opportunity for discussion and questions

Follow-Up Physical Exam
• Weight: Usually done at each visit, as optimal weight gain (Table 2.4) is associated with better outcomes. Excessive fast weight gain can be a sign of preeclampsia.
• Blood pressure: Should be performed and recorded at each visit.
• Fetal heart tones: Should be performed and recorded at each visit after the first trimester.
• Symphyseal-fundal height measurement: Can be performed at each visit from the 24th through 41st weeks. Fundal height measurement may help to detect fetal growth restriction (FGR) and macrosomia, but there is poor intra- and interuser reliability. There is probably some value in evaluating trends and although it will not impact on the underlying condition, it may affect decision making on fetal surveillance. There is insufficient evidence to show whether this measurement has any impact, beneficial or not, on pregnancy outcomes, with no effect in the only one trial [19] (see also Chapter 45 in Maternal-Fetal Evidence Based Guidelines).
• Cervical examination: Routine digital examination of the cervix is not recommended as a screening measure for prevention of PTB (see Chapter 17 in Maternal-Fetal Evidence Based Guidelines).
• Sweeping or “stripping” of membranes during cervical exam at ≥38 weeks reduces the rate of late-term delivery (see Chapter 21). Cervical examination may assist in the identification of abnormal presentation, and therefore the opportunity to offer appropriate intervention (i.e., version).
• Fetal movement: There is no evidence that formalized kick counts reduce the incidence of fetal death in the healthy singleton [20] (see Chapter 56 in Maternal-Fetal Evidence Based Guidelines). Nonetheless, women may be instructed to be aware of daily fetal movements from at or around 28 weeks.
• Leopold’s maneuvers: Perform at each visit from 34 weeks to estimate fetal weight and determine presentation. Ultrasound can be used to confirm findings, and interventions may be offered [21,22].
• Clinical pelvimetry: Measurement of the bony birth canal is of limited, unproven value in predicting dystocia during delivery (see Chapters 7 and 8).
• Routine evaluation for edema: Edema has traditionally been a part of the evaluation for preeclampsia, but by itself, it is neither specific nor sensitive.
Follow-Up Laboratory Screening

- 10 6/7–13 6/7 weeks: Serum aneuploidy screening, with nuchal translucency screening by ultrasound (see below), should be offered to every pregnant woman. Consider cell-free DNA aneuploidy testing (also called noninvasive prenatal testing, NIPT) in high-risk women (see Chapter 5) (Table 2.1).
- 14–21 weeks: The second part of serum aneuploidy screening (best at 16–18 weeks) should be offered to all pregnant women interested in prenatal diagnosis of aneuploidy (see Chapter 5). Counseling regarding the variety of screening options and the limitations of testing should be made available to all pregnant women.
- 24–28 weeks: Women with risk factors for GDM should be screened with either one-step or two-step tests, since intervention (diet, exercise, glucose monitoring, and, as necessary, medical therapy) prevents maternal and perinatal morbidities (see Chapter 5 in Maternal-Fetal Evidence Based Guidelines). Universal glucose challenge screening for GDM is the most sensitive approach, but the following women are at low risk and less likely to benefit from testing (must meet all of the following criteria): age <25 years; ethnic origin of low-risk (not Hispanic, African, Native American, South or East Asian, or Pacific Islander); BMI <25; no previous personal or family history of impaired glucose tolerance; and no previous history of adverse obstetric outcomes associated with GDM. Antibody screening and hemoglobin and hematocrit are also repeated. Repeat screening of rapid plasma reagin (RPR) (or venereal disease research laboratory [VDRL]) and HIV in the early third trimester and at delivery can be considered for high-risk populations (see Chapters 32 and 35 in Maternal-Fetal Evidence Based Guidelines).
- 35–37 weeks: Group B Streptococcus (GBS) is a significant cause of morbidity and mortality in neonates. Approximately 10%–30% of pregnant women are asymptptomatically colonized with GBS in the vagina or rectum. Vertical transmission of this organism from mother to fetus occurs most commonly after onset of labor or rupture of membranes. All women should be screened for GBS colonization by rectovaginal culture at 35–37 weeks of gestation. Colonized women should be treated with IV antibiotics (penicillin is first choice if not allergic) in labor or with rupture of membranes (see Chapter 37 in Maternal-Fetal Evidence Based Guidelines).

Ultrasoundography

Ultrasound has not been proven harmful to mother or fetus (see Chapter 4).

- First-trimester “fetal dating” ultrasoundography (before 14 weeks): First-trimester ultrasound is more accurate than LMP to determine gestational age. First-trimester ultrasound also allows earlier detection of multiple pregnancies, aneuploidy screening with nuchal translucency, and diagnosis of nonviable pregnancies.
- Second-trimester “fetal anatomy” ultrasound: Generally, women are offered an ultrasound at 18–22 weeks to screen for structural anomalies. Routine use of ultrasound reduces the incidence of postterm pregnancies and rates of induction of labor for postterm pregnancy, increases early detection of multiple pregnancies, increases earlier detection of major fetal anomalies when termination of pregnancy is possible, increases detection rates of fetal malformations, and decreases admission to special care nursery [23,24]. Given the benefits mentioned, all pregnant women should be offered a second-trimester ultrasound. No significant differences are detected for substantive clinical outcomes such as perinatal mortality, possibly because of insufficient data. Transvaginal ultrasound (TVU) cervical length (CL) screening of all singletons gestations, even those without a prior spontaneous PTB, can be offered (ACOG 2012, SMFM 2012), and is recommended by experts [25] (see also Chapter 17).
- Third-trimester “fetal growth” ultrasound: In low-risk or unselected populations, routine third-trimester ultrasound has not been associated with improvements in perinatal mortality [26]. Selective ultrasound in later pregnancy is of benefit in specific situations, such as calculation of interval growth for suspected FGR, assessment of amniotic fluid index for suspect oligohydramnios or hydramnios, and assessment of malpresentation (see Chapter 4). A large prospective cohort study showed that screening of nulliparous women with universal third-trimester fetal biometry roughly tripled the detection of SGA infants and that the combined analysis of fetal biometry and fetal growth velocity identified a subset of SGA fetuses that were at increased risk of neonatal morbidity [27].
- Routine umbilical artery or other Doppler ultrasound in low-risk or unselected patients has not been shown to be of benefit.

CONTENT OF PREGNATAL CARE

The content of prenatal care is extensive and reviewed in detail not only in this chapter but also in most other chapters in this book, as well as its companion, Maternal-Fetal Evidence Based Guidelines. In Chapter 1, see Table 1.2 for topics to be reviewed, Table 1.3 for screening, Table 1.4 for laboratory tests, Table 1.5 for vaccinations, Table 1.6 for interventions for all women, and Table 1.7 for interventions for women with risk factors. Prenatal care usually incorporates, among other things, the following:

- Prenatal education and reassurance (regarding drugs, environment, lifestyle, nutrition, supplements, vaccinations, preventive measures, preparation for labor and delivery, depression, breastfeeding, etc.)
- Provision of evidence-based screening tests at appropriate intervals (Table 2.1)
- Risk assessment
- Problem-oriented visits as needed
- Condition-specific care for high-risk patients

Content issues that should be included in prenatal care such as drugs and environment, lifestyle, nutrition, supplements, vaccinations, prenatal education, and others are described below.

Drugs and Environment

Substance Abuse

Screening for use and counseling for cessation of tobacco, alcohol, and recreational or illicit drug use is recommended (see Chapters 22 and 23 in Maternal-Fetal Evidence Based Guidelines). Maternal smoking as well as exposure to secondhand smoke is dangerous to both the woman and her fetus. Provider support and educational material tailored to
pregnancy are shown to increase smoking cessation by 70% and reduce LBW and PTB [28–30] as well as the number of women who continue to smoke in late pregnancy [30].

**Alcohol use** at any level in pregnancy cannot be supported although deleterious effects at low-moderate levels are difficult to quantify [31]. The evidence from the limited number of studies suggests that psychological and educational interventions may result in increased abstinence from alcohol, and a reduction in alcohol consumption among pregnant women. However, results were not consistent, and the paucity of studies, the number of total participants, the high risk of bias of some of the studies, and the complexity of interventions limit our ability to determine the type of intervention that would be most effective in increasing abstinence from, or reducing the consumption of, alcohol among pregnant women [32]. Counseling may be effective in reducing substance abuse in pregnancy, although women with addictions will need specialized interventions. **Screening and brief intervention (SBI)** for unhealthy alcohol use has demonstrated efficacy in some trials. There is some evidence regarding the acceptability and efficacy of computer-delivered SBI plus tailored mailings in women who screened positive for alcohol risk [33]. There is insufficient evidence to recommend the routine use of home visits for women with a drug or alcohol problem [34]. However, a cluster randomized controlled trial among urban South African mothers showed that a home-visiting intervention improved the emotional health of low-income mothers and that relative to standard care, intervention mothers were significantly less likely to report depressive symptoms and alcohol abuse [35].

**Over-the-Counter, Alternative/Complementary, and Prescription Medications**

Because of the possibility of adverse fetal effects, medication use, including alternative remedies, should be limited to circumstances where benefit outweighs risk. Beneficial medications should be continued in pregnancy when safe for both mother and fetus (see specific disease guidelines in *Maternal-Fetal Evidence Based Guidelines*).

**Environmental/Occupational Risks and Exposures**

In general, working is not associated with poor pregnancy outcome. Some workplace exposures, such as toxic chemicals, radiation (>5 rad), heavy repeated lifting, prolonged (>8 hours) standing, excessive (>80/week) work hours, and high fatigue score may be associated with pregnancy complications, but there is insufficient evidence on the effect of avoidance of these risks (see also Chapter 17). There is insufficient safety data for paint, solvents, hair dyes, fumes, anesthetic drugs, etc., with no absolute evidence of harm. Hot tubs, saunas should avoid temperatures >102°F to avoid risk of dehydration, especially in the first trimester.

**Domestic Violence**

Domestic violence against pregnant women is associated with an increased risk of PTB, LBW, second- and third-trimester bleeding, and fetal injury. Domestic violence may escalate during pregnancy. As such, providers need to be alert to signs and symptoms of abuse and provide opportunities for private disclosure. However, so far, there is insufficient evidence to assess the effectiveness of interventions for domestic violence on pregnancy outcome [36].

**Lifestyle**

**Work**

There is insufficient evidence to recommend exact work hours and when to take off from work before delivery (if at all). Work accommodations are often necessary and helpful to allow a pregnant woman to continue working and earning an income. Pregnant women should not be discriminated against by their employers just because they are pregnant. The website www.pregnantatwork.org provides online tools that health-care professionals can use to prepare notes drafted using language that increases the likelihood that a patient will receive the accommodations she needs to continue doing her job safely. Occupational lifting guidelines have been published [37].

**Exercise**

Regular exercise during low-risk pregnancies is beneficial as it increases overall maternal fitness and sense of well-being. Exercise is an effective tool in maternal weight gain control in pregnancy [38–40].

Aerobic exercise for 35–90 minutes 3–4 times per week can be safely performed by normal-weight women with singleton, uncomplicated gestations because this is not associated with an increased risk of PTB or with a reduction in mean gestational age at delivery [39], and was associated with a significantly higher incidence of vaginal delivery and a significantly lower incidence of cesarean delivery [39,40], with a significantly lower incidence of gestational diabetes mellitus and hypertensive disorders and therefore should be encouraged [39].

Structured physical exercise programs appear also to be safe for the neonate [41] and reduce the risk of having a large newborn without a change in the risk of having a small newborn [42]. Furthermore, there is some evidence that exercise may be effective in treating depression during pregnancy [43]. Diet or exercise, or both, during pregnancy can reduce the risks of: excessive gestational weight gain (GWG), cesarean section, maternal hypertension, macrosomia, and neonatal respiratory morbidity, particularly for high-risk women receiving combined diet and exercise interventions [39]. However, most of the studies included in the meta-analysis were carried out in developed countries and therefore it is not clear if these findings are widely applicable [44]. In another meta-analysis, exercise was associated with a lower (by 600 g) GWG [45]. Possible maternal benefits include improved cardiovascular function, limited pregnancy weight gain, decreased musculoskeletal discomfort, reduced incidence of muscle cramps and lower limb edema, mood stability, and attenuation of GDM and gestational hypertension. Fetal benefits include decreased fat mass, improved stress tolerance, and advanced neurobehavioral maturation [46]. For most pregnant women, at least 30 minutes of moderate exercise is recommended on most days all of the week. There is no target heart rate that is right for every pregnancy woman. Walking, swimming, and other sports with low chance of loss of balance are recommended (Table 2.5) [47]. Avoid contact sports and sports with high chance of loss of balance. Special considerations may be made for professional athletes at the patient and provider’s discretion. Avoid hypoglycemia and dehydration. It is important to advise women to be careful while stretching, as the hormone relaxin can leave joints vulnerable to overstretching and injury [47]. It is important for clinicians to keep emphasizing that exercise is medicine [48].
The following activities are safe to initiate or continue:
- Walking
- Swimming
- Stationary cycling
- Low-impact aerobics
- Yoga, modified
- Pilates, modified
- Running or jogging
- Racquet sports
- Strength training

The following activities should be avoided:
- Contact sports (e.g., ice hockey, boxing, soccer, and basketball)
- Activities with a high risk of falling (e.g., downhill snow skiing, water skiing, surfing, off-road cycling, gymnastics, and horseback riding)
- Scuba diving
- Sky diving
- Running or jogging
- Pilates, modified
- Yoga, modified
- Hot yoga or hot pilates

Yoga

In pregnancy is associated with lower pain and discomfort, as well as lower perceived stress and improved quality of life in physical domains in the three RCTs evaluating its effects [49].

Travel

Counseling should include the proper use of passenger restraint systems in automobiles with the lap belt below the abdomen, reduction of risk of venous thromboembolism during long-distance air travel by walking and exercise, and provision of care and prevention of illness during travel abroad.

Sex and Sexuality

Intercourse has not been associated with adverse outcomes in pregnancy. Some women have a progressive decrease in sexual desire during the pregnancy, most markedly in the third trimester. Couples are often concerned that intercourse may harm the pregnancy. This is associated with progressively decreasing frequency of sexual intercourse in pregnancy [50]. Most women desire more communication regarding sex in pregnancy by their care providers. Health-care provider counseling should be reassuring, in the absence of pregnancy complications. Semen is a source of prostaglandin, pyospermia is associated with preterm premature rupture of membranes (PPROM), and orgasms and nipple stimulation do increase contractions [51]. Therefore, sexual intercourse may be detrimental in women with cervical dilatation and/or shortening but this is not well studied. PTB and other complications of pregnancy do not seem increased in most studies of sex in pregnancy. Most studies report that sexual activity is associated with better pregnancy outcomes, probably because women who are sexually active are healthier to begin with compared with women with less sexual activity [52].

Nutrition

Energy (Calorie)/Protein Supplementation

A meta-analysis of 17 RCTs provided evidence that antenatal nutritional education with the goal of increasing energy and protein intake in pregnant women appears to be effective in reducing the risk of PTB and of LBW and effective in increasing the head circumference at birth and the birth weight among undernourished women [53]. Balanced energy and protein supplementation seems to improve fetal growth, and may reduce the risk of stillbirth and infants born SGA. However, high-protein supplementation does not seem to be beneficial and may be harmful to the fetus increasing the risk of SGA. Balanced-protein supplementation alone has no significant effects on perinatal outcomes [53].

Cholesterol-Lowering Diet

A cholesterol-lowering diet with omega-3 fatty acids and dietary counseling does not affect cord or neonatal lipids but is associated with a 90% reduction in PTB <37 weeks in one trial [54] (see also Chapter 17). More evidence is needed for a recommendation.

Low-Glycemic Index Diet

A low-glycemic index diet appears to be beneficial to both mother and child in reducing the incidence of abnormal glucose tolerance tests, large-for-gestational-age (LGA) infants, and ponderal indices. The numbers of studies and subjects are small, however, and therefore considered inconclusive [55]. Studies evaluating the effects of different types of dietary advice for women with gestational diabetes mellitus did not find any significant benefits for the diets investigated [56].

Antigen Avoidance Diet

Prescription of an antigen avoidance diet (e.g., avoiding chocolate or nuts) to a pregnant woman is unlikely to reduce her child's risk of atopic disease and such a diet may adversely affect maternal or fetal nutrition [57].

Probiotics

A probiotic capsule intervention among women with abnormal glucose tolerance had no impact on glycemic control [58].
Food Safety

Food safety and prevention of food-borne illness and infection are suggested in Tables 2.3 and 2.6.

BMI and Weight Gain

BMI is utilized in counseling a woman on optimal weight gain in pregnancy (Table 2.2) [59–71].

Suggested weight gain in pregnancy is shown in Table 2.4 [59]. Women who are underweight are at increased risk for LBW and PTB and have better outcomes with a higher total weight gain [67]. Excessive weight gain in women with normal BMI can be prevented with dietary and lifestyle counseling [62–69]. For example, a program of education on recommended GWG, application of personalized weight graph, formalized prescription of exercise, and regular monitoring of GWG at every antenatal visit is associated with a significant reduction in GWG [70]. Obesity is associated with cardiovascular disease, diabetes, hypertension, stroke, osteoarthritis, gallstones, endometrial, breast, and colon cancers, cardiomyopathy, fatty liver, obstructive sleep apnea, urinary tract infections, other complications, and most importantly, mortality. Prepregnancy obesity and excessive gestational weight gain are associated with increased risk of childhood obesity for the fetus. Obese pregnant women are specifically at increased risk for miscarriage, congenital malformations, GDM, hypertension, preeclampsia, stillbirth, cesarean birth, labor abnormalities, macrosomia, anesthesia complications, wound infection, and thromboembolism. These women have better maternal outcomes with lower (or no) total weight gain [60,61,67,68,71] (Table 2.7) (see also Chapter 3 in Maternal-Fetal Evidence Based Guidelines). Even if some studies have reported some small increased risk of SGA with weight loss in obese women, this is really NOT an increase. What happens is that obese women who gain weight have larger babies (incidence of SGA ≤5%), while those who lose weight have a normal incidence of SGA (i.e., ≤10%) [72]. Moreover, all other neonatal outcomes are the same or better with no weight gain or some moderate weight loss in obese pregnant women (Table 2.7) [61,72].

Caffeine

Moderate caffeine consumption (<200 mg/day) does not appear to be a major contributing factor in miscarriage or PTB.

Reducing the caffeine intake of regular coffee drinkers (3+ cups/day) during the second and third trimester by an average of 182 mg/day did not affect birth weight or length of gestation in one RCT [73]. A meta-analysis from two RCTs concluded that there is insufficient evidence to confirm or refute the effectiveness of caffeine avoidance on birth weight or other pregnancy outcomes; moreover, they found that reducing the caffeine intake of regular coffee drinkers (3+ cups/day) during the second and third trimester did not affect PTB or SGA rate [74].

Supplements

**Multivitamin**

There is insufficient evidence to suggest replacement of iron and folate supplementation with a multiple-micronutrient supplement. A reduction in the number of LBW and SGA babies and maternal anemia has been found with a multiple-micronutrient supplement compared with supplementation with two or less micronutrients or none or a placebo, but analyses revealed no added benefit of multiple-micronutrient supplements compared with iron and FA supplementation [75,76]. These results are limited by the small number of studies available. There is also insufficient evidence to identify which micronutrients are more effective, to assess adverse effects, and to say that excess multiple-micronutrient supplementation during pregnancy is harmful to the mother or the fetus [75,76]. Therefore, there is insufficient evidence to recommend routine multivitamin supplementation for all women, or even only for women who are underweight, have poor diets, smokers, substance abusers, vegetarians, multiple gestations, or others. Excess (>1) prenatal vitamin intake per day should be avoided. No prenatal multivitamin supplement has been shown to be superior to another. Use of multivitamin supplement not specific for pregnancy should be discouraged, as often excess doses can pose risks to the pregnancy. Each supplement, including each vitamin supplement, should be studied for safety and efficacy individually.

**Folic Acid**

Folic acid supplementation is recommended, with minimum 400 µg/day for all women (93% decrease in NTDs), and 5 mg/day for women with prior children with NTD (69% decrease in NTD) [77,78]. Supplementation should start at least 1 month before conception and continue until at least 28 days after conception (time of neural tube closure). Given the unpredictability of conception and that 50% of pregnancies are unplanned, all reproductive-age women should be on FA supplementation. Because in several countries the baseline serum folate level is only 5 ng/mL, and increases in this level are directly proportional with a decrease in the incidence of NTD, some experts have advocated 5 mg of FA per day as optimal supplementation [79]. No increase in ectopic pregnancy, miscarriage, or stillbirth has been associated with folate supplementation, but it might increase (nonsignificant trend) the incidence of multiple gestations by 40% [77–82]. However, a multicenter prospective cohort study showed that children whose mothers used FA supplement dosages higher than 5 mg/mL had a lower mean psychomotor scale score than children whose mothers used a recommended FA supplements dosages (i.e., 400 µg/day) [82]. Folic acid supplementation has been associated (one non-RCT study) with decrease in severe language delay at 3 years of age [81]. Fortifying basic foods such as grains with added folate is associated with an increase in supplementation of only 140–200 µg/day, and with only a

<table>
<thead>
<tr>
<th>Table 2.7 Weight Gain Suggestions for Overweight and Obese Women</th>
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</thead>
<tbody>
<tr>
<td><strong>Prepregnancy weight category</strong></td>
</tr>
<tr>
<td>--------------------------------</td>
</tr>
<tr>
<td>Overweight (BMI 25–29.9 kg/m²)</td>
</tr>
<tr>
<td>Class I obesity (BMI 30–34.9 kg/m²)</td>
</tr>
<tr>
<td>Class II obesity (BMI 35–39.9 kg/m²)</td>
</tr>
<tr>
<td>Class III obesity (BMI &gt;40 kg/m²)</td>
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Abbreviation: IOM, Institute of Medicine.
20%–50% decrease in incidence of NTD, with the potential for large-scale prevention [80]. Women taking antiseizure medications, other drugs that might interfere with FA metabolism, those with homozygous methylenetetrahydrofolate reductase (MTHFR) enzyme mutations, or those who are obese may need higher doses of folate supplementation. Women with first-trimester diabetes mellitus or exposure to valproic acid or high temperatures might not experience the decrease in NTD risk with folate supplementation due to these risks (see also Chapter 1).

Vitamin A
In pregnancy, some extra vitamin A is required for growth and tissue maintenance in the fetus, for providing fetal reserves, and for maternal metabolism. However, vitamin A in its synthetic form as well as in large doses as retinol (preformed vitamin A found in cod liver oil and chicken or beef liver) is teratogenic. It is recommended that pregnant women ingest vitamin A found in cod liver oil and chicken or beef liver) is teratogenic form as well as in large doses as retinol (preformed vitamin A in its synthetic form). Vitamin A is beneficial during pregnancy for either low- or high-risk women. There may be an associated increased risk of PTB with vitamin C supplementation (RR 1.38, 95% CI 1.04–1.82, 3 trials, 583 women) [87]. No other difference in outcome is noted between vitamin C supplementation and no treatment or placebo. There are very limited trials available to assess whether vitamin C supplementation may be useful for all pregnant women. Usually the women involved in the trials were either at high risk of preeclampsia or PTB or the women had established severe early-onset preeclampsia (see also Chapter 1 in Maternal-Fetal Evidence Based Guidelines). No difference is seen between women supplemented with vitamin C alone and those supplemented with vitamin C in combination with other supplements compared with placebo for the risk of stillbirth, neonatal death, LBW, or intrauterine growth restriction (IUGR) [87].

Vitamin D
There is insufficient evidence to evaluate vitamin D supplementation during pregnancy [88–92]. Vitamin D 1000 IU/day in the third trimester is associated with no consistent effect on incidence of LBW [88]. Neonatal hypocalcemia is less common with vitamin D supplementation compared with placebo [88]. Vitamin D supplementation during pregnancy is associated with increase circulating 25(OH)D levels, birth weight and birth length but with no effects on maternal-fetal outcomes [89]. There are limited data to assess any benefit of vitamin D supplements for complete vegetarians and women with extremely limited exposure to sunlight. Vitamin D supplementation of vitamin D deficient pregnant women prevents neonatal vitamin D deficiency [90]. Vitamin D plus calcium have no effect on duration of pregnancy, type of delivery, and infant anthropometric indicators [91]. However, low maternal vitamin D levels in pregnancy ≤50 nmol/L may be associated with an increased risk of preeclampsia, gestational diabetes, PTB, and SGA [92].

Vitamin B6 (Pyridoxine)
There is insufficient evidence to evaluate pyridoxine supplementation during pregnancy [85]. There are few trials, reporting few clinical outcomes and mostly with unclear trial methodology and inadequate follow-up. There is not enough evidence to detect clinical benefits of vitamin B6 supplementation in pregnancy and/or labor other than one trial suggesting protection against dental decay [86]. For the aim of decreasing dental decay or missing/filled teeth, pyridoxine supplementation 20 mg/day (lozenges or capsules) is associated with decreased incidence of these outcomes in pregnant women [86]. Pyridoxine has been used in the management of nausea and vomiting in pregnancy. It is now considered Category A in combination with doxylamine as “Diclegis” which is the only U.S. Food and Drug Administration (FDA) approved treatment for nausea and vomiting of pregnancy. Studies done for FDA approval of the drug showed no adverse outcomes and demonstrated safety and good tolerance by women when used in the recommended dose of up to 4 pills (10 mg/10 mg) per day (see Chapter 9 in Maternal-Fetal Evidence Based Guidelines).

Vitamin C
The data are insufficient to assess if vitamin C supplementation either alone or in combination with other supplements is beneficial during pregnancy for either low- or high-risk women. There may be an associated increased risk of PTB with vitamin C supplementation (RR 1.38, 95% CI 1.04–1.82, 3 trials, 583 women) [87]. No other difference in outcome is noted between vitamin C supplementation and no treatment or placebo. There are very limited trials available to assess whether vitamin C supplementation may be useful for all pregnant women. Usually the women involved in the trials were either at high risk of preeclampsia or PTB or the women had established severe early-onset preeclampsia (see also Chapter 1 in Maternal-Fetal Evidence Based Guidelines). No difference is seen between women supplemented with vitamin C alone and those supplemented with vitamin C in combination with other supplements compared with placebo for the risk of stillbirth, neonatal death, LBW, or intrauterine growth restriction (IUGR) [87].
of preeclampsia and all other outcomes are similar. In the analysis of one high-quality trial, no differences between magnesium and placebo groups are seen. Poor-quality trials are likely to have resulted in a bias favoring magnesium supplementation.

Calcium
Calcium supplementation is associated with a reduction of the incidence of preeclampsia in pregnancy in all women, particularly for women at high risk of hypertension and in women with low dietary calcium intake (e.g., <600 mg/day) [95]. The minimum dose in the Cochrane review was 1 g/day. Further research is needed to determine whether dietary sources of calcium confer the same benefit and at what amount. There is insufficient evidence to determine optimum dosage and the effect on other important maternal and fetal outcomes. There is no overall reduction in PTB, although there is a reduction in PTB among women at high risk of developing hypertension. Benefits are considered to outweigh an anomalous increase in the risk of HELLP syndrome, which was small in absolute numbers. There is no evidence of any effect of calcium supplementation on stillbirth or death before discharge from hospital. In women at high risk of hypertension, calcium supplementation is associated with fewer babies with birth weight <2500 g. In one study, childhood systolic blood pressure >95th percentile was reduced [95] (see also Chapter 1 in Maternal-Fetal Evidence Based Guidelines).

Iron
There is no evidence to advice against a policy of routine iron and folate supplementation in pregnancy. Iron supplementation is associated with prevention of low hemoglobin at birth or at 6 weeks postpartum [96]. Iron supplementation, however, has no detectable effect on any substantive measures of either maternal or fetal outcome. One trial, with the largest number of participants of selective versus routine supplementation, shows an increased likelihood of cesarean section and postpartum blood transfusion, but a lower perinatal mortality rate (up to 7 days after birth). There are few data derived from communities where iron deficiency is common and anemia is a serious health problem. There is limited evidence for daily versus intermittent supplementation. High-dose supplementation (80 mg daily) has no clinical advantage over low-dose supplementation (20 mg daily) and is associated with more gastrointestinal (GI) side effects. One RCT suggests adverse effects of hemocoagulation from iron supplementation in nonanemic women. For iron supplementation for women with anemia, see chapter 1 and 4 and 25 months of age [98]. There is little data, however, on the safety of routine iodine supplementation in populations with normal or low normal iodine levels. Some data suggest an increased risk of fetal and maternal hypothyroidism from iodine supplementation. The upper levels of safety have not been established [98,99].

Omega-3
Pregnancy is a time of increased risk for omega-3 deficiency as omega-3 is used for the developing fetus. Thirty-four RCTs have been performed to assess whether omega-3 supplementation during pregnancy affects maternal-fetal outcomes. Pooled results from the 34 studies [100] show lack of evidence to support the routine use of omega-3 supplementation during pregnancy, as omega-3 supplementation did not affect PTB, preeclampsia, IUGR, gestational diabetes, SGA, post-partum depression, children development or other maternal or fetal outcomes. Meta-analyses also found that omega-3 supplementation during pregnancy did not prevent PTB in low-risk women [101], or in women with prior PTB [102], and did not prevent recurrent IUGR [103].

Vaccinations
Immunity to rubella, varicella, hepatitis B, influenza, tetanus, and pertussis should be assessed at the first prenatal visit. Ideally, needed vaccinations would be provided preconception. There is no vaccine that is more dangerous to a pregnant woman or her fetus than the disease it is designed to prevent. Recombinant, inactivated, and subunit vaccines, as well as toxoids, and immunoglobulins pose no threat to a developing fetus. Inactivated influenza vaccine should be given (by injection, as killed virus) to all pregnant women during the influenza season. The live attenuated form of the vaccine (intranasal spray) should not be given during pregnancy. Hepatitis B vaccine can be safely given in pregnancy. Tdap or “whooping cough” vaccine is recommended for all pregnant women after 28 weeks (see Table 1.5 in Chapter 1, and Chapter 38 in Maternal-Fetal Evidence Based Guidelines).
Abdominal Decompression
Abdominal decompression consists of a rigid dome placed about the abdomen and covered with an airtight suit, with the space around the abdomen decompressed to ~50 to ~100 mmHg for 15–30 seconds out of each minute for 30 minutes once to thrice daily, or with uterine contractions during labor. This is thought to “pump” blood through the intervillous space. There is no evidence to support the use of abdominal decompression in normal pregnancies. There is no difference between the abdominal decompression groups and the control groups for LBW, admission for preeclampsia, low Apgar score, perinatal mortality, and childhood development [105].

Prevention of Complications
Please see specific diseases in each chapter of this book, and its companion, Maternal-Fetal Evidence Based Guidelines. Here are reported only some general, nonspecific interventions.

Antibiotic prophylaxis of pregnant women with no specific risk factor or infection is associated with similar incidence of PPROM, PTB, and postpartum endometritis [106,107] (see also Chapter 17).

Programs offering additional social support (caring family members, friends, and health professionals) for at-risk (e.g., for PTB and LBW) pregnant women are not associated with improvements in any perinatal outcomes, but there is a reduction in the likelihood of antenatal hospital admission (RR 0.79, 95% CI 0.68–0.92) and cesarean birth (RR 0.87, 95% CI 0.78–0.97) [108].

For issues such as mild hypertension or preeclampsia, small studies suggest that there are no major differences in clinical outcomes for mothers or babies between antenatal day units and hospital admission, but women may prefer day care [109] (see also Chapter 1 in Maternal-Fetal Evidence Based Guidelines).

Prenatal Education
There is insufficient evidence to assess the effectiveness of formal prenatal education programs. Prenatal education directed at specific objectives (e.g., promoting breastfeeding and avoiding planned induction of labor) has been demonstrated to be effective [110–113]. Individualized prenatal education directed toward avoidance of a cesarean delivery does not increase the rate of vaginal birth after cesarean section. As a part of prenatal care, women should be provided with information and instruction regarding their health, including risk avoidance, breastfeeding, what to expect during labor and birth (see Section “Preparation for Labor and Delivery”), how to obtain care when labor begins, and the value of a support person during the labor process (see Chapters 7 and 8).

Community Interventions
There is encouraging evidence of the value of integrating maternal and newborn care in community settings through a range of interventions that can be packaged effectively for delivery through a range of community health workers and health promotion groups. Such evidence-based available interventions as immunization to mothers, clean and skilled care at delivery, newborn resuscitation, exclusive breastfeeding, clean umbilical cord care, and management of infections in newborns require facility-based and outreach services. Implementation of community-based interventional care packages is associated with a trend for reduction in maternal mortality (RR 0.77, 95% CI 0.59–1.02) and with significant reductions in maternal morbidity (RR 0.75, 95% CI 0.61–0.92), neonatal mortality (RR 0.76; 95% CI 0.68–0.84), stillbirths (RR 0.84, 95% CI 0.74–0.97), and perinatal mortality (RR 0.80; 95% CI 0.71–0.91). It also increases the referrals to health facility for pregnancy-related complication by 40% and improves the rates of early breastfeeding by 94% [113].

Preparation for Labor and Delivery
Perineal massage with sweet almond oil for 5–10 minutes daily from 34 weeks until delivery is associated with a significantly higher chance of intact perineum compared with no massage in nulliparous, but probably not multiparous women [114–116]. The type of the oil used during the second stage of labor for prevention of perineal tears has no effect on the integrity of the perineum; accordingly it seems that there is no perfect oil [117]. For perineal massage in labor, see Chapter 8.

Women should be provided with written information and instruction regarding what to expect during labor and delivery, how to obtain care when labor begins, and the value of a support person during the labor process (see Chapters 7 and 8).

Labor and delivery classes should be encouraged. Compared with no such training, 9 hours of antenatal classes with training to prepare for labor and delivery are associated with arriving to L and D ward more often in active labor (RR 1.45, 95% CI 1.26–1.65) and using less epidural analgesia (RR 0.84, 95% CI 0.73–0.97) [118].

Compared with standard antenatal education, antenatal education focusing on natural childbirth preparation with training in breathing and relaxation techniques is not associated with any effects on maternal or perinatal outcomes, including similar incidences of epidural analgesia, childbirth, or parental stress, in nulliparous women and their partners [119].

Compared with conventional therapy, intensive counseling therapy for fear of childbirth does not affect the incidence of cesarean but is associated with reduced pregnancy-and birth-related anxiety and concerns, and shorter labors in one RCT [120].

In a small RCT, a specific antenatal education program is associated with a reduction in the mean number of visits to the labor suite before the onset of labor [4]. It is unclear whether this results in fewer women being sent home because they are not in labor [120] (see also Chapter 7). One trial study, comparing the use of an educational technique based on patient participation with routine instructions to prepare patients to recognize the onset of active labor, showed that, without any increase in time, nurses can prepare patients to make judgments about the need for hospitalization [121].

Depression in Pregnancy and the Postpartum Period
Between 14% and 23% of pregnant women will experience a depressive mood disorder while pregnant [122–129]. Maternal anxiety, life stress, history of depression, lack of social support, unintended pregnancy, public insurance, domestic violence, lower income, lower education, smoking, single status, and poor relationship quality were associated with a greater likelihood of antepartum depressive symptoms in bivariate analyses. Life stress, lack of social support, and domestic violence continued to demonstrate a significant association in multivariate analyses [123,124]. Identification of risk factors and
screening for depression will facilitate referral for treatment (see Chapter 21 in Maternal-Fetal Evidence Based Guidelines).

Between 5% and 7% of women will experience postpartum depression. Risk factors include antenatal depressive symptoms, a history of major depressive disorder, or previous postpartum major depression [123]. If left untreated, postpartum major depression can lead to poor mother-infant bonding, delays in infant growth and development, and an increased risk of anxiety or depressive symptoms in the infant later in life. Identifying mothers at-risk assists the prevention of postpartum depression compared with intervening on the general population. The provision of intensive postpartum support provided by public health nurses or midwives is associated with 32% less postpartum depression. Interventions with only a postnatal component appeared to be more beneficial than interventions that also incorporated an antenatal component. Individual-based interventions may be more effective than those that are group based. Women who received multiple-contact intervention are just as likely to experience postpartum depression as those who received a single-contact intervention [125]. There is insufficient evidence to assess the effectiveness of antidepressants given immediately postpartum in preventing postnatal depression in all women or just in high-risk women [126]. Norethisterone enantate, a synthetic progestogen, 200 mg intramuscularly (IM) administered once within 48 hours of delivery to unselected women is associated with a significantly higher risk of developing postpartum depression at 6 weeks [127]. A pilot randomized controlled trial showed that prenatal yoga may be of benefit in prevention of postpartum depression in low-risk women [128]. Moreover, prenatal yoga was also found to be a feasible and acceptable intervention and was associated with reductions in symptoms in women with symptoms of anxiety and depression [129] (see Chapter 21 in Maternal-Fetal Evidence Based Guidelines).

Breastfeeding
Breastfeeding is the best feeding method for most infants and should be strongly encouraged (see Chapter 30). Counseling and education during pregnancy have been shown to facilitate breastfeeding success [130]. Breastfeeding education and/or support increased exclusive breastfeeding rates and decreased no breastfeeding rates at birth and at 1–5 months. Combined individual and group counseling appeared to be superior to either individual counseling alone or group counseling alone. Attitudes of the health-care provider are highly associated with breastfeeding success. Antigen avoidance diet during lactation by high-risk women may reduce the child’s risk of developing atopic eczema, and may reduce atopic eczema in children already with atopic eczema during the first 12–18 months, although more trials are needed.

INTERVENTIONS FOR COMMON PREGNANCY COMPLAINTS

Itching
The differential diagnosis of itching in late pregnancy (>32 weeks) is presented in Chapter 10 in Maternal-Fetal Evidence Based Guidelines. If the itching is not due to liver disease, and if there is no rash, aspirin (600 mg qid) has been reported to decrease itching [124], but because of potential detrimental fetal effects (closure of ductus arteriosus and oligohydramnios) should not be used. If there are both itching and a rash, chlorpheniramine 4 mg tid decreased itching in a small trial [131]. However, aspirin 600 mg four times a day appears to be more effective than chlorpheniramine 5 mg three times a day for relief of itching when no rash is present in a small crossover trial [132].

Stretch Marks
Some stretch marks (striae gravidarum) develop in about 50% of women by the end of pregnancy. There is no high-quality evidence to support the use of any topical preparation in the prevention of stretch marks during pregnancy [133,134]. There is also no proven treatment for stretch marks once they have developed [134]. Olive oil is not effective in preventing the occurrence of striae gravidarum or affecting its severity [134]. There is no available product that has been definitively shown to prevent the formation of SG. Massage with either Trofolastin cream or Verum ointment is associated in small RCTs with a decrease in the development of SG [134]. A small randomized trial showed that a specific anti-stretch mark cream (emollient and moisturizer containing hydroxyprolisilane C, rosehip oil, Centella asiatica, triterpenes, and vitamin E) had a small effect in reducing severity (but not the incidence) of striae during pregnancy [135] (see Chapter 43 in Maternal-Fetal Evidence Based Guidelines).

Leg Cramps
Leg cramps are reported to occur in a reported 34% of pregnant women in the midtrimester [136,137]. Magnesium lactate or citrate chewable tablets 5 mmol in the morning and 10 mmol in the evening for 3 weeks are associated with one-third of women not having persistent leg cramps compared with 94% of placebo controls having persistent cramps. Multivitamin with mineral supplement might decrease leg cramps, but it is unclear which one of the 12 ingredients (or combination) is beneficial. Sodium chloride is associated with a slight reduction although consideration must be given to potential effect on blood pressure. Calcium supplements do not decrease leg cramps compared with placebo. However, it is unclear whether any of the interventions studied (i.e., oral magnesium, oral calcium, oral vitamin B or oral vitamin C) provide an effective treatment for leg cramps due to poor study design and trials being too small to address the question satisfactorily. Calf stretching prior to bedtime does not decrease nocturnal leg cramps in nonpregnant patients [137].

Back and Pelvic Pain
Back pain is common in pregnancy, given weight gain and its uneven distribution as well as the softening effects of pregnancy hormones on the musculature.

There is evidence that exercise (any exercise on land or in water) may reduce pregnancy-related low-back pain, improve functional disability and reduce sick leave. Water gymnastics for 1 hour weekly starting at <19 weeks reduces back pain in pregnancy and allows more women to continue to work, with no adverse effects [138].

Pregnancy-specific exercises, physiotherapy, and acupuncture starting <32 weeks for 10 sessions appear to reduce back and pelvic pain; individual acupuncture sessions are more beneficial than group physiotherapy sessions. Education, other exercises, massage, heat therapy, support belts, analgesic therapy, etc. have not been studied in a trial in pregnancy for back pain relief.
Constipation
Constipation is common in pregnancy, probably because of decreased bowel peristalsis (possibly related to increased progesterone). It is reported by nearly 70% of women in the mid trimester. In nonpregnant adults, exercise, increase in water intake, dietary counseling, and certain foods (e.g., prunes) have been shown to relieve constipation. If these self-help measures are inadequate, the pregnant woman should then try daily bran or wheat fiber supplements. There is insufficient evidence to comprehensively assess the effectiveness and safety of interventions (pharmacological and nonpharmacological) for treating constipation in pregnancy, due to limited data (few studies with small sample size and no meta-analyses). Compared with bulk-forming laxatives, stimulant laxatives (e.g., Senna 14 mg, or dioctyl sodium succinate 120 mg and dihydroxyanthroquinone 100 mg—Normax) appear to be more effective in improvement of constipation (moderate quality evidence), but are accompanied by an increase in diarrhea and abdominal discomfort. Docusate sodium is a similar stimulant laxative, and it is widely available. Additionally, dietary fiber supplements (e.g., 10 mg/day of either corn-based biscuits—“Fibermed”—or 23 g wheat bran) increase the frequency of defecation and are associated with softer stools [139]. These findings in pregnant women are consistent with nonpregnant evidence.

Varicosities and Leg Edema
A small RCT (n = 69) shows that rutoside capsules improve leg edema symptoms; however, there are insufficient data to confirm rutoside safety in pregnancy. Another small RCT (n = 43) demonstrates a reduction in leg edema with reflexology. Compression stockings are not effective compared with simple resting, but studies do not compare compression stockings to no compression stockings. Leg elevation, compression hosiery, and swimming have not been studied for leg edema/varicosities relief in pregnancy [140].

Hemorrhoids
Hemorrhoids are common during pregnancy with 13% of women complaining of them in the mid trimester. Oral hydroxyethylrutosides decrease symptoms compared with placebo group in women with hemorrhoids and reduce the signs identified by the health-care provider [141]. Rutosides are associated with mild side effects such as GI discomfort, and their safety data in pregnancy are still insufficient. Constipation is a predisposing factor for hemorrhoids and should be treated. Sitz baths, ice, or ointments have been insufficiently studied for treatment of hemorrhoids in pregnancy. A small RCT showed that Hai’s Perianal Support toilet seat device reduced the symptoms of hemorrhoids in pregnancy and improved the well-being of pregnant women [142].

Heartburn
Heartburn is common during pregnancy with 53% of women complaining of it in the mid trimester. There is no large-scale RCT to assess heartburn relief in pregnancy [143].

A consensus document has recommended that lifestyle and dietary modifications should remain the first-line treatment for heartburn in pregnancy. The measures include reducing and avoiding intake of reflux-inducing foods (e.g., greasy and spicy foods, tomatoes, highly acidic citrus products, and carbonated drinks) and substances such as caffeine. Nonsteroidal anti-inflammatory drugs (NSAIDs) should also be avoided. Other lifestyle changes to reduce the risk of reflux, such as avoiding lying down within 3 hours after eating, are advised. However, if heartburn is severe enough to warrant this action, medication should begin after consultation with a health-care professional. Antacids, H2 blockers, and proton pump inhibitors all have acceptable safety profiles for the pregnant woman [143–145].

REFERENCES

A prospective randomized controlled trial. BJOG. 1990;97: 675–680. [RCT, n = 1,639] [I]


Khalifeh A, Berghella V. Ten reasons why universal cervical length screening should be recommended. AJOG. 2016 (in press). [III]


Physiologic changes

Jason Baxter and Colleen Horan

KEY POINTS
• The normal physiologic changes of pregnancy are several and listed in part in Table 3.1.
• Normal laboratory values for pregnant women are presented in Table 3.2.
• Failure to understand these physiologic changes of pregnancy may result in both undue alarm and costly evaluation of normal symptoms of pregnancy or in the neglect of pathologic conditions due to which the presentation is dismissed as another "discomfort of pregnancy."
• The physician should carefully address the pregnant patient keeping in mind the question "how is this presentation affected by the physiology of pregnancy?"

BACKGROUND
Over the course of human pregnancy, the significance of physiologic changes that occur is such that it often becomes no longer appropriate for the physician to evaluate her according to standards that have been set through the observation and study of men and nonpregnant women. Some of these physiologic changes are advantageous to the growth and survival of the fetus. Others enhance the ability of the maternal system to compensate for demands of pregnancy, prepare for stress of delivery, and recover from delivery.

Understanding physiologic changes in pregnancy is important in evaluating common symptoms associated with pregnancy, interpreting laboratory values in the parturient, and understanding pathologic conditions to which pregnant women are susceptible. Failure to understand the normal physiologic changes of pregnancy may result in both undue alarm and costly evaluation of normal symptoms of pregnancy or in the neglect of pathologic conditions due to which the presentation is dismissed as another discomfort of pregnancy. The patient will most likely be better served by the physician who carefully addresses her symptoms while keeping in mind the questions "how is this presentation affected by the physiology of pregnancy?" and "what pathologic conditions may be represented by this scenario?" than by the physician who uncritically memorizes laboratory values and makes a diagnosis without considering the interplay between pregnancy and underlying pathophysiology.

Two summary tables are provided. Table 3.1 summarizes the commonly accepted pregnancy-related changes in various physiologic parameters and their importance in the evaluation and management during pregnancy [1]. Table 3.2 provides a summary of laboratory values in each trimester of pregnancy versus in the nonpregnant state based on a recent systematic review [2]. The reader will note that some values show significant overlap between pregnant and nonpregnant states. Some show little change between pregnant and nonpregnant states. Others show trends that clearly increase or decrease with pregnancy. In most cases, it is not possible to consider the effect of pregnancy on laboratory values in a simplistic or formulaic way. Rather, it is the understanding of the underlying physiology that these laboratory values reflect that is the most important in their evaluation during pregnancy.

CARDIOVASCULAR/HEMODYNAMIC
An understanding of pregnancy-related hemodynamic changes is crucial in the management of both benign and life-threatening complications of pregnancy ranging from near-syncopal episodes experienced by many pregnant women to hypotension from obstetric hemorrhage and gestational hypertension, both of which are leading causes of maternal intensive care unit admissions and mortality [3]. Even the ultimate clinical emergency of cardiac arrest is complicated by hemodynamic changes which are unique to pregnancy.

Hemodynamic changes in pregnancy that have been well established include an increase in cardiac output and a decrease in both systemic and pulmonary vascular resistance. There is an overall increase in the heart rate and a decrease in the blood pressure. Blood volume, plasma volume, and erythrocyte volume increase, with a greater relative increase in the plasma volume resulting in a dilution lowering of hematocrit and other blood indices. There is also a redistribution of cardiac output with an increase in flow to the uterus, kidneys, skin, and breasts [1]. The increase in stroke volume and cardiac output creates a more audible physiologic flow murmur and splitting of the S2 sound during pregnancy, which may be striking upon physical examination.

One longitudinal study followed maternal hemodynamics as measured by thoracic electrical bioimpedance monitoring in 50 healthy pregnant women [4]. The results showed an increase in the mean heart rate from 87 ± 2 beats per minute (bpm) at 10–18 weeks to 92 ± 1 bpm at 34–42 weeks. Mean arterial pressure (MAP) decreased significantly after 14 weeks and increased after 29 weeks. Systemic vascular resistance (SVR) increased during the last trimester. This study also found a significantly higher mean cardiac output in nulliparous women compared with multiparous women. Mean cardiac output and stroke volume, which show an overall increase during pregnancy, were found to decrease in the third trimester in this study [4]. However, the change in cardiac output during the third trimester showed significant individual variation and has not been consistent in other longitudinal studies, demonstrating the need for further research for a more conclusive understanding of how this parameter changes across gestation [5].

Another longitudinal study performed serial echocardiography studies on 35 healthy pregnant women from the early second trimester to 6–12 weeks postpartum [6]. This study showed a significant increase in the cardiac output that peaked in the early third trimester and was maintained until term.
### Table 3.1 Summary of Physiologic Adaptations during Pregnancy

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Expected change</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Cardiovascular system</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Heart size</td>
<td>Increases 12%</td>
<td>Increased diastolic filling and muscle hypertrophy</td>
</tr>
<tr>
<td>Murmurs</td>
<td>Physiologic systolic</td>
<td>Ejection murmurs attributable to increased stroke volume usually occur in early or midsystole and are best heard along the left sternal edge</td>
</tr>
<tr>
<td>EKG</td>
<td>Positional heart shifts result in changes that resemble ischemia</td>
<td>Heart is pushed upward and forward, deviating electrical axis to the left by 15°–20°, causing flattened or inverted T in lead III</td>
</tr>
<tr>
<td>Cardiac output</td>
<td>Increases 30%–50%</td>
<td>Greatest increase occurs immediately after delivery with redistribution of blood flow from uterus. Cardiac output then decreases SVT not infrequent</td>
</tr>
<tr>
<td>Rhythm</td>
<td>Increases atrial and ventricular extrasystole</td>
<td>Significant change in second half of pregnancy</td>
</tr>
<tr>
<td>Mean arterial pressure</td>
<td>Decreased soon after beginning of pregnancy and midpregnancy (100–110/60–70 mean levels); then returns to prepregnant values by third trimester and term</td>
<td>Supine hypotension—decreased venous return due to compression from gravid uterus; increased blood flow through alternative pathways such as paravertebral azygous veins</td>
</tr>
<tr>
<td>Pulse pressure</td>
<td>Increases</td>
<td></td>
</tr>
<tr>
<td>Venous pressure</td>
<td>Increases in femoral system; unchanged in arms</td>
<td></td>
</tr>
<tr>
<td>Systemic vascular resistance</td>
<td>Decreases</td>
<td></td>
</tr>
<tr>
<td>Pulmonary BP</td>
<td>Unchanged</td>
<td></td>
</tr>
<tr>
<td>Blood flow to uterus</td>
<td>Increases by 500 mL/min</td>
<td>No autoregulation</td>
</tr>
<tr>
<td>Blood flow to kidneys</td>
<td>Increases by 400 mL/min</td>
<td></td>
</tr>
<tr>
<td>Blood flow to skin</td>
<td>Increases by 300–400 mL/min</td>
<td></td>
</tr>
<tr>
<td>Blood flow to breasts</td>
<td>Increases by 200 mL/min</td>
<td></td>
</tr>
<tr>
<td><strong>Respiratory system</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anatomy</td>
<td>Increases subcostal angle from 68° to 103° → 3 cm increase in transthoracic diameter</td>
<td>Increased upper respiratory capillary engorgement can cause increased congestion, epistaxis, and intubation trauma</td>
</tr>
<tr>
<td>Tidal volume</td>
<td>Increases by 300 mL or 40%</td>
<td>Cephalad displacement of diaphragm</td>
</tr>
<tr>
<td>Expiratory reserve volume</td>
<td>Decreases by 200 mL</td>
<td></td>
</tr>
<tr>
<td>Inspiratory capacity</td>
<td>Increases by 300 mL</td>
<td></td>
</tr>
<tr>
<td>Functional residual volume</td>
<td>Decreases by 500 mL</td>
<td></td>
</tr>
<tr>
<td>Minute volume</td>
<td>Increases by 40% or 3 L/min beginning in first trimester</td>
<td>Increased rate of induction of, and emergence from, inhaled anesthetics</td>
</tr>
<tr>
<td>Maximum breathing capacity</td>
<td>Unchanged</td>
<td></td>
</tr>
<tr>
<td>Forced expiratory volume</td>
<td>Unchanged</td>
<td></td>
</tr>
<tr>
<td>Peak expiratory flow rate</td>
<td>Unchanged</td>
<td></td>
</tr>
<tr>
<td>Pulmonary diffusing capacity</td>
<td>Decreases by 4 mL/min/mmHg</td>
<td></td>
</tr>
<tr>
<td>Oxygen requirement</td>
<td>Increases by 30–40 mL/min</td>
<td></td>
</tr>
<tr>
<td>Carbon dioxide output</td>
<td>Increases; expressed as respiratory quotient</td>
<td></td>
</tr>
<tr>
<td>Carbon dioxide pressure</td>
<td>Decreases from 35–40 mmHg to 28–30 mmHg</td>
<td></td>
</tr>
<tr>
<td>Oxygen pressure</td>
<td>Increases</td>
<td></td>
</tr>
<tr>
<td>pH</td>
<td>Mild increase (7.40–7.44 is normal)</td>
<td>Compensated respiratory alkalosis</td>
</tr>
<tr>
<td>PCO₂</td>
<td>Decrease (30–31 mmHg is normal)</td>
<td></td>
</tr>
<tr>
<td>PO₂</td>
<td>Mild increase (100–104 mmHg is normal)</td>
<td></td>
</tr>
<tr>
<td><strong>Renal system and homeostasis</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>GFR</td>
<td>Increases from 97 mL/min to 128 mL/min by 10 weeks</td>
<td>Increased renal clearance of certain drugs and vitamins</td>
</tr>
<tr>
<td>Glucose excretion</td>
<td>Increased</td>
<td>Random glycosuria</td>
</tr>
<tr>
<td>Protein excretion</td>
<td>Increased (up to 300 mg/24 hours is normal)</td>
<td>Diminished vascular response causes less pressor effect from angiotensin</td>
</tr>
<tr>
<td>Renin, angiotensins I and II</td>
<td>Increased</td>
<td>Increased risk of pyelonephritis; screen for asymptomatic bacteriuria</td>
</tr>
<tr>
<td>Anatomic changes</td>
<td>Dilation of renal calyces and ureters to pelvic brim; “physiologic” hydronephrosis, right &gt; left</td>
<td></td>
</tr>
<tr>
<td>Potassium</td>
<td>Increased retention</td>
<td></td>
</tr>
<tr>
<td>Sodium</td>
<td>Increased retention</td>
<td></td>
</tr>
<tr>
<td>Urine output</td>
<td>No significant change</td>
<td></td>
</tr>
</tbody>
</table>

(Continued)
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Expected change</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vitamin excretion</td>
<td>Increased loss of folate, B12, and ascorbic acid</td>
<td></td>
</tr>
<tr>
<td>Osmolality</td>
<td>Decreases 10 mOsm/kg first trimester, then stable</td>
<td>Increased intestinal absorption of calcium and increased bone turnover</td>
</tr>
<tr>
<td>Sodium</td>
<td>Decreases 3 mEq/L first trimester, then stable</td>
<td></td>
</tr>
<tr>
<td>Potassium</td>
<td>Decreases 0.5 mEq/L</td>
<td></td>
</tr>
<tr>
<td>Calcium</td>
<td>Decreased total and ionized</td>
<td></td>
</tr>
<tr>
<td>Magnesium</td>
<td>Decreases 10%–20% in first half of pregnancy</td>
<td></td>
</tr>
<tr>
<td>Chloride</td>
<td>Unchanged</td>
<td></td>
</tr>
<tr>
<td>Bicarbonate</td>
<td>Decreases markedly (18–22 mEq/L is normal)</td>
<td>Compensates for decrease in PCO₂</td>
</tr>
<tr>
<td>Total protein</td>
<td>Decreases from 72 to 62 g/L</td>
<td></td>
</tr>
<tr>
<td>Albumin</td>
<td>Decreases from 47 to 36 g/L</td>
<td></td>
</tr>
<tr>
<td>Urea, creatinine, and uric acid</td>
<td>Decrease in first trimester; stabilize in second trimester; increase toward term</td>
<td></td>
</tr>
<tr>
<td>Vitamin B6</td>
<td>Decreases</td>
<td></td>
</tr>
<tr>
<td>Blood glucose</td>
<td>Fasting levels decrease in first trimester, then unchanged</td>
<td>Postprandial levels remain elevated longer, prolonging return to fasting state. Increased glucose levels allow passive diffusion across placenta to fetus</td>
</tr>
<tr>
<td>Folate</td>
<td>Decreases 50% toward term</td>
<td></td>
</tr>
<tr>
<td>Vitamin B12</td>
<td>Decreases 50% or more</td>
<td>B12 levels in folate-deficient women will increase with folate supplementation alone</td>
</tr>
</tbody>
</table>

**Endocrine system**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Expected change</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pituitary gland</td>
<td>Increases to 50% greater than adult male</td>
<td>Attributable to increase of prolactin-secreting cells in anterior lobe</td>
</tr>
<tr>
<td>Prolactin</td>
<td>Increases from 300 to 5000 mIU/L</td>
<td>Estrogen stimulates hyperplasia and hypertrophy of pituitary lactotrophs</td>
</tr>
<tr>
<td>Follicle-stimulating hormone and luteinizing hormone</td>
<td>Decrease to nearly undetectable</td>
<td></td>
</tr>
<tr>
<td>Melanocyte-stimulating hormone</td>
<td>Increased</td>
<td>Likely responsible for linea nigra, chloasma, and increased areolae pigmentation</td>
</tr>
<tr>
<td>Thyroid-stimulating hormone</td>
<td>Normal range is unchanged</td>
<td>May be suppressed during late first to early second trimester due to hCG-mediated increase in TH production—subtle effect may be exacerbated by conditions that increase hCG levels, such as hyperemesis, molar pregnancies, and multiples</td>
</tr>
<tr>
<td>Thyroid-binding globulin</td>
<td>Increases—doubles by end of first trimester; triples by term</td>
<td>Due to estrogen effect on liver</td>
</tr>
<tr>
<td>Thyroxine (T4) and triiodothyronine (T3)</td>
<td>Increased total circulating amount; unchanged free fraction</td>
<td>Fetal thyroid hormone production commences at 18 weeks</td>
</tr>
<tr>
<td>Reverse T₃</td>
<td>Unchanged in maternal circulation; increased in cord blood</td>
<td>Episodic pattern of release is maintained</td>
</tr>
<tr>
<td>Cortisol</td>
<td>Increased to 3 × nonpregnant values</td>
<td>Tenfold increase rate of transformation to estradiol and estrone</td>
</tr>
<tr>
<td>Aldosterone</td>
<td>Increased twofold by term</td>
<td>Precursor for steroid hormone synthesis by placenta</td>
</tr>
<tr>
<td>Deoxycorticosterone</td>
<td>Increased by 20–100 times</td>
<td>Proportionally less increase of glucagons compared with insulin; so insulin:glucagon ratio is increased</td>
</tr>
<tr>
<td>Testosterone</td>
<td>Increased total amount; decreased free fraction</td>
<td>Especially fasting levels</td>
</tr>
<tr>
<td>Androstenedione</td>
<td>Increases</td>
<td>Maintains calcium levels in face of increased renal absorption and transfer to fetus</td>
</tr>
<tr>
<td>DHEA</td>
<td>Unchanged or small decrease</td>
<td>Production originates in corpus luteum over first 7–8 weeks; then placenta takes over</td>
</tr>
<tr>
<td>Pancreas</td>
<td>Hypertrophy of islets due to hyperplasia of B cells</td>
<td></td>
</tr>
<tr>
<td>Insulin</td>
<td>Increases fasting levels toward term; hyperplasia of pancreatic B cells</td>
<td></td>
</tr>
<tr>
<td>Glucagon</td>
<td>Increases fasting levels</td>
<td></td>
</tr>
<tr>
<td>Glucose</td>
<td>Slight decrease</td>
<td></td>
</tr>
<tr>
<td>Parathyroid hormone</td>
<td>Increased during end of pregnancy</td>
<td></td>
</tr>
<tr>
<td>Progesterone</td>
<td>Markedly increases</td>
<td></td>
</tr>
<tr>
<td>Estradiol</td>
<td>Markedly increases</td>
<td></td>
</tr>
<tr>
<td>17-Hydroxyprogesterone</td>
<td>Increases</td>
<td></td>
</tr>
<tr>
<td>Estriol</td>
<td>Increases</td>
<td></td>
</tr>
<tr>
<td>Human placental lactogen (hPL)</td>
<td>Increases 5000-fold from 0.002 μU/mL to 10 μU/mL</td>
<td>Peak 93 U/mL; term 14 U/mL</td>
</tr>
<tr>
<td>hCG</td>
<td>Increases to maximum values by 8–10 weeks</td>
<td></td>
</tr>
<tr>
<td>Relaxin</td>
<td>Peaks at 10 weeks</td>
<td></td>
</tr>
</tbody>
</table>

(Continued)
The detected 46%–51% increase in the cardiac output was attributed to a 15% increase in heart rate and a 24% increase in stroke volume [6]. The changes in heart rate occur early in pregnancy, whereas those of stroke volume occur later, with the net effect of a progressively increasing cardiac output as gestation progresses. Again, significant variation in cardiac output changes in the late third trimester was attributed to patient factors, precluding confident conclusions regarding the behavior of cardiac output at the very end of pregnancy. Maternal cardiac output was found to correlate with maternal body surface area and with fetal birth weight. Left ventricular mass and left ventricular mass index increased to maximal levels at term but remained well below the cutoff for a diagnosis of left ventricular hypertrophy. This increase corresponded to an increase in the mean blood pressure at term as well as to an increase in the left atrial size and a decrease in the left ventricular diastolic filling value. These changes may explain some of the variations in cardiac output in the third trimester and may be relevant to the

## Table 3.1 Summary of Physiologic Adaptations during Pregnancy (Continued)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Expected change</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Gastrointestinal system</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Appetite</td>
<td>Increases</td>
<td></td>
</tr>
<tr>
<td>Gastric reflux</td>
<td>Increases</td>
<td>Cardiac sphincter laxity and anatomic displacement; treated during labor and delivery/anesthetic procedures with nonparticulate oral antacids</td>
</tr>
<tr>
<td>Gastric secretion</td>
<td>Decreased acidity; increased volume</td>
<td>“Full stomach” effect increases risk of aspiration</td>
</tr>
<tr>
<td>Gastric motility</td>
<td>Decreases</td>
<td></td>
</tr>
<tr>
<td>Intestinal absorption</td>
<td>Increased</td>
<td></td>
</tr>
<tr>
<td>Intestinal transit time</td>
<td>Delayed</td>
<td></td>
</tr>
<tr>
<td>Large intestine</td>
<td>Greater absorption; slower transit time</td>
<td></td>
</tr>
<tr>
<td>Gallbladder</td>
<td>Larger due to passive dilation</td>
<td></td>
</tr>
<tr>
<td><strong>Hematologic system</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Plasma volume</td>
<td>40%–60% increase from 12 to 36 weeks; 70%–100% increase in multiple gestations</td>
<td>Offsets blood loss at delivery</td>
</tr>
<tr>
<td>Total erythrocyte volume</td>
<td>Increases 15%–30%</td>
<td>Greater increase with iron supplementation</td>
</tr>
<tr>
<td>Hematocrit</td>
<td>Decreases 3%–5% by 36 weeks</td>
<td>Physiologic anemia due to greater proportionate increase plasma volume compared with erythrocyte volume; less change with iron supplementation</td>
</tr>
<tr>
<td>Hemoglobin</td>
<td>Decreases</td>
<td>Best indicator of iron status. Slight increase with iron supplementation</td>
</tr>
<tr>
<td>Mean corpuscular volume</td>
<td>Unchanged</td>
<td>Provides little diagnostic value; combined with increase in white blood cell (WBC) can cause false suspicion for infection</td>
</tr>
<tr>
<td>Erythrocyte sedimentation rate</td>
<td>Significantly increased</td>
<td></td>
</tr>
<tr>
<td>WBC count</td>
<td>Increases 8% by term and may increase further postpartum</td>
<td>Predominantly due to increase in neutrophils</td>
</tr>
<tr>
<td>Serum iron</td>
<td>Decreases 35% by term</td>
<td></td>
</tr>
<tr>
<td>Serum transferrin</td>
<td>Increases by 100% or more by second trimester</td>
<td></td>
</tr>
<tr>
<td>TIBC</td>
<td>Increases by 25%–100%</td>
<td></td>
</tr>
<tr>
<td>Serum ferritin</td>
<td>Decreases markedly (even with iron supplementation)</td>
<td></td>
</tr>
<tr>
<td>Erythropoietin</td>
<td>Increases fourfold</td>
<td></td>
</tr>
<tr>
<td>α-Fetoprotein</td>
<td>Increases</td>
<td></td>
</tr>
<tr>
<td>Glutamic-oxaloacetic transaminase and glutamic-pyruvic transaminase</td>
<td>Unchanged</td>
<td></td>
</tr>
<tr>
<td>Alkaline phosphatase</td>
<td>Increases</td>
<td>Heat stable fraction formed by placenta</td>
</tr>
<tr>
<td>Lipids</td>
<td>Increase</td>
<td>Triglycerides, cholesterol, phospholipids, and free fatty acids each increase progressively</td>
</tr>
<tr>
<td>Fibrinogen</td>
<td>Increases 2 g/L by term</td>
<td>Overall increased tendency toward thrombosis</td>
</tr>
<tr>
<td>Factors VII, VIII, IX, and X</td>
<td>Increase</td>
<td></td>
</tr>
<tr>
<td>Factors XI and XIII</td>
<td>Decrease by about 30%</td>
<td></td>
</tr>
<tr>
<td>Antithrombin III</td>
<td>Decreases</td>
<td></td>
</tr>
<tr>
<td>Fibrin, FDP</td>
<td>Increase progressively</td>
<td></td>
</tr>
<tr>
<td>Protein C</td>
<td>Unchanged</td>
<td></td>
</tr>
<tr>
<td>Protein S</td>
<td>Decreases</td>
<td>Protein S–binding protein levels fluctuate in pregnancy, making screening more reliable in nonpregnant women</td>
</tr>
</tbody>
</table>


Abbreviations: BP, blood pressure; CO, cardiac output; TH, thyroid hormone; DHEA, dehydroepiandrosterone; FDP, fibrin degradation products; TIBC, total iron binding capacity.
<table>
<thead>
<tr>
<th>Table 3.2 Normal Reference Ranges in Pregnant Women</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Hematology</strong></td>
</tr>
<tr>
<td>Erythropoietin (U/L)</td>
</tr>
<tr>
<td>Ferritin (ng/mL)</td>
</tr>
<tr>
<td>Folate, red blood cell (ng/mL)</td>
</tr>
<tr>
<td>Folate, serum (ng/mL)</td>
</tr>
<tr>
<td>Hemoglobin (g/dL)</td>
</tr>
<tr>
<td>Hematocrit (%)</td>
</tr>
<tr>
<td>Iron, total binding capacity (µg/dL)</td>
</tr>
<tr>
<td>Iron, serum (µg/dL)</td>
</tr>
<tr>
<td>Mean corpuscular hemoglobin (pg/cell)</td>
</tr>
<tr>
<td>Mean corpuscular volume (µm³)</td>
</tr>
<tr>
<td>Platelet (×10⁹/L)</td>
</tr>
<tr>
<td>Mean platelet volume (µm³)</td>
</tr>
<tr>
<td>Red blood cell count (×10⁶/mm³)</td>
</tr>
<tr>
<td>Red cell distribution width (%)</td>
</tr>
<tr>
<td>White blood cell count (×10⁹/mm³)</td>
</tr>
<tr>
<td>Neutrophils (×10⁹/mm³)</td>
</tr>
<tr>
<td>Lymphocytes (×10⁹/mm³)</td>
</tr>
<tr>
<td>Monocytes (×10⁹/mm³)</td>
</tr>
<tr>
<td>Eosinophils (×10⁹/mm³)</td>
</tr>
<tr>
<td>Basophils (×10⁹/mm³)</td>
</tr>
<tr>
<td>Mean corpuscular hemoglobin (pg/cell)</td>
</tr>
<tr>
<td>Mean platelet volume (µm³)</td>
</tr>
<tr>
<td>Red blood cell count (×10⁶/mm³)</td>
</tr>
<tr>
<td>Red cell distribution width (%)</td>
</tr>
<tr>
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<tr>
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<tr>
<td>Lymphocytes (×10⁹/mm³)</td>
</tr>
<tr>
<td>Monocytes (×10⁹/mm³)</td>
</tr>
<tr>
<td>Eosinophils (×10⁹/mm³)</td>
</tr>
<tr>
<td>Basophils (×10⁹/mm³)</td>
</tr>
<tr>
<td>Mean corpuscular hemoglobin (pg/cell)</td>
</tr>
<tr>
<td>Mean platelet volume (µm³)</td>
</tr>
<tr>
<td>Red blood cell count (×10⁶/mm³)</td>
</tr>
<tr>
<td>Red cell distribution width (%)</td>
</tr>
<tr>
<td>White blood cell count (×10⁹/mm³)</td>
</tr>
<tr>
<td>Neutrophils (×10⁹/mm³)</td>
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<tr>
<td>Lymphocytes (×10⁹/mm³)</td>
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<tr>
<td>Monocytes (×10⁹/mm³)</td>
</tr>
<tr>
<td>Eosinophils (×10⁹/mm³)</td>
</tr>
<tr>
<td>Basophils (×10⁹/mm³)</td>
</tr>
<tr>
<td>Mean corpuscular hemoglobin (pg/cell)</td>
</tr>
<tr>
<td>Mean platelet volume (µm³)</td>
</tr>
<tr>
<td>Red blood cell count (×10⁶/mm³)</td>
</tr>
<tr>
<td>Red cell distribution width (%)</td>
</tr>
<tr>
<td>White blood cell count (×10⁹/mm³)</td>
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<tr>
<td>Neutrophils (×10⁹/mm³)</td>
</tr>
<tr>
<td>Lymphocytes (×10⁹/mm³)</td>
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<tr>
<td>Monocytes (×10⁹/mm³)</td>
</tr>
<tr>
<td>Eosinophils (×10⁹/mm³)</td>
</tr>
<tr>
<td>Basophils (×10⁹/mm³)</td>
</tr>
</tbody>
</table>

(Continued)
### Table 3.2 Normal Reference Ranges in Pregnant Women (Continued)

<table>
<thead>
<tr>
<th>Test</th>
<th>Nonpregnant adult</th>
<th>First trimester</th>
<th>Second trimester</th>
<th>Third trimester</th>
</tr>
</thead>
<tbody>
<tr>
<td>Magnesium (mg/dL)</td>
<td>1.5–2.3</td>
<td>1.6–2.2</td>
<td>1.5–2.2</td>
<td>1.1–2.2</td>
</tr>
<tr>
<td>Osmolality (mOsm/kg H₂O)</td>
<td>275–295</td>
<td>275–280</td>
<td>276–289</td>
<td>278–280</td>
</tr>
<tr>
<td>Phosphate (mg/dL)</td>
<td>2.5–4.3</td>
<td>3.1–4.6</td>
<td>2.5–4.6</td>
<td>2.8–4.6</td>
</tr>
<tr>
<td>Potassium (mEq/L)</td>
<td>3.5–5</td>
<td>3.6–5</td>
<td>3.3–5</td>
<td>3.3–5–5.1</td>
</tr>
<tr>
<td>Prealbumin (mg/dL)</td>
<td>17–34</td>
<td>15–27</td>
<td>20–27</td>
<td>14–23</td>
</tr>
<tr>
<td>Protein, total (g/dL)</td>
<td>6.7–8.6</td>
<td>6.2–7.6</td>
<td>5.7–6.9</td>
<td>5.6–6.7</td>
</tr>
<tr>
<td>Sodium (mEq/L)</td>
<td>136–146</td>
<td>133–148</td>
<td>129–148</td>
<td>130–148</td>
</tr>
<tr>
<td>Urea nitrogen (mg/dL)</td>
<td>7–20</td>
<td>7–12</td>
<td>3–13</td>
<td>3–11</td>
</tr>
<tr>
<td>Uric acid (mg/dL)</td>
<td>2.5–5.6</td>
<td>2–4.2</td>
<td>2.4–4.9</td>
<td>3.1–6.3</td>
</tr>
</tbody>
</table>

#### Metabolic and endocrine tests

<table>
<thead>
<tr>
<th>Test</th>
<th>Value (mg/dL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aldosterone (ng/dL)</td>
<td>2–9</td>
</tr>
<tr>
<td>Angiotensin-converting enzyme (U/L)</td>
<td>9–67</td>
</tr>
<tr>
<td>Cortisol (µg/dL)</td>
<td>0–25</td>
</tr>
<tr>
<td>Hemoglobin A1c (%)</td>
<td>4–6</td>
</tr>
<tr>
<td>Parathyroid hormone (pg/mL)</td>
<td>8–51</td>
</tr>
<tr>
<td>Parathyroid hormone-related protein (pmol/L)</td>
<td>&lt;1.3</td>
</tr>
<tr>
<td>Renin, plasma activity (ng/mL/hour)</td>
<td>0.3–9</td>
</tr>
<tr>
<td>TSH (µIU/mL)</td>
<td>0.34–4.25</td>
</tr>
<tr>
<td>Thyroxine-binding globulin (mg/dL)</td>
<td>1.3–3</td>
</tr>
<tr>
<td>Thyroxine, free (ng/dL)</td>
<td>0.8–1.7</td>
</tr>
<tr>
<td>Thyroxine, total (µg/dL)</td>
<td>5.4–11.7</td>
</tr>
<tr>
<td>Triiodothyronine, free (pg/mL)</td>
<td>2.4–4.2</td>
</tr>
<tr>
<td>Triiodothyronine, total (ng/dL)</td>
<td>77–135</td>
</tr>
</tbody>
</table>

#### Vitamins and minerals

<table>
<thead>
<tr>
<th>Test</th>
<th>Value (µg/dL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Copper (µg/dL)</td>
<td>70–140</td>
</tr>
<tr>
<td>Selenium (µg/L)</td>
<td>63–160</td>
</tr>
<tr>
<td>Vitamin A (retinol) (µg/dL)</td>
<td>20–100</td>
</tr>
<tr>
<td>Vitamin B12 (µL)</td>
<td>279–966</td>
</tr>
<tr>
<td>Vitamin C (ascorbic acid) (mg/dL)</td>
<td>0.4–1</td>
</tr>
<tr>
<td>Vitamin D, 1,25-dihydroxy (pg/mL)</td>
<td>25–45</td>
</tr>
<tr>
<td>Vitamin D, 24,25-dihydroxy (pg/mL)</td>
<td>0.5–3</td>
</tr>
<tr>
<td>Vitamin D, 25-hydroxy (mg/mL)</td>
<td>14–80</td>
</tr>
<tr>
<td>Vitamin E (α-tocopherol) (mg/mL)</td>
<td>5–18</td>
</tr>
<tr>
<td>Zinc (µg/dL)</td>
<td>75–120</td>
</tr>
</tbody>
</table>

#### Autoimmune and inflammatory mediators

<table>
<thead>
<tr>
<th>Test</th>
<th>Value (mg/dL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C3 complement (mg/dL)</td>
<td>83–177</td>
</tr>
<tr>
<td>C4 complement (mg/dL)</td>
<td>16–47</td>
</tr>
<tr>
<td>C-reactive protein (mg/L)</td>
<td>0.2–3</td>
</tr>
<tr>
<td>Erythrocyte sedimentation rate (mm/hour)</td>
<td>0–20</td>
</tr>
<tr>
<td>Immunoglobulin A (mg/dL)</td>
<td>70–350</td>
</tr>
<tr>
<td>Immunoglobulin G (mg/dL)</td>
<td>700–1700</td>
</tr>
<tr>
<td>Immunoglobulin M (mg/dL)</td>
<td>50–300</td>
</tr>
</tbody>
</table>

#### Sex hormones

<table>
<thead>
<tr>
<th>Test</th>
<th>Value (µmol/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dehydroepiandrosterone sulfate (µmol/L)</td>
<td>1.3–6.8</td>
</tr>
<tr>
<td>Estradiol (pg/mL)</td>
<td>&lt;20–443</td>
</tr>
<tr>
<td>Progesterone (ng/mL)</td>
<td>&lt;1–20</td>
</tr>
<tr>
<td>Prolactin (ng/mL)</td>
<td>0–20</td>
</tr>
<tr>
<td>Sex hormone-binding globulin (nmol/L)</td>
<td>18–114</td>
</tr>
<tr>
<td>Testosterone (ng/dL)</td>
<td>6–86</td>
</tr>
<tr>
<td>17-Hydroxyprogesterone (nmol/L)</td>
<td>0.6–10.6</td>
</tr>
</tbody>
</table>

#### Lipids

<table>
<thead>
<tr>
<th>Test</th>
<th>Value (mg/dL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cholesterol, total (mg/dL)</td>
<td>&lt;200</td>
</tr>
<tr>
<td>High-density lipoprotein cholesterol (mg/dL)</td>
<td>40–60</td>
</tr>
<tr>
<td>Low-density lipoprotein cholesterol (mg/dL)</td>
<td>&lt;100</td>
</tr>
<tr>
<td>Very-low-density lipoprotein cholesterol (mg/dL)</td>
<td>6–40</td>
</tr>
<tr>
<td>Triglycerides (mg/dL)</td>
<td>&lt;150</td>
</tr>
<tr>
<td>Apolipoprotein A-I (mg/dL)</td>
<td>119–240</td>
</tr>
<tr>
<td>Apolipoprotein B (mg/dL)</td>
<td>52–163</td>
</tr>
</tbody>
</table>

#### Cardiac

<table>
<thead>
<tr>
<th>Test</th>
<th>Value (µg/mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Atrial natriuretic peptide (pg/mL)</td>
<td>28.1–70.1</td>
</tr>
<tr>
<td>B-type natriuretic peptide (pg/mL)</td>
<td>&lt;167</td>
</tr>
<tr>
<td>Creatine kinase (U/L)</td>
<td>39–238</td>
</tr>
<tr>
<td>Creatine kinase-MB (U/L)</td>
<td>&lt;6</td>
</tr>
<tr>
<td>Troponin I (ng/mL)</td>
<td>0–0.8</td>
</tr>
</tbody>
</table>

(Continued)
vulnerability of pregnant women to pulmonary edema during hypertensive crisis [6].

Perhaps the most common hemodynamic complaint that must be evaluated during pregnancy is that of syncope or near-syncope, which provides a useful example of how an understanding of pregnancy physiology is useful in clinical evaluation. Syncope is defined as a transient loss of consciousness and posture, caused by decreased cerebral perfusion that may result from hypotension, changes in heart rate, or changes in blood volume or redistribution. The decreased SVR of pregnancy makes pregnant women particularly susceptible to this condition, with 28% of gravidas experiencing at least one episode of presyncope, 10% experiencing recurrent presyncopal episodes, and 5% experiencing outright syncope [7]. The overwhelming majority of syncopal episodes are benign neurocardiogenic syncope but there are also several potentially dangerous conditions in the differential diagnosis of syncope. An understanding of the vasovagal reflex at the root of most syncopal episodes helps the clinician to manage benign syncopal episodes while being aware of conditions that are reflected in pulmonary function tests and in acid–base balance. These are important in the evaluation of acute pulmonary complications of pregnancy.

The above events that are precipitated by a decrease in venous return can also explain the occurrence of supine hypotension in pregnancy. The commonly recommended “leftward tilt” position is intended to displace the uterus off of the inferior vena cava, which runs to the right of midline. This position should be used to avoid supine hypotension when recumbent as well as when performing surgery on the parturient in the second half of pregnancy. A more extreme application of this physiology comes in the performance of perimortem cesarean section during maternal cardiac arrest. The procedure is purported to allow fetal survival and also the evacuation of the uterus, which may allow an increase in venous return and cardiac output that may increase the chance of maternal survival [8]. In order to optimize maternal and fetal survival, it is recommended that the procedure should be performed within 4 minutes of cardiac arrest due to the inadequacy of chest compressions in producing adequate cardiac output during pregnancy and the susceptibility of both mother and fetus to anoxic brain injury [8] (see also Chapters 1 and 2 in Maternal-Fetal Evidence Based Guidelines).

**RESPIRATORY**

During pregnancy the respiratory system undergoes alterations that are reflected in pulmonary function tests and in acid–base balance. These are important in the evaluation of dyspnea in pregnancy, the management of pregnancy with coexisting pulmonary diseases such as asthma, and the recognition of acute pulmonary complications of pregnancy.

Pregnancy is associated with a significant increase in ventilatory drive both at rest and during exercise [9]. Minute ventilation increases mostly due to an increase in tidal volume with little or no increase in respiratory rate [39,10]. Alveolar ventilation increases, along with an increase in arterial partial pressure of oxygen (PaO₂) and alveolar partial

**Table 3.2 Normal Reference Ranges in Pregnant Women (Continued)**

<table>
<thead>
<tr>
<th>Blood gas</th>
<th>Nonpregnant adult</th>
<th>First trimester</th>
<th>Second trimester</th>
<th>Third trimester</th>
</tr>
</thead>
<tbody>
<tr>
<td>pH</td>
<td>7.38–7.42</td>
<td>7.36–7.52</td>
<td>7.4–7.52</td>
<td>7.41–7.53</td>
</tr>
<tr>
<td>PO₂ (mmHg)</td>
<td>90–100</td>
<td>93–100</td>
<td>90–98</td>
<td>89–107</td>
</tr>
<tr>
<td>PCO₂ (mmHg)</td>
<td>38–42</td>
<td>38–42</td>
<td>36–42</td>
<td>35–42</td>
</tr>
<tr>
<td>Bicarbonate (HCO₃⁻) (mEq/L)</td>
<td>22–26</td>
<td>22–26</td>
<td>22–26</td>
<td>22–26</td>
</tr>
</tbody>
</table>

**Renal function tests**

| Effective RPF (mL/min) | 492–696 | 696–985 | 600–800 | 595–945 |
| Filtration fraction (%) | 106–132 | 131–166 | 135–170 | 117–182 |
| Osmolarity, urine (mOsm/kg) | 500–800 | 526–975 | 500–1066 | 238–1034 |
| 24-hour albumin excretion (mg/24 hours) | <30 | 5–15 | 4–18 | 3–22 |
| 24-hour calcium excretion (mmol/24 hours) | <7.5 | 1.6–5.2 | 0.3–6.9 | 0.8–4.2 |
| 24-hour creatinine clearance (mL/min) | 91–130 | 131–166 | 135–170 | 117–182 |
| 24-hour creatinine excretion (mmol/24 hours) | 8.8–14 | 10.6–11.6 | 10.3–11.5 | 10.2–11.4 |
| 24-hour potassium excretion (mmol/24 hours) | 91–130 | 131–166 | 135–170 | 117–182 |
| 24-hour sodium excretion (mmol/24 hours) | 25–100 | 19–141 | 47–186 | 46–185 |

Source: Adapted from Abbassi-Ghanavati M et al., Obstet Gynecol, 114(6), 1326–1331, 2009.
pressure of oxygen (PAO₂), and a decrease in arterial partial pressure of carbon dioxide (PaCO₂), with a compensatory decrease in serum bicarbonate with an overall mild increase in pH, reflecting a state of compensated respiratory alkalosis [9]. These changes occur early in pregnancy and are almost fully established by 7–8 weeks’ gestation [9]. This may be due to stimulation of the ventilatory drive by progesterone and/or estrogen. Ventilatory equivalents for CO₂ and O₂ are increased both at rest and during exercise throughout pregnancy. The underlying mechanism for the increased ventilatory drive during pregnancy is not fully understood, but theories have included an increased sensitivity to chemoreflexive drives to breathe (due to hypercapnia or hypoxia) versus a hormone-mediated increase in the neural drive to breathe [9].

Uterine enlargement and abdominal distension result in a 4- to 5-cm cephalad displacement of the diaphragm and a 5- to 7-cm increase in thoracic circumference. This results in a decrease in expiratory reserve volume, residual volume, and functional residual capacity. There is a compensatory increase in inspiratory capacity, while total lung capacity and vital capacity do not change [9]. Chest wall compliance is increased but inspiratory muscle strength is preserved with an overall increase in the oxygen cost of breathing [9]. However, it is important to recognize that there is no significant change in the parameters of forced vital capacity, peak expiratory flow rate (PEFR), or forced expiratory volume in 1 second (FEV₁) during pregnancy.

Physiologic dyspnea of pregnancy, experienced by 60%–70% of healthy pregnant women, must be clinically distinguished from more serious respiratory conditions. Physiologic dyspnea tends to be an isolated symptom that begins in early pregnancy, plateaues, or improves as pregnancy progresses, and does not interfere with daily activities [11]. The mechanism for physiologic dyspnea has not been conclusively defined, but pregnant women with dyspnea have been demonstrated to have an increase in minute ventilation and tidal volume and a decrease in end-tidal CO₂ pressure compared with pregnant women who do not report dyspnea [11]. The perception of physiologic dyspnea during pregnancy has been associated with increased sensitivity to hypoxia and hypercapnia, suggesting an increased chemosensitivity causing an increased central inspiratory drive in pregnant women who experience dyspnea. However, the chemical stimuli of hypoxia and hypercapnia are both reduced in pregnancy, causing others to suggest a neural mechanism [9]. Despite the common symptom of physiologic dyspnea, pregnancy has not been found to be associated with a decrease in aerobic work capacity or with an increased perception of breathlessness during exercise [9].

While measurement of FEV₁ requires a spirometer, measurement of PEFR correlates well with FEV₁, and can be measured with a relatively inexpensive spirometer (peak flow meter), which patients can be taught to use at home. Again, these parameters do not change due to pregnancy, so any detected worsening should be treated appropriately and not attributed to pregnancy or to physiologic dyspnea. In the evaluation of severe acute asthma exacerbations with the potential for impending respiratory arrest, knowledge of physiologic changes of pregnancy is particularly important in the interpretation of blood gases (Table 3.1). The normal parturient lives in a state of compensated respiratory alkalosis with a lower partial pressure of carbon dioxide (PCO₂) compared with that of nonpregnant patients. Thus, significant CO₂ retention may be present despite values that are high normal for nonpregnant patients.

While physiologic dyspnea and asthma exacerbation are two of the most common causes of dyspnea in pregnancy, the obstetrician must also be alert to other pulmonary complications to which the parturient is susceptible such as pulmonary embolism and pulmonary edema. Pulmonary edema may occur as a result of preeclampsia, peripartum cardiomyopathy, or the use of certain tocolytics. It is important that the prevalence of pulmonary symptoms in pregnancy should not be met with complacency by the obstetrician, for it may signify a life-threatening condition for the pregnant woman.

ENDOCRINE

Pregnancy-related endocrine alterations include the production of hormones that are specific to pregnancy, an increase in other reproductive hormones, and alterations in the level and function of nonreproductive hormones, especially thyroid hormones. There is a significant contribution of steroid hormone secretion by the fetal-placental unit. This section provides a brief description of the changes in reproductive hormones during gestation followed by a more in-depth review of the behavior and clinical application of thyroid hormones during pregnancy.

Pregnancy-specific hormones include human chorionic gonadotropin (hCG) and relaxin. Relaxin is detectable in maternal serum by the time of missed menses and peaks at 10 weeks’ gestation then declines over the course of the second and third trimesters [12]. Relaxin is secreted by the corpora lutea of pregnancy and thought to have an important role in early pregnancy maintenance that has not yet been clearly elucidated [13]. hCG also peaks at approximately 10 weeks’ gestation. The reproductive hormones estradiol, progesterone, testosterone, prolactin, and 17-hydroxyprogesterone, all increase significantly during gestation. Initially the corpus luteum and maternal ovarian tissue make the greatest contribution to steroid hormone concentrations, but as of 9 weeks’ gestation aromatization of dehydroepiandrosterone sulfate by the placenta becomes the predominant source of maternal steroids [14]. The elevated estradiol levels stimulate increased hepatic production of sex hormone–binding globulin and thyroxin-binding globulin. Estrogen also induces hypertrophy and hyperplasia of pituitary lactotrophs with a resultant increase in prolactin levels corresponding to the increase in estradiol levels throughout gestation [14]. Meanwhile, there is a reflexive decrease in follicle-stimulating hormone and luteinizing hormone to almost undetectable levels, as would be expected.

One longitudinal study assayed reproductive hormone levels in the blood of 60 healthy women drawn during the first, second, and third trimesters of uncomplicated pregnancies [14]. Mean progesterone levels increased steadily from 49 nmol/L at 5 weeks’ gestation to 384 nmol/L at term. Mean 17-hydroxyprogesterone levels are more stable during the first and second trimesters at 12.2 nmol/L but then increase threefold to 36 nmol/L by term. Mean testosterone increased from 3.3 nmol/L at 5 weeks to 5.7 nmol/L at 40 weeks. Mean serum estradiol levels increased during the first trimester from 1.64 nmol/L at 5 weeks to 11.13 nmol/L at 16 weeks and then increased fivefold to 53.44 nmol/L at 40 weeks. Mean sex hormone-binding globulin levels increase rapidly during the first half of gestation from 71 nmol/L at 5 weeks to 392 nmol/L at 25 weeks, and then remain relatively constant until 40 weeks. Mean levels of dehydroepiandrosterone sulfate decreased from 5.8 mmol/L at 5 weeks to 2.7 mmol/L
at midgestation, which remained constant until term. Mean prolactin concentration rose from 294 milli-international units (mIU) to 1106 mIU at 16 weeks. Prolactin levels then continued to increase to a mean of 4092 mIU at 35 weeks and to 4293 mIU at 40 weeks. Mean androstenedione levels increase gradually from 8.1 nmol/L at 5 weeks to 10.6 nmol/L at 40 weeks [14].

Other hormonal alterations include an increase in aldosterone, cortisol, parathyroid hormone, parathyroid-related hormone, and renin [2]. Deoxycorticosterone increases. Androstenedione increases with an increase in the transformation to estrone and estradiol [1]. Fasting levels of both insulin and glucagon increase [1]. There is an increase in melanocyte-stimulating hormone to which can be attributed the pregnancy-related increases in pigmentation seen in the areola, the linea nigra, and in chloasma [1].

The function of the thyroid gland is crucial to a healthy gestation (see also Chapters 6 and 7 in Maternal-Fetal Evidence Based Guidelines). The interplay between maternal and fetal thyroid function can cause confusion for the obstetrician. Early fetal development is dependent on maternal thyroid function, and both hypothyroidism and hyperthyroidism can have important maternal and fetal effects and risks of thyroid dysfunction extend well into the postpartum period. The effects of subclinical thyroid disease are more controversial. Symptoms of hyperthyroidism and hypothyroidism can mimic symptoms of normal pregnancy. For example, symptoms such as fatigue, muscle cramps, palpitations, thymomgaly, and constipation can be common in normal pregnancy, but progressive symptoms of insomnia, intellectual slowness, or weight loss should be evaluated [15].

Thyroid-binding globulin increases due to stimulation of synthesis by estrogen as well as decreased hepatic clearance. Total thyroxine (TT4) and total triiodothyronine (TT3) both increase while resin triiodothyronine uptake (RT3U) decreases. Structural similarities between hCG and thyroid-stimulating hormone (TSH) may result in an hCG-mediated increase in free thyroxine (FT4) and the free thyroxine index as well as a decrease in TSH in the first trimester. However, these changes, sometimes referred to as gestational transient thyrotoxicosis, are typically self-limited and do not tend to result in values that are outside the normal range for nonpregnant individuals [16] (see also Chapters 6 and 7 in Maternal-Fetal Evidence Based Guidelines).

Iodine requirements during pregnancy increase by greater than 50% due to increased maternal thyroid production to maintain maternal and fetal euthyroidism and increased renal iodine clearance [17]. Plasma iodine levels decrease. This is associated with an increase in the size of the maternal thyroid gland. Longitudinal studies of thyroid ultrasonography in pregnancy show a mean increase in the thyroid size of 18%, which is noticeable in most women but not associated with abnormalities in thyroid function tests [16]. Iodine supplementation results in a less substantial increase in the thyroid gland size [17]. While ultrasound or laboratory evaluation is not necessary in the pregnant patient with a mild diffuse increase in the thyroid size, a significant goiter or thyroid nodule must be evaluated as in any patient. A woman who is marginally iodine deficient may be able to compensate with increased thyrotropin stimulation of the thyroid to achieve euthyroidism, but become hypothyroid when faced with the increasing iodine requirements of pregnancy [17].

Thyroid homeostasis is important for healthy fetal development. The fetal thyroid begins to concentrate iodide at 10–12 weeks. Thyroid hormone necessary for fetal brain development before this time is provided by the maternal system [15,18]. Thyroid hormone synthesis in the fetus is controlled by the fetal pituitary gland by 20 weeks. Small amounts of T4 and T3 pass the placenta but TSH does not cross the placenta. Thyroid-releasing hormone (TRH) and iodide do cross the placenta [16]. Maternal hypothyroidism has been associated with abnormal intelligence quotient testing and pediatric neurodevelopment in offspring, particularly when untreated [18]. While severe maternal iodine deficiency can lead to cretinism in the offspring, it is less clear whether mild-to-moderate iodine deficiency leads to more subtle cognitive or neurologic dysfunction. Iodine supplementation in iodine-deficient populations has been found to substantially reduce the relative risk of cretinism and to improve psychomotor and cognitive test scores in the offspring [17].

Hyperemesis gravidarum is associated with elevated levels of hCG, an increase in FT4, and a decrease in TSH (biochemical hyperthyroidism). However, this is largely transitory and rarely associated with clinical hyperthyroidism. Thus, routine measuring of thyroid function in hyperemesis is not indicated in the absence of other signs of hyperthyroidism such as weight loss or persistent tachycardia [16]. Furthermore, treatment of transient hyperthyroidism associated with elevated hCG and hyperemesis should not be undertaken in the absence of evidence of intrinsic thyroid disease [19] (see also Chapter 9 in Maternal-Fetal Evidence Based Guidelines). A large prospective observational study of 25,765 pregnant women who underwent thyroid screening in pregnancy showed no difference in pregnancy complications or in perinatal morbidity and mortality in women with subclinical hyperthyroidism [20].

HEMATOLOGIC

Pregnancy is characterized by both quantitative and qualitative changes in the hematologic system. These changes can be adaptive to normal pregnancy but can also put the pregnant women at increased risk for certain pathologic conditions. Anemia and thrombocytopenia are commonly diagnosed during pregnancy, as will be discussed below. Measurements of the acute phase response such as erythrocyte sedimentation rate, C-reactive protein, and white blood cell count have been found to increase during pregnancy. This presents a challenge in the evaluation of pregnant women suspected to have various infectious or inflammatory conditions, but careful clinical assessment allows the practitioner to distinguish between physiologic and pathologic abnormalities in these tests. Pregnancy also has important effects on the coagulation system, with the creation of an overall hypercoagulable state. This section also reviews the theories and limitations of the evidence surrounding the diagnosis of thrombophilia during pregnancy.

Anemia is usually defined as a hemoglobin less than 11 g/dL and hematocrit less than 33% in the first trimester, hemoglobin less than 10.5 g/dL and hematocrit less than 32% in the second trimester, and hemoglobin less than 11 g/dL and hematocrit less than 33% in the third trimester [21] (see also Chapter 14 in Maternal-Fetal Evidence Based Guidelines). Anemia may be caused by decreased production of red blood cells, by increased destruction of red blood cells, or by blood loss. Anemia in pregnancy is complicated by increased iron requirements and an expanded blood volume. Blood volume increases by about 50% while red blood cell mass increases by
about 25%, resulting in an anemia of dilution, as measured by hemoglobin and hematocrit, that is physiologic in pregnancy. **Iron requirements increase** in order to support the increase in red blood cell mass, support the requirements of the fetus and placenta, and prepare for blood loss during delivery. **Iron-deficiency anemia** is characterized by microcytosis and hypochromatosis. Iron studies reveal a decrease in total iron and ferritin and an increase in total iron-binding capacity (TIBC). **Ferritin levels** have the highest sensitivity and specificity for iron deficiency, with levels less than 10–15 μg/dL being diagnostic of iron deficiency [21]. Iron supplementation as well as screening for iron deficiency during pregnancy is recommended by the Centers for Disease Control and Prevention (CDC). The typical diet contains 15 mg of elemental iron per day, while the recommended dietary daily allowance during pregnancy is 27 mg/day [21]. Iron-deficiency anemia in pregnancy has been associated with an increased risk of low birth weight, prematurity, perinatal mortality, postpartum depression, and poor mental and psychomotor testing in offspring. Severe anemia (less than 6 g/dL) has been associated with abnormal fetal oxygenation, abnormal fetal heart rate patterns, reduced amniotic fluid volumes, cerebral vasodilation, and fetal death. Transfusion may be indicated for fetal indications in the case of anemia of this severity [21]. Iron supplementation has been found to decrease the incidence of anemia at delivery, although its effect on healthy, non-iron-deficient women is not clear. Factors that increase the risk for iron deficiency in pregnancy include young maternal age, heavy menses, short interpregnancy interval, low socioeconomic status, and non-Hispanic black race [21]. Dietary factors can have a significant effect on iron levels, not only due to levels of consumption of iron but also due to consumption of foods that significantly enhance or inhibit iron absorption.

**Megaloblastic macrocytic anemia** may be caused by deficiency of folic acid and vitamin B12 and by pernicious anemia. Nonmegaloblastic macrocytic anemia may be caused by alcoholism, hypothyroidism, liver disease, aplastic anemia, or increased reticulocyte count. The most common cause of onset of macrocytic anemia during pregnancy in the United States is folic acid deficiency [21]. During pregnancy, daily folic acid requirements increase from 50 to 400 μg [21]. Women who have had gastrectomy and those with Crohn’s disease may be at risk of vitamin B12 deficiency in pregnancy [21].

The performance of complete blood counts as a part of routine prenatal screening results in a frequent diagnosis of **thrombocytopenia** in asymptomatic pregnant women. The mean platelet count in pregnant women is lower than in nonpregnant women, with about 8% of pregnant women meeting criteria for the diagnosis of thrombocytopenia [22]. Manifestations of thrombocytopenia include epistaxis, petechiae, and ecchymosis, although frequently there are no clinically significant effects. Clinically significant spontaneous bleeding is rare as long as platelet counts are greater than 10,000/μL, and even excessive bleeding associated with trauma or surgery is unlikely with platelet counts greater than 50,000/μL [22]. The most common cause of thrombocytopenia in pregnancy is gestational thrombocytopenia, which is an apparently benign condition whose underlying physiology is not well understood. However, thrombocytopenia in pregnancy may also be caused by more severe underlying conditions such as preeclampsia, human immunodeficiency virus infection, immune thrombocytopenic purpura (ITP), systemic lupus erythematosus, antiphospholipid antibody syndrome, hypersplenism, disseminated intravascular coagulation, thrombotic thrombocytopenic purpura, hemolytic uremic syndrome, congenital thrombocytopenia, or medication effect [22]. Most of these conditions can be ruled out by history, physical exam, and exclusion of underlying diagnoses.

The most difficult differential usually comes down to gestational thrombocytopenia versus ITP. These conditions cannot be reliably differentiated with antiplatelet antibody testing or any other diagnostic test. Differentiation can be obtained by documenting platelet counts less than 70,000/μL, which suggests ITP, or by documenting a return to normal platelet counts after delivery, suggestive of gestational thrombocytopenia [22]. To be classified as gestational thrombocytopenia, there are several conditions that must be satisfied. Gestational thrombocytopenia is mild, with platelet counts greater than 70,000/μL. There is no history of significant bleeding and no history of thrombocytopenia prior to pregnancy. Platelet counts generally return to normal within 2–12 weeks following delivery, and there is an extremely low risk of fetal or neonatal thrombocytopenia. Many women with ITP have a history of abnormal bleeding prior to pregnancy, although this is not universal. Findings suggestive of ITP include persistent platelet counts less than 100,000/μL, normal or increased megakaryocytes in the bone marrow, absence of splenomegaly, and exclusion of other systemic disorders known to be associated with thrombocytopenia [22].

Women with gestational thrombocytopenia are not at risk for maternal or fetal hemorrhage or bleeding complications [22]. Immunologic thrombocytopenia such as ITP and neonatal alloimmune thrombocytopenia (NAIT) have the potential for fetal complications. Both are characterized by increased platelet destruction. Twelve to fifteen percent of neonates born to mothers with ITP may develop platelet counts less than 50,000/μL. This may result in findings such as purpura, ecchymosis, or melena. Less commonly, fetal intracranial hemorrhage may develop, unrelated to mode of delivery. The incidence of serious bleeding complications in neonates of women with ITP is estimated at 3% and the rate of intracranial hemorrhage at 1% [22].

Onset of thrombocytopenia during the third trimester should prompt consideration of gestational hypertensive disorders, which are associated with about 20% of maternal thrombocytopenia, and decreasing platelet count is considered a sign of worsening of disorders of this spectrum (see also Chapter 1 in *Maternal-Fetal Evidence Based Guidelines*). When combined with hemolytic anemia and elevated liver tests, thrombocytopenia is indicative of the diagnosis of hemolysis, elevated liver enzymes, and low platelet count (HELLP) syndrome. These disorders are associated with an increase in platelet destruction but the underlying physiology is not known. Platelet function may be reduced even if platelet counts are normal, and thrombocytopenia may occur prior to other manifestations of gestational hypertension. Hemorrhage is uncommon in the absence of disseminated intravascular coagulation. Neonatal thrombocytopenia following gestational hypertension is increased in premature infants but not in term infants [22].

Pregnancy is associated with significant alterations in the coagulation system. There is a decrease in protein S levels as well as in coagulation factors XI and XIII. There is an increase in coagulation factors I, VII, VIII, IX, and X. D-dimer and fibrinogen levels also increase. The overall result is the creation of a hypercoagulable state that is exacerbated by venous stasis and compression of the inferior vena cava and pelvic veins by the enlarging uterus [3, 23]. This places the pregnant women at increased risk for such phenomena as deep venous
thrombosis and pulmonary embolism, which are further elevated when pregnancy is associated with other high-risk states such as obesity, prolonged immobility (bed rest), or surgery (see also Chapter 28 in Maternal-Fetal Evidence Based Guidelines). Consideration of these risks is important in the evaluation of the pregnant patient with unilateral lower extremity edema or acute dyspnea. They also warrant caution and the responsibility to practice evidence-based medicine rather than erroneously recommending “bed rest” for treatment of conditions ranging from threatened abortion to preterm contractions to gestational hypertension.

GASTROINTESTINAL

Changes in gastrointestinal physiology lead to some of the most commonly described discomforts of pregnancy ranging from nausea and vomiting in early pregnancy to more persistent symptoms of gastroesophageal reflux and constipation. However, gastrointestinal symptoms may reflect coexisting diseases or may even herald life-threatening complications of pregnancy such as severe preeclampsia and HELLP syndrome or acute fatty liver of pregnancy (AFLP). Once again, it becomes crucial for the obstetric care provider to be skilled in recognizing signs and symptoms that result from normal pregnancy physiology and distinguishing those of more serious conditions.

There are several recognizable effects of pregnancy on gastrointestinal function. Gastric secretion acidity declines but volume increases and relaxation of the cardiac sphincter leads to greater esophageal reflux [1]. Combined with a decrease in gastric and intestinal motility, this leads to the “full stomach” effect that puts pregnant women at increased risk of aspiration. This, combined with increased airway edema, increases the risks of general endotracheal anesthesia during pregnancy. Thus, for anesthesia purposes, all pregnant women are considered to have a full stomach. Precautions to reduce aspiration risk include the use of non particulate oral antacids prior to induction of anesthesia, the use of rapid sequence induction methods, and the use of a cuffed endotracheal tube [1].

During pregnancy there is a decrease in colonic motility and an increase in water absorption, leading to constipation as a common complaint among pregnant women. A prospective study of constipation in pregnancy found that one in two women reports constipation at some point in pregnancy, with rates of 24%, 26%, 16%, and 24% in first, second, third trimesters, and postpartum, respectively. Constipation is more likely in women with a prior history of constipation and in women taking iron supplements. The study did not include nonpregnant controls, but historic controls indicate a constipation rate of 7% in a similar age group [24].

The most common gastrointestinal symptom of pregnancy is nausea and vomiting (see also Chapter 9 in Maternal-Fetal Evidence Based Guidelines). As many as 80% of women report nausea during pregnancy and it is also the most common reason for hospitalization in the first trimester [19,25]. Nausea is generally considered a normal symptom of early pregnancy, and morning sickness has been associated with improved pregnancy outcomes such as reduced risk of miscarriage, preterm birth, low birth weight, and perinatal death. This is theorized to be due to a placental etiology for nausea and vomiting, which is increased by early development of a healthy and robust placenta [19]. However, the exact etiology of this symptomatology is not known. It has been hypothesized that nausea and restricted intake in the mother create an environment that is favorable for early placental development [26] or that it confers an evolutionary advantage by causing the mother to avoid the ingestion of foods that may be dangerous to the developing fetus [19]. Numerous psychological theories have also been proposed to explain the phenomenon of nausea and vomiting in pregnancy. Nausea in pregnancy is commonly attributed to hCG levels, but conclusive evidence of the underlying physiology is lacking. Experience of nausea is also correlated with elevated estradiol levels and inversely correlated with prolactin levels [25]. Estrogens in oral contraceptive pills have shown a dose-related effect of nausea and vomiting. Smoking decreases both hCG and estrogen, and a reduced rate of nausea and vomiting of pregnancy has been demonstrated in smokers [19]. Increased placental mass, as found in multiple gestations and gestational trophoblastic disease, has been found to increase the risk of nausea and vomiting and of hyperemesis gravidarum.

It is important for nausea and vomiting of pregnancy to be distinguished from that resulting from other pathologic conditions. Complacency in the evaluation of pregnant patients with nausea and vomiting may result in undertreatment of distressing symptoms, development of hyperemesis gravidarum, or failure to diagnose a coexisting underlying disease. Nausea and vomiting of pregnancy typically start before 9 weeks’ gestation and are not accompanied by fever, abdominal pain, or headache [19]. Deviation from this presentation should prompt evaluation for other etiologies. The differential diagnosis includes gastrointestinal disorders such as gastroenteritis, gastroparesis, achalasia, biliary tract disease, hepatitis, intestinal obstruction, peptic ulcer disease, pancreatitis, and appendicitis. Genitourinary conditions that may cause nausea and vomiting include pyelonephritis, uremia, ovarian torsion, kidney stones, and degenerating myoma. Nausea and vomiting may also be due to metabolic disorders such as diabetic ketoacidosis, porphyria, Addison’s disease, and hyperthyroidism or neurologic disorders such as pseudotumor cerebri, vestibular lesions, migraines, or central nervous system tumors. Finally, drug-related toxicity, psychologic factors, or pregnancy-related complications such as preeclampsia and AFLP may present with nausea and vomiting [19].

Patients who are taking a multivitamin at the time of conception have reduced rates of nausea and vomiting [21]. There is good evidence to support the use of vitamin B6 alone or combined with doxylamine for the treatment of nausea and vomiting in pregnancy. Ginger supplements have also been shown to reduce severity of nausea and vomiting [19,27]. Numerous antiemetics have also shown acceptable safety and efficacy against nausea and vomiting. Hospitalization, intravenous fluids, and enteral nutrition may be used in rare cases of continued weight loss in spite of these therapies (see also Chapter 9 in Maternal-Fetal Evidence Based Guidelines).

RENAL SYSTEM AND HOMEOSTATIC

Pregnancy-related changes in the urinary tract include dilatation of calyces, pelvis, and ureters. Ureteral dilatation may be noted as early as the first trimester and is present in 90% of gravidas by term. Obstructive and humoral mechanisms have been proposed for this dilatation, with obstruction by the gravid uterus and ovarian venous plexus likely causing the dilatation above the pelvic brim [28]. Dextrorotation of the gravid uterus, likely due to the sigmoid colon on the left and
posterior to the uterus, causes the mechanical obstruction at the pelvic brim on the right more than the left. One important adverse consequence of this ureterocolic dilatation is the increased incidence of pyelonephritis among gravidas with asymptomatic bacteriuria.

There are significant increases in renal blood flow and in glomerular filtration rate (GFR) in pregnancy [1]. Indeed, GFR increases up to 50% higher than in the nonpregnant state. As a result, serum urea and creatinine levels decline in pregnancy [29]. This can have significant effects on renal clearance of vitamins and pharmaceutical agents. There is a lowering of the threshold for glucose excretion, which may result in significant random glucosuria even in the absence of gestational diabetes. This glucosuria may also contribute to the increased susceptibility of pregnant women to urinary tract infections.

There is also a marked increase in ureteral pressure in the third trimester while standing or sitting that is decreased when in the lateral recumbent position [28]. This has implications for collection of 24-hour urine samples, as retention of urine in the dilated collecting system may result in an incomplete sample. This may be alleviated by instructing the patient to lie in the lateral recumbent position for about 45 minutes before the discard void prior to starting the collection and again before the final void of the sample [28]. Proteinuria is considered abnormal if excessive of 300 mg/24 hours in pregnant patients [28].

Relaxin and nitric oxide (NO) have been implicated as key factors in mediating the renal vasodilatation and glomerular hyperfiltration that is characteristic of normal human pregnancy. In vivo, relaxin administration to male and non-pregnant female rats produced physiologic changes that mimicked normal pregnancy, with decrements in SVR along with significant increases in effective renal plasma flow (RPF) and, hence, GFR [29].

Pregnancy is associated with altered tubular function and therefore altered reabsorption of protein, glucose, amino acids, and uric acid. In contrast to tubular function, our knowledge of the factors that govern gestational changes in serum electrolytes is somewhat more definitive. Total body sodium increases on an average by 3–4 mEq/d, ultimately producing a net balance of 900–1000 mEq, and total body potassium also increases by up to 320 mEq by the end of gestation [29]. Despite the net increase in body stores of sodium and potassium Table 3.1, serum levels of both electrolytes decrease during pregnancy Table 3.2. Therefore, pregnancy is characterized by increments in total body electrolyte stores, albeit with decrements in serum levels. Clinicians must recognize that increments in serum electrolytes that still fall within the range of normal may constitute meaningful aberrations in electrolyte balance. Furthermore, conditions prone to either electrolyte retention or loss may be exacerbated during pregnancy [29].

Absorption is affected by changes in gastric pH, gastric emptying, and small intestine motility. Increased cardiac output and heightened blood flow to the stomach and small intestine can increase absorption [30]. Changes in plasma volume, body fat, and total body water affect drug distribution; the proportion of unbound fraction of certain drugs may increase due to a decrease in protein concentration. Metabolism is affected by upregulation or downregulation of various enzymes. For example, liver cytochrome p-450 activities increase while extrahepatic cholinesterase activity decreases [30]. Drug excretion is affected by the increased renal blood flow and elevated GFR of pregnancy as well as by increased respiratory elimination [30]. This has important implications for both the maintenance of therapeutic drug levels and the avoidance of toxicity. Clinical evidence to guide pharmacologic therapy in pregnancy is limited, in part due to the frequent elimination of women of reproductive age from pharmacokinetic trials. A review of the National Library of Medicine database shows that there are a very limited quantity of pharmacokinetic data for pregnancy, and thus evidence-based recommendations for dosing and scheduling of drugs during pregnancy are sparse [30].

REFERENCES


PHARMACOKINETICS

It may be helpful to conclude with a final clinical topic that illustrates many of the previously described physiologic changes: that of pharmacokinetics. The four major events involved in pharmacokinetics (absorption, distribution, metabolism, and excretion) are potentially altered by physiologic change of pregnancy in a number of organ systems.

46 OBSTETRIC EVIDENCE BASED GUIDELINES
KEY POINTS
- There is no evidence that ultrasound examination during pregnancy is harmful. Prenatal exposure to ultrasound is not associated with adverse influence on school performance, physical or neurological function.
- Ultrasound should be performed by trained and experienced professionals, with continuing education and ongoing quality-monitoring programs.
- Routine use of ultrasound in pregnancy increases early detection of multiple pregnancies, and major fetal anomalies.
- Ultrasound examination is the best method to estimate accurate gestational dating in pregnancy.
- Ultrasound dating in the first trimester is most accurate for gestational age assessment. Ultrasound examination at first prenatal visit (usually first trimester) versus at 18–20 weeks provides more precise estimate of gestational age, and may be associated with less maternal worry. First trimester ultrasound allows earlier detection of multiple pregnancies, screening for Down’s syndrome with nuchal translucency, and diagnosis of nonviable pregnancies.
- All pregnant women should be offered a second trimester ultrasound for optimal anatomy evaluation. If only one ultrasound will be done in pregnancy, it should be done in the second trimester at about 18–22 weeks.
- In low-risk or unselected populations, routine third trimester (≥24 weeks) pregnancy ultrasound has not been associated with improvements in perinatal mortality. Routine use of ultrasound in the third trimester is associated with higher detection of small-for-gestational-age babies.
- In low-risk or unselected populations, routine Doppler ultrasound examination in the third trimester does not result in reduced perinatal mortality.
- In high-risk pregnancies with fetal growth restriction, umbilical artery Doppler assessment is associated with a reduction in perinatal deaths and obstetric interventions.
- Measurement of cervical length (CL) by transvaginal ultrasound (TVU) has been shown to be an effective predictor of preterm birth (PTB). When a short CL is detected before 24 weeks, interventions such as vaginal progesterone in singletons without prior PTB (using TVU CL ≤ 20 mm), and cerclage in singletons with prior PTB (using TVU CL ≤ 25 mm), have been associated with decrease in PTB and perinatal morbidity and mortality.

SAFETY OF ULTRASONOGRAPHY
The main concern about the safety of ultrasound is about tissue temperature elevation from energy transfer and its possible effect of cavitations, or the formation of microbubbles in the tissues exposed to ultrasound waves. The effect of ultrasound on tissues has been studied with animal experimentation and has suggested an adverse effect. In humans, however, the information comes from epidemiological data and population studies. No epidemiologic studies have shown harmful effects in humans.

There is no consistent evidence that ultrasound examination during pregnancy is harmful. Studies have shown that prenatal exposure to ultrasound is not associated with adverse effects on children’s physical or cognitive development [1]. Even multiple ultrasound exams during pregnancy were not shown to adversely affect speech, language, behavior or neurologic function on postnatal follow up at 8 years of age [2]. In a randomized trial comparing those receiving a second trimester ultrasound to those who remained unexposed, there was no significant difference in school performance up to age 15–16 [3]. Overall, ultrasound in pregnancy is not associated with adverse maternal or perinatal outcome, however, there may be a weak association between exposure to ultrasound and nonright handedness in boys [4].

Despite the lack of evidence suggesting harmful effect, ultrasound is a form of energy and may produce secondary effects in the tissues it traverses. Obstetrical ultrasound in pregnancy should be considered a medical procedure for the evaluation of the fetus and maternal pelvic organs. Current expert consensus recommends that ultrasound be only performed with valid medical indications, with the shortest duration possible and at the lowest settings. Adhering to the as low as reasonably achievable (ALARA) principle helps to avoid unnecessary exposure to ultrasonic waves. Sonographers and sonologists should familiarize themselves with the mechanical and thermal indices during ultrasound examinations. Exposing the fetus to ultrasonography with no anticipation of medical benefit is thus not justified [5,6].

QUALITY
Levels of expertise vary between different health care centers. Since ultrasound efficiency is operator dependent, continuing education and ongoing quality-monitoring programs are important strategies in each center offering ultrasound diagnosis. The ongoing risks of “false-negative” tests and/or misinterpretation of the images obtained (either false-positives, or wrong diagnoses) can be minimized if those examinations are carried out and interpreted by trained and experienced professionals. Sensitivity of ultrasound screening for pregnancy varies widely. Appropriate ultrasound laboratory accreditation, certification of staff, documentation of findings, and continuous careful quality control are important components of ultrasound competency [5,6].

INFORMED CONSENT AND PATIENTS’ EXPECTATIONS
Even though a formal written informed consent is not always needed before the examination, every pregnant woman should be informed on expectations about the obstetric
ultrasound, as well as its benefits and risks. The patients should know that ultrasound evaluation is a screening test with wide variations in detection rates for fetal anomalies, and that all ultrasound diagnoses, especially false-positive and false-negative ones, can put both mother and fetus at risk.

Whether the sex of the fetus should be revealed to the patient with a singleton gestation should be addressed. It may be harmful for the physician–patient relationship to withhold this information, especially if the patient previously requested it. Although a moral conflict may exist in some cultures around the world where this information is used by the patient for voluntary abortions based on sex selection and sex preferences [7], in general disclosing fetal gender during ultrasound can benefit not only the doctor–patient relation but also parent–child relationship [8].

During high-feedback ultrasound scans, women can see the screen and they receive detailed explanations of the images. In low-feedback ultrasound scans, only the operator can see the screen and the women are told the results at the end of the scan. Compared with low-feedback ultrasound, women who had high-feedback fetal ultrasound are significantly more likely to stop smoking and avoid alcohol during pregnancy, with a trend favoring the women’s increased use of positive adjectives to describe their feelings after the ultrasound [9].

**ROUTINE VERSUS SELECTIVE USE OF ULTRASOUND**

**Routine** (i.e., performed on every pregnant woman) ultrasound examination is associated with the following, compared with **selective** ultrasound examination (i.e., performed only on women with specific indications) [1]:

1. Increases the early detection of multiple pregnancies.
2. Increases detection of major fetal anomalies.
3. Reduces the incidence of late-term and post-term pregnancies and rates of induction of labor for late-term pregnancy by allowing a more precise estimation of gestational age.
4. No significant differences are detected for clinical outcomes such as perinatal mortality. The effect of ultrasound on perinatal mortality is dependent on the detection rate of fetal malformations and on the uptake of pregnancy termination for anomalies in the population at study.

If one routine ultrasound examination is done, it is usually performed at 18–22 weeks (<24 weeks). Earlier examination provides more accurate assessment of gestational age; later examination (e.g., 20–22 weeks) allows more complete inspection of fetal anatomy. In the obese population, transabdominal ultrasound screening may have better completion rates if delayed by 2 weeks (20–22 weeks gestation) [10] and TVU at 12–16 weeks may offer better fetal anatomy screening (see later in this chapter).

**GESTATIONAL AGE DATING IN PREGNANCY**

Precise estimation of gestational age is extremely important for optimal obstetric care, including evaluation of fetal growth, interpretation of maternal screening markers, choosing the appropriate gestational age to perform interventions, and management of preterm and late-term pregnancies.

For gestational age estimation, cardinal numbers should be preferred to ordinal numbers to avoid confusion. So week 1 is 1–7 days after LMP, week 2 is 8–14 days, etc. In clinical practice, gestational weeks are used to estimate dating, and not months. If a lay person asks “How many months am I?,” then 6 weeks of gestation can be equated approximately to 1 month, etc., and 38 weeks = 9 months. **Definitions of gestational age, while not uniformly accepted, are shown in Table 4.1** [10].

Ultrasound examination is the best method to determine gestational age and estimated due date (EDD) [10]. The first day of the last menstrual period (LMP) should be asked of all pregnant women to determine when the dating ultrasound should be performed. Compared with LMP, ultrasound-based gestational age is more precise due to errors in patient recall, and variations in cycle length and timing of ovulation [10,11]. The error, even with certain LMP, is due often to late ovulation (>14 days after LMP). Some have stated that there is no reason to use LMP for dating when adequate ultrasound data is available by 24 weeks [10,11].

Ultrasound-based gestational age estimates are lower than LMP-based gestational age estimates, and generate a higher rate of PTB and lower rate of post-term birth. The Naegle rule (add 7 days to first day of LMP, add 1 year, take back 3 months), manual assessment of uterine size, quickening, etc., should not be used unless ultrasound dating is unavailable.

In general, the earlier the ultrasound, the more accurate the dating. Multiple parameters and equations have been evaluated to estimate gestational age. The crown-rump length is associated with the most accurate estimation up to and including 13 6/7 weeks of gestation with an accuracy of ±2–7 days. For pregnancies in the second trimester, beyond 14 weeks gestation, the head circumference (HC) or biparietal diameter (BPD) appear to be the best single-measurement predictor of gestational dating. BPD is most accurate for dating early, between 12 and 14 weeks. Combining three or more parameters improved dating slightly over single biometric parameter [12]. A combination of BPD, HC, abdominal circumference (AC), and femur length (FL) is commonly used for dating by ultrasound in the second and third trimesters [13]. Repeated examinations improve the prediction only marginally, and the EDD should always be set by the earliest ultrasound due to a smaller prediction error. While prediction of gestational age by ultrasound can be very accurate, prediction of date of delivery remains less accurate, with an error of usually ≥7–8 days, given other biologic factors.

Other parameters that may play a role in estimating gestational age include trans-cerebellar diameter (TCD) and long bone measurements. The TCD is an accurate predictor of gestational age, and can be used between 14 and 28 weeks reliably with the use of nomograms. There is some reliability in gestational age prediction even up to 35 weeks, and TCD is spared effects from intrauterine growth restriction, so can be used to assess pregnancies at risk for this complication [14]. The presence of epiphyseal lysis in the lower extremities usually signifies a gestational age (GA) of >32 weeks [15].

**Table 4.1 Definition of Gestational Age Periods in Pregnancy**

<table>
<thead>
<tr>
<th>Period</th>
<th>Gestational age (weeks)</th>
</tr>
</thead>
<tbody>
<tr>
<td>First trimester</td>
<td>0–13 6/7</td>
</tr>
<tr>
<td>Second trimester</td>
<td>14 0/7–27 6/7</td>
</tr>
<tr>
<td>Third trimester</td>
<td>28 0/7–36 6/7</td>
</tr>
<tr>
<td>Preterm</td>
<td>28 0/7 to delivery</td>
</tr>
<tr>
<td>Late preterm</td>
<td>34 0/7–36 6/7</td>
</tr>
<tr>
<td>Term</td>
<td>37 0/7–41 6/7</td>
</tr>
<tr>
<td>Early term</td>
<td>37 0/7–38 6/7</td>
</tr>
<tr>
<td>Full-term</td>
<td>39 0/7–40 6/7</td>
</tr>
<tr>
<td>Late-term</td>
<td>41 0/7–41 6/7</td>
</tr>
<tr>
<td>Post-term</td>
<td>≥42 0/7</td>
</tr>
</tbody>
</table>
The American College of Obstetricians and Gynecologists (ACOG) have published specific guidelines to guide amendment of dating when ultrasound and LMP are discrepant. Ultrasound dating is adopted when discrepancy is noted. Knowledge about LMP, regularity of cycles, OCP use and unusual bleeding are clinically helpful, but imprecise regarding dating. Ultrasound dating is best, and often corrects dating by even a “certain” LMP [16]. Table 4.2 shows dating criteria based on ultrasound results.

### ULTRASOUND EXAMINATIONS BY TRIMESTER

#### First Trimester

Ultrasonographic evaluation in the first trimester (0–13 6/7 weeks) is the most accurate method to determine exact gestational age, as discussed above. Ultrasound examination at the first prenatal visit versus at 18–20 weeks provides more precise estimate of gestational age, and may be associated with less maternal worry. First trimester ultrasound also allows earlier detection of multiple pregnancies, nonviable pregnancies, certain fetal anomalies, and screening for Down’s syndrome and other aneuploidy with nuchal translucency (NT) [1,6,17].

No other important maternal or perinatal outcome differences are detected, with insufficient data to accurately assess some rare outcomes such as perinatal mortality. Current guidelines do not recommend the routine use of ultrasound in the first trimester in the absence of indications [10], but several experts advocate its routine use for the benefits listed above.

Transvaginal scanning is preferred for dating early in the first trimester. It is also useful in cases of a pregnancy resulting from ovulation induction or other assisted reproductive technologies, first trimester bleeding, or increased risk of aneuploidy, and should be used if transabdominal examination is inconclusive for diagnosis. First trimester screening for congenital defects by TVU is an option for pregnant women who meet certain criteria, such a very high risk for congenital anomalies (e.g., very elevated hemoglobin A1c) and who may elect termination based on ultrasound results.

The NT is a physiologic fluid-filled space at the back of the fetal neck measured for aneuploidy screening between weeks 10 6/7 and 13 6/7 of gestation. Increase in NT has been associated with chromosomal and anatomic abnormalities in the fetus. First trimester screening for aneuploidy between 10 6/7 and 13 6/7 weeks (or crown–rump length or CRL 45–84 mm) combines NT measurement with maternal serum markers to provide individualized risk assessment. Early risk determination permits choosing the most appropriate definitive diagnostic procedures like chorionic villus sampling, allowing women to prepare for a child with health problems and also providing the option of earlier pregnancy termination [6] (see Chapter 5).

Indications by ACOG and American Institute of Ultrasound in Medicine (AIUM) for first trimester ultrasound are shown in Table 4.3 and essential elements of first trimester ultrasound in Table 4.4 [5,6].

#### Ultrasound Diagnosis of Anembryonic Pregnancy or Embryonic Demise

Diagnostic criteria for the diagnosis of nonviable pregnancy in the first trimester is a subject that has undergone recent revision. Balancing risk of harming a viable intrauterine pregnancy must be balanced with intervention for a nonviable one. Some criteria are routinely used (Table 4.5), usually based on TVU: absence of cardiac activity with an embryo of certain length, absence of embryo with gestational sac of a certain size, and absence of embryo by a certain time in pregnancy [20].

A gestational sac is normally noted within the uterus by 5 weeks of gestation. Shortly thereafter at approximately 5 1/2 weeks in a normal pregnancy, the yolk sac appears, followed by the fetal pole at 6 weeks. When measuring the gestational sac, dimensions are recorded in three orthogonal planes; the average of the measurements is the mean sac diameter. **A mean gestational sac diameter of ≥25 mm without an embryo is diagnostic of pregnancy failure (e.g., anembryonic pregnancy, or blighted ovum) with positive predictive value approaching 100%** [21]. An intrauterine gestational sac should be visible by TVU with a serum beta-human chorionic gonadotropin (B-HCG) of >1500 mIU/mL. If this is not the case, ectopic pregnancy should be suspected.

Fetal cardiac activity is usually seen once the fetal pole is visible around 6 weeks of gestation. A CRL cutoff of 5 mm without cardiac activity was previously used to diagnose nonviable pregnancy, but literature review showed that there have been pregnancies that met this criterion that went on to be viable [21,22]. Interobserver variability in measurements can also lead to inaccurate diagnosis of failed pregnancy. Adopting a **CRL cutoff of 7 mm with no visible cardiac activity** brings the specificity of this finding for diagnosing failed pregnancy (e.g., embryonic demise, or missed abortion) close to 100% [20].

Since the presence of intrauterine structures in a viable pregnancy appear in a predictable sequence at standard time intervals [23], aberrations in this sequence can indicate abnormal pregnancy. Abnormal pregnancy should thus be suspected in the absence of an embryo with a heartbeat ≥14 days.
**Table 4.3** Indications for First Trimester Ultrasound

- To confirm the presence of an intrauterine pregnancy
- To evaluate suspected ectopic pregnancy
- To define the cause of vaginal bleeding
- To evaluate pelvic pain
- To estimate gestational age
- To diagnose or evaluate multiple gestations
- To confirm cardiac activity and identify nonviable pregnancies
- As an adjunct to chorionic villus sampling, embryo transfer, and localization and removal of an intrauterine device
- To evaluate maternal pelvic masses and/or uterine anomalies
- To evaluate suspected hydatidiform mole
- To screen for certain anomalies such as anencephaly in patients at high risk
- To measure NT when part of a screening program for fetal aneuploidy

**Table 4.4** Essential Elements of First Trimester Ultrasound

- Gestational sac (location, mean diameter).
- Yolk sac (diameter).
- CRL of embryo.  
- Development of fetal anatomy in early pregnancy.
- Fetal viability (cardiac activity should be seen in embryo >7 mm).
- Fetal number (amnioncystic and chorionicity has to be reported for multiples).
- Ultrasound features of early pregnancy failure, e.g., ectopic pregnancy, hydatidiform mole.
- Uterus, adnexa, cervix, and cul de sac.
- If possible, the appearance of the nuchal region should be assessed, and specific measurement of Nuchal translucency (NT) measured as part of desired screening.
- Any other abnormalities (e.g., leiomyomata).

**Table 4.5** Criteria for Diagnosis of Embryonic or Anembryonic Demise

<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>Criteria</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anembryonic pregnancy</td>
<td>Mean gestational sac diameter ≥ 25 mm without an embryo</td>
</tr>
<tr>
<td>Anembryonic demise</td>
<td>CRL ≥7 mm with no visible cardiac activity</td>
</tr>
<tr>
<td></td>
<td>No embryo with cardiac activity</td>
</tr>
<tr>
<td></td>
<td>≥14 days after gestational sac without yolk sac</td>
</tr>
<tr>
<td></td>
<td>or</td>
</tr>
<tr>
<td></td>
<td>≥11 days after gestational sac with yolk sac</td>
</tr>
</tbody>
</table>

After seeing a gestational sac without a yolk sac, or ≥11 days after presence of a gestational sac with a yolk sac [20]. The presence of normal embryonic cardiac activity in the uterine cavity in the first trimester has a >90% prediction for a live birth in both symptomatic and asymptomatic pregnancies.

**Precautions and Pitfalls**

Physiologic midgut herniation is normal at 7–11 weeks; it resolves ≥12 weeks; do not confuse with omphalocele. The rhomboencephalon can appear as a cystic mass up until 8–10 weeks, and should not be confused with a central nervous system (CNS) anomaly; ventriculomegaly cannot be assessed well in the first trimester. The amnion and chorion are expected to be fused by 14 weeks.

**Second Trimester (aka “Anatomy,” “Morphology,” or “Standard” Ultrasound)**

If just a single exam is to be done during pregnancy, the best timing for such ultrasound screening of the fetal anatomy and dating is the early to mid-second trimester (18–22 weeks) [1], to obtain an accurate estimation of gestational age and a satisfactory inspection of the fetal anatomy. This is therefore usually called the anatomy, morphology, or standard ultrasound examination (nomenclature such as Levels I, II, etc. ultrasound is controversial and less descriptive). All pregnant women should be offered a second trimester ultrasound, whether or not they have had a first trimester ultrasound. The estimated sensitivity to detect fetal anomalies varies widely, with higher rates of detection for major anomalies than for minor anomalies, and some organs (e.g., neural tube defect) versus others (e.g., heart) [19].

The detection of major fetal anomalies is often reported at about 50%–70% even in the best centers [18]. The detection of fetal cardiac malformations is particularly poor, with often not more than 20%–30% of major heart malformations detected [24].

A “detailed” ultrasound can be performed with the aim to detect anomalies or markers associated with fetal aneuploidy [25].

While the best time for detection of most malformations is around 20–24 weeks it is important for women to obtain this information as early as possible. In some circumstances more than one second trimester ultrasound is necessary, especially if the first second trimester ultrasound is performed at 18–19 weeks. Experts have suggested essential elements for second trimester ultrasound (Table 4.6) [5,6].

There can be other types of follow-up ultrasounds. Limited ultrasound examination is performed when a specific question requires investigation. Examples include assessment of amniotic fluid volume, fetal viability, biophysical profile, to guide amniocentesis, to localize the placenta in antepartum bleeding, or to evaluate fetal position. A limited ultrasound is only appropriate if a prior complete standard ultrasound examination has been done. A detailed ultrasound examination should be considered for a patient who, by history, clinical evaluation, or prior scanning evaluation, is at an increased risk for a fetal anatomic or physiological abnormality. This ultrasound examination must be done by personnel with expertise in obstetric ultrasonography, and maternal and fetal diseases [25].

Other specialized examinations include but are not limited to fetal Doppler studies, biophysical profile, CL, three-dimensional (3D) imaging, and fetal echocardiography (see the section Fetal Echocardiography).

**Third Trimester**

The potential benefit of a third trimester ultrasound examination greatly depends on the quality of prior ultrasounds and maternal indications. If the first and only ultrasound is in the third trimester, it probably has similar benefits to the routine second trimester ultrasound, with the exception of accurate dating and early anomaly detection of the latter. Ultrasound evaluations in the third trimester generally involve assessments of fetal growth,
lished nomograms. Consecutive ultrasound examinations for growth evaluation should typically be performed no less than 3 weeks apart.

of internal os is found.

previa are suspected early in gestation, a TVU is indicated, with verification in the third trimester by a TVU if previa or placental edge within 20 mm

c First trimester CRL measurement is the most accurate means for sonographic dating.

b The apparent position early in pregnancy may not correlate well with its location at the time of delivery. Therefore, if low-lying placenta or placenta

DC, 2009.

American Institute of Ultrasound in Medicine,

Sources:

• Evaluation of the maternal uterus and adnexal structures should be performed.

• Fetal weight estimation can be calculated by obtaining measurements, such as BPD, HC, AC, and FL. d

• TVU may be offered for detection of short CL (see Chapter 17).

• Gestational age assessment. e

• Fetal weight estimation can be calculated by obtaining measurements, such as BPD, HC, AC, and FL. d

• Evaluation of the maternal uterus and adnexal structures should be performed.

• Fetal anatomy survey: Fetal anatomy is best assessed by ultrasound ≥18 weeks. Essential elements of a standard examination:

  • Head and neck: Cerebellum, choroids plexus, cisterna magna, lateral cerebral ventricles, midline falx, cavum septi pellucidi, upper lip.
  • Chest: The basic cardiac inspection includes a four-chamber view of fetal heart with visualization of the right and left ventricular outflow tracts.
  • Abdomen: stomach (presence, size, situs), kidneys, bladder, umbilical cord (insertion site into fetal abdomen and vessel number).
  • Spine: Cervical, thoracic, lumbar, and sacral spine.
  • Extremities: Legs and arms (presence or absence).
  • Gender: For evaluation of multiple gestations and when medically indicated.

Table 4.6 Essential Elements for Second Trimester Ultrasound d

- Fetal cardiac activity (abnormal heart rate or rhythm should be documented).
- Fetal number (multiple pregnancies require additional information: chorionicity, amnionycity, comparison of fetal sizes, estimation of amniotic fluid volume at each side of the membranes, and fetal gender).
- Presentation.
- A qualitative or semi-quantitative assessment of amniotic fluid (e.g., amniotic fluid index, single deepest pocket, 2-diameter pocket) (see Chapter 21 in Maternal Fetal Evidence Based Guidelines).
- The placental location, appearance and relationship to the internal cervical os should be recorded. 
- The umbilical cord should be imaged to confirm number of vessels. Placental cord insertion site should be documented when technically feasible.
- TVU may be offered for detection of short CL (see Chapter 17).
- Gestational age assessment. e
- Fetal weight estimation can be calculated by obtaining measurements, such as BPD, HC, AC, and FL. d
- Evaluation of the maternal uterus and adnexal structures should be performed.
- Fetal anatomy survey: Fetal anatomy is best assessed by ultrasound ≥18 weeks. Essential elements of a standard examination:
  - Head and neck: Cerebellum, choroids plexus, cisterna magna, lateral cerebral ventricles, midline falx, cavum septi pellucidi, upper lip.
  - Chest: The basic cardiac inspection includes a four-chamber view of fetal heart with visualization of the right and left ventricular outflow tracts.
  - Abdomen: stomach (presence, size, situs), kidneys, bladder, umbilical cord (insertion site into fetal abdomen and vessel number).
  - Spine: Cervical, thoracic, lumbar, and sacral spine.
  - Extremities: Legs and arms (presence or absence).
  - Gender: For evaluation of multiple gestations and when medically indicated.


d If not performed in second trimester, a third trimester ultrasound is indicated.

e The apparent position early in pregnancy may not correlate well with its location at the time of delivery. Therefore, if low-lying placenta or placenta previa are suspected early in gestation, a TVU is indicated, with verification in the third trimester by a TVU if previa or placental edge within 20 mm of internal os is found.

f First trimester CRL measurement is the most accurate means for sonographic dating.

None of the several equations for estimating fetal weight based on such fetal biometric measurements is superior to others; ideally, the equations should be derived by actual fetal weights of the local or institutional population. Results can be compared with fetal weight percentiles from published nomograms. Consecutive ultrasound examinations for growth evaluation should typically be performed no less than 3 weeks apart.

amniotic fluid volume, evaluation of the placenta, and evaluation of fetal wellbeing (possibly biophysical profile or Doppler studies). Examples of possible indications for third trimester ultrasound based on maternal and fetal risk factors are shown in Table 4.7.

In low-risk women, ultrasound examinations at 30–32 weeks and at 36–37 weeks significantly decrease the likelihood of newborns with growth restriction, though they do increase the rate of antenatal intervention. This randomized controlled trial included 1998 women, and investigators calculate over 30,000 women are required for a trial to show a significant decrease in neonatal mortality [26]. In a meta-analysis, there was no difference in antenatal, obstetric and neonatal intervention in women screened with >24 weeks (late) ultrasound versus those not screened. There was a slightly higher caesarean section rate in women screened with late ultrasound, but this difference did not reach statistical significance. Routine late pregnancy ultrasound was not associated with improvements in overall perinatal mortality [27]. In a recent RCT, performing a growth ultrasound at 36 versus 32 weeks is more sensitive (61% vs. 32%) in detecting severe FGR, but not associated with significant differences in perinatal outcomes [28]. Routine screening for fetal growth restriction in the third trimester has been investigated also in a large prospective cohort study and may increase detection of fetuses that will go on to be small for gestational age infants, to 57% in routine screening from 20% in selected screening [29]. Currently there is insufficient data to recommend routine screening for growth restriction in the third trimester without indication, but many experts advocate its routine use based on the data just described. There is insufficient data about the potential psychological effects of routine ultrasound in late pregnancy, and limited data about its effects on both short- and long-term neonatal and childhood outcome.

Placental grading as an adjunct to third trimester ultrasound examination was associated with a significant reduction in the stillbirth rate in one trial [30]. In one study 15,122 patients were evaluated for Grannum grade III placental calcifications prior to 28 weeks of gestation. Grade III placental appearance prior to term was independently associated with increased risk of stillbirth after controlling for tobacco use [31]. More research is needed in placental grading before recommendation can be made for its routine use for prediction of poor perinatal outcome.

DOPPLER Umbilical Artery

In low-risk or unselected populations, routine Doppler ultrasound examination, usually of the umbilical artery and fetal vessels at around 28–34 weeks, does not result in increased antenatal, obstetric and neonatal interventions, and no overall differences are detected for substantive short term clinical outcomes such as perinatal mortality [32]. On the other hand, the use of umbilical artery Doppler ultrasound in pregnancies with fetal growth restriction is associated with a reduction in perinatal deaths and obstetric interventions [33]. Guidelines published by the Society for Maternal Fetal Medicine confirm a decrease in induction of labor, cesarean delivery and perinatal death with use of umbilical artery Doppler assessment in high-risk pregnancies with fetal growth restriction. Surveillance with umbilical artery Doppler studies should be started once growth restriction is
suspected in the viable fetus [34]. Guidelines for the technical aspects of Doppler use in pregnancy are available [35] (see Chapter 45 in Maternal-Fetal Evidence Based Guidelines).

**Middle Cerebral Artery**

Fetal middle cerebral artery (MCA) peak systolic velocity (PSV) Doppler has been used to evaluate fetal anemia in cases of maternal red cell alloimmunization, parvovirus infection, or twin-twin transfusion syndrome in monochorionic twins. Fetal MCA Dopplers are considered a screening test that requires a confirmatory test for diagnosis (fetal blood sampling) at the initiation of therapy (transfusion). MCA-PSV is regarded as the best noninvasive screening test for fetal anemia [36,37] (see Chapter 53 in Maternal-Fetal Evidence Based Guidelines).

**Ductus Venosus**

The ductus venosus (DV) is a vascular shunt that connects the umbilical vein to the inferior vena cava in the fetus. This waveform is reflective of downstream pressure in the right atrium. In high resistance states (increased uteroplacental resistance), the DV shows absent or reversed flow in late diastole. A small retrospective study showed that reversed flow in the DV in addition to increased MCA is associated with perinatal mortality in fetuses less than 32 weeks gestation [38]. A meta-analysis including data from 2267 patients confirmed these data, showing moderate predictive value for fetal outcomes in high-risk pregnancies with placental insufficiency [39]. A randomized trial did not show significant perinatal benefit from adding DV screening to fetal heart rate monitoring alone for antepartum monitoring of the growth restricted fetus [40]. Therefore, there insufficient data to currently recommend the use of DV Doppler in the routine evaluation and management of the fetus with intrauterine growth restriction (IUGR) [34] (see Chapter 45 in Maternal-Fetal Evidence Based Guidelines).

**Uterine Artery**

The Doppler waveform of the uterine artery has been shown to reflect high impedance placental circulation by the presence of a waveform notch and low diastolic flow in association with hypertension and preeclampsia. Studies have shown an association between abnormal uterine artery Doppler and early onset preeclampsia, but predictive value is low. Current evidence has not shown a benefit to performing routine mid-pregnancy utero-placental Doppler ultrasound for prevention of preeclampsia, intrauterine growth restriction or adverse pregnancy outcome [41,42] (see Chapter 45 in Maternal-Fetal Evidence Based Guidelines). Furthermore, there is currently paucity of data to recommend the use of uterine artery Doppler in the clinical management of hypertensive pregnancies [42].

**BIOPHYSICAL PROFILE**

A fetal biophysical profile score (BPS) is a specialized obstetric ultrasound consisting of monitoring of fetal movements, tone and breathing, and assessment of amniotic fluid volume, with or without fetal heart rate monitoring. BPS has been used to identify fetuses that may be at high risk of poor pregnancy outcome. While information gained from a biophysical profile (BPP) regarding fetal status can help guide clinical management, available evidence from randomized trials does not support the use of BPS as an isolated test of fetal well-being in high-risk pregnancies. Additional evidence from larger trials is needed [43] (see Chapter 56 in Maternal-Fetal Evidence Based Guidelines).

**CERVICAL LENGTH**

TVU for the measurement of CL has been shown to be predictive of spontaneous PTB in all populations studied so far, including singletons and multiple gestations, either asymptomatic or with symptoms of preterm labor (PTL). TVU CL can be effective in evaluating the need for such interventions as cervical cerclage or progesterone supplementation, as well as more effective management of women with symptoms of PTL [44–47].

TVU CL screening is considered universal when offered routinely to singleton gestations without a prior spontaneous PTB at the time of the anatomy ultrasound, i.e., 18–24 weeks. Both ACOG and Society for Maternal-Fetal Medicine (SMFM) have published guidelines indicating that universal
TVU CL screening is deemed reasonable, but not mandatory [48,49]. About 1% of singletons without a prior spontaneous PTB develop a TVU CL ≤ 20 mm before 24 weeks [50], and should be started on vaginal progesterone daily until 36 weeks [51,52]. Over two-thirds of academic maternal-fetal medicine units in the United States perform universal CL screening [53].

Women with history of spontaneous PTB prior to 37 weeks gestation should be offered intramuscular progesterone supplementation [54]. In addition, current recommendations are for screening of these high risk women (history of spontaneous PTB) with serial TVU CL ultrasounds every 2 weeks from 16 to 24 weeks gestation. In those (about 40%) with TVU CL ≤ 25 mm, a cerclage can be offered [55].

TVU CL screening of singletons presenting between 24 and 34 weeks with symptoms of PTL has been associated with both lower evaluation and triage time, and significantly less incidence of PTB [56] (see Chapter 17).

THREE-DIMENSIONAL ULTRASOUND
3D ultrasound examination is not considered a required modality for all pregnant women at this time [4], but it can add accuracy in the assessment of the fetus identified to have anomalies by 2D ultrasound (especially facial anomalies, neural tube defects, and skeletal malformations). 3D ultrasound allows the acquisition of volume measurements, which can depict topographic anatomy not able to be seen on 2D imaging. New technology allows for 3D reconstruction of vascular structures, further characterizing vessel relationship [57], vascular malformations and vascular invasion. It has not been shown to have a clear clinical advantage over traditional ultrasound in routine settings [6].

Routine 3D ultrasound (versus the traditional 2D ultrasound) among low-risk women has not shown a significant impact on maternal-fetal bonding [58]. Additionally, 3D/4D ultrasound in women at risk for having a fetus with congenital anomalies does not reduce maternal anxiety compared with conventional 2D ultrasound alone [59].

FETAL ECHOCARDIOGRAPHY
The incidence of moderate to severe congenital heart disease (CHD) has been reported at 6–13 per 1000 live births [60,61], with most affected infants born to pregnancies without identifiable risk factors [62]. Fetal cardiac evaluation is an important part of the prenatal ultrasound examination. Basic cardiac screening to be completed during the mid-trimester ultrasound during every pregnancy includes the four-chamber view and evaluation of the right and left outflow tracts [5,10].

An in depth evaluation of fetal cardiac structures should be performed if the screening exam is abnormal or incomplete, or when there are maternal or fetal indications (Table 4.8) [63,64]. Fetal echocardiography is usually performed between 18 and 22 weeks. Main areas of evaluation include visceral situs, atrial and ventricular anatomy, valvular structure and function, and the orientation and morphology of the great vessels. Gray scale and color Doppler imaging are required while spectral Doppler and M-mode should be used as needed to evaluate suspected anomalies [65]. These structures are usually best seen in the second trimester, but experienced technicians and sonologists may be able to detect cardiac anomalies in the first trimester. One prospective observational study showed a sensitivity and specificity of 88% and 100%, respectively, for detection of cardiac anomalies when using the four-chamber view, three-vessel view, and three-vessel trachea view to screen an unselected patient population in the first trimester [66].

Table 4.8 Indications for Fetal Echocardiography

<table>
<thead>
<tr>
<th>Maternal indications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Autoimmune antibodies, anti-Ro/anti-La</td>
</tr>
<tr>
<td>Familial inherited disorders (e.g., 22q11.2 deletion syndrome)</td>
</tr>
<tr>
<td>In vitro fertilization</td>
</tr>
<tr>
<td>Metabolic disease (e.g., diabetes mellitus and phenylketonuria)</td>
</tr>
<tr>
<td>Teratogen exposure (e.g., retinoids, lithium)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Fetal indications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abnormal cardiac screening exam</td>
</tr>
<tr>
<td>First degree relative of a fetus with CHD</td>
</tr>
<tr>
<td>Abnormal heart rate or rhythm</td>
</tr>
<tr>
<td>Fetal chromosome anomaly</td>
</tr>
<tr>
<td>Extracardiac anomaly</td>
</tr>
<tr>
<td>Hydrops</td>
</tr>
<tr>
<td>Increased nuchal translucency</td>
</tr>
<tr>
<td>Monochorionic twins</td>
</tr>
</tbody>
</table>


REFERENCES


27. Bricker L, Medley N, Pratt JJ. Routine ultrasound in late pregnancy (after 24 weeks’ gestation). *Cochrane Library*. 2015. [meta-analysis; 13 RCTs, n = 34,980]


Prenatal diagnosis and screening for aneuploidy
Dawnette Lewis

KEY POINTS
- All women should be offered aneuploidy screening, ideally starting in the first trimester. Population screening should have genetic counseling services available to discuss the different modalities, advantages and disadvantages, and the time frame for each test prior to screening. All women should have counseling available if desired or if an "abnormal" result occurs. The difference between a screening and a diagnostic test should be clearly explained before any testing.
- Issues of sensitivity, specificity, positive, and negative predictive values are vital to the interpretation of a screening test. Positive predictive value of a screening test is greatly influenced by prevalence rates in the population tested.
- The performance of a screening test depends on the age of the women screened (which determines prevalence of trisomies), women's preference of screening methods, their choice of invasive testing, and their attitudes toward pregnancy termination. There is as of now no definitive noninvasive prenatal diagnostic test. The only diagnostic tests are invasive, i.e., chorion villus sampling (CVS) and amniocentesis.
- Compared with a Down's screening policy of amniocentesis for age ≥35 years of age and universal ultrasound at 18 weeks, a policy of nuchal translucency (NT) screening is associated with similar numbers of Down's neonates born and a decrease in invasive tests.
- First-trimester screening (STS) includes NT, pregnancy associated plasma protein-A (PAPP-A), and human chorionic gonadotropin (hHCG), and can be performed at 10 3/7 to 13 6/7 weeks, with the best detection rate achieved at 11 weeks. The overall detection rate (sensitivity) is about 84–87% (false positive rate or FPR, 5%). FTS should be offered only if
  - Appropriate training and ongoing quality monitoring programs are in place for both ultrasound (NT) and laboratory assays of analytes
  - Sufficient information and resources are available to provide comprehensive counseling to women regarding the different options and limitations of these tests
  - Access to an appropriate diagnostic test (i.e., CVS) is available when screening results are positive
- Compared with management using second-trimester screening (STS), management using FTS is associated with a significant reduction in induction for post-term pregnancy because of better dating with first-trimester ultrasound.
- Analyte screening (quadruple marker screen) can detect approximately 70%–81% of affected pregnancies (FPR 5%).
- Integrative screening has the best detection rate for Down's syndrome (95%), with a low (1%–4%) FPR, but results are available only in the second trimester. Stepwise or contingent sequential screening offer the same 95% detection rate, a reasonable 5% FPR, and availability of results in the first trimester.
- Cell-free fetal DNA screens for Trisomy 21, 18, and 13 and can be performed at ≥10 weeks. In a mixed population of low and high risk women the sensitivity of cell free DNA (cfDNA) for trisomy 21 was reported between 99% and 100%, for trisomy 18 between 90% and 99%, and for trisomy 13 greater than 99%, with a very low (usually <1%) FPR. Experts in general currently still recommend first- and second-trimester noninvasive aneuploidy screening over cfDNA in low-risk populations, but this is an area of intense research currently. Indications for cfDNA screening are currently maternal age ≥35 years at delivery; fetal ultrasound findings that indicate an increased risk for aneuploidy, specifically for trisomies 21, 18, 13; previous pregnancy with a trisomy detectable by cfDNA screening; positive screen results for aneuploidy; and parental balanced translocation with increased risk of fetal trisomy 13 or 21.
- Second-trimester "genetic" ultrasound has an impact, among other things, on dating, induction rates, and anatomic evaluation of the fetus. As a modality for genetic screening, the data is more limited compared with other available tests. Major anomalies (e.g., congenital cardiac defects and duodenal atresia), and some markers (especially nuchal thickening, short humerus or femur, and echogenic bowel) are associated with a significantly higher risk for Down's syndrome.
- Second-trimester amniocentesis is safer than transcervical (TC) CVS or early amniocentesis (<15 weeks). Early amniocentesis should never be performed. With expert operators (>400 CVS), CVS by any route may be as safe as second-trimester amniocentesis.
- If earlier diagnosis is required, transabdominal (TA) CVS is preferable to early amniocentesis or TC CVS. In circumstances where TA CVS may be technically difficult, the preferred options are TC CVS in the first trimester with expert operator, or second-trimester amniocentesis, per patients' preference.
- Chromosomal microarray analysis (CMA) should be recommended as the primary test (replacing conventional karyotype) to patients undergoing prenatal diagnosis in whom a structural abnormality is detected by ultrasound.
- Trisomy 21 is the most common trisomy at birth. Its incidence increases with increasing maternal age.

DEFINITION
Prenatal diagnosis: Prenatal diagnosis incorporates screening for fetal aneuploidies and anomalies, with many different
modalities, including population screening, individual risk assessment, genetic counseling and diagnostic testing.

SCREENING VERSUS DIAGNOSTIC TESTS
Prior to screening a population with an available test, test specifics should be assessed. Issues of prevalence, sensitivity, specificity, and positive and negative predictive values are vital to the interpretation of a screening test and are a large part of the problem that is in interpreting the value of a test by either practitioners or the public. Sensitivity of a screening test can also be called detection rate. Table 5.1 lists the characteristics of an ideal perinatal screening test. High sensitivity and specificity are preferable; however, the prevalence of a condition (based upon the population tested) will ultimately determine the value of a positive or negative result (Figure 5.1). With lower prevalence, the chance of a particular “positive” test to be a true finding is much less. For example, based upon the numbers from Figure 5.1, if all the women with a positive test from group “A” had a chorionic villus sampling (CVS), there would be one positive result for every 55 CVS performed. If all the women in “B” had a CVS, one out of every 2.4 tests would yield a positive result.

The performance of a screening test depends on the age of the women screened (which determines prevalence of

Table 5.1 Ideal Perinatal Screening Test

<table>
<thead>
<tr>
<th>Test</th>
<th>Disease</th>
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<tbody>
<tr>
<td>+</td>
<td>+</td>
<td>A</td>
<td>+</td>
</tr>
<tr>
<td>-</td>
<td>+</td>
<td>B</td>
<td>+</td>
</tr>
<tr>
<td>-</td>
<td>-</td>
<td>C</td>
<td>+</td>
</tr>
<tr>
<td>+</td>
<td>-</td>
<td>D</td>
<td>+</td>
</tr>
</tbody>
</table>

Sensitivity = A/A + C
Specificity = D/B + D
Positive predictive value (PPV) = A/A + B
Negative predictive value (NPV) = D/C + D

Effect of prevalence

N = 100,000
Sensitivity 90%
False positive = 5%
Prevalence = 0.1%

<table>
<thead>
<tr>
<th>Prevalence</th>
<th>Sensitivity</th>
<th>Specificity</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.1%</td>
<td>90</td>
<td>10</td>
</tr>
<tr>
<td>4%</td>
<td>3,600</td>
<td>400</td>
</tr>
</tbody>
</table>

PPV = 1.8%  NPV = 99.9%

Abbreviation: FPR, false positive rate.

Table 5.2 Screening Tests for Down’s Syndrome

<table>
<thead>
<tr>
<th>Test</th>
<th>FPR (%)</th>
<th>Sensitivity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>5</td>
<td>25–30</td>
</tr>
<tr>
<td>First trimester (11–14 weeks) NT</td>
<td>5</td>
<td>65–80</td>
</tr>
<tr>
<td>PAPP-A and β-HCG</td>
<td>5</td>
<td>60–80</td>
</tr>
<tr>
<td>Age, NT, PAPP-A, and β-HCG (FTS)</td>
<td>5</td>
<td>85</td>
</tr>
<tr>
<td>Age, NT, PAPP-A, and HCG (FTS)</td>
<td>5</td>
<td>80–85</td>
</tr>
<tr>
<td>Cell-free DNA</td>
<td>.5</td>
<td>99.9</td>
</tr>
<tr>
<td>Second trimester (15–21 weeks) Age, MSAFP, HCG, uE₃ (TS)</td>
<td>5</td>
<td>60–70</td>
</tr>
<tr>
<td>Age, MSAFP, HCG, uE₃, inhibin (QS)</td>
<td>5</td>
<td>70–81</td>
</tr>
<tr>
<td>Integrative (nondisclosure of FTS) Integrated (NT, PAPP-A, QS)</td>
<td>4</td>
<td>95</td>
</tr>
<tr>
<td>Serum integrated (PAPP-A, QS)</td>
<td>5</td>
<td>85–90</td>
</tr>
<tr>
<td>Sequential (disclosure of FTS) Independent</td>
<td>11</td>
<td>95</td>
</tr>
<tr>
<td>Stepwise</td>
<td>5</td>
<td>95</td>
</tr>
<tr>
<td>Contingent</td>
<td>5</td>
<td>88–94</td>
</tr>
<tr>
<td>Genetic ultrasound</td>
<td>5</td>
<td>50–70</td>
</tr>
<tr>
<td>Extended ultrasound</td>
<td>5</td>
<td>80–85</td>
</tr>
</tbody>
</table>

Abbreviations: uE₃, unconjugated estriol; TS, triple screen; *, genetic ultrasound and serum screening; QS, quadruple screen; FTS, first trimester screen; FPR, false positive rate.
diagnostic testing in higher risk populations can be offered.
Several studies have shown that most pregnant women prefer early (vs. later) screening for Down’s syndrome.

Many women still believe that the whole purpose behind screening for aneuploidy is so that a couple can terminate a pregnancy prior to viability. While it is true that couples faced with the reality of an aneuploid pregnancy may opt for termination, the purpose of prenatal diagnosis and screening is to provide information. If an abnormality is found, depending on the specifics, couples can be provided with specific information regarding their situation. When a couple decides to carry an abnormal pregnancy to term, the antenatal, intrapartum, and postpartum care can be performed under more ideal circumstances, hopefully altering the outcome. Also, one can not underestimate the effect that preparation can have for the individuals involved (Table 5.2).

ANTENATAL NONINVASIVE ANEUPLOIDY SCREENING

History
Langdon Down, in 1866, reported that the skin of individuals with trisomy 21 appears enlarged. In the 1970’s data became available on the relationship with maternal age and increased risk for aneuploidy. A statistically relevant difference was seen between the 30- to 34-year-old group, and the 35- to 39-year-old group, so that this difference led to the offering of women 35 years of age diagnostic evaluation for karyotype. Maternal serum alpha-fetoprotein (MSAFP) was originally found to be elevated in women carrying fetuses with neural tube defects (NTDs), then in 1984 a low MSAFP was associated with a higher risk of Down’s syndrome. NT first-trimester ultrasound screening was introduced by in the early 1990s [1]. While live births to women >35 years of age continue to increase, due to better, more diffuse screening the number of Down’s neonates is decreasing.

Principles
All women should be offered aneuploidy screening, ideally in the first trimester. There is presently a general consensus in the United States that invasive testing for Down’s syndrome be offered to those with a second-trimester risk of ≥1:270 or higher (live-born risk of 1:380). The cut-off level and subsequent public policy was determined over 30 years ago and was based on a maternal age risk of 35 years at delivery. Factors considered in determining this value included the prevalence of disease, a perceived significant increase in the trisomy 21 risk after this age, the risk of invasive testing, the availability of resources, and a cost benefit analysis. Since that time, a number of additional screening tests for Down’s syndrome have become available that enhance the validity of maternal age as a single indication for invasive testing. There are a limited amount of randomized control trials (RCTs) for the evaluation of different tests. Most data comes from cohort studies or cross-sectional analysis.

Age
The risk of fetal trisomy 21 increases with maternal age, but decreases with the gestational age (GA) at assessment in determining the risk, secondary to in utero death rates (Table 5.3) [2].

<table>
<thead>
<tr>
<th>Maternal age</th>
<th>Gestational age (weeks)</th>
<th>Live-born</th>
</tr>
</thead>
<tbody>
<tr>
<td>20</td>
<td>1/1068</td>
<td>1/1200</td>
</tr>
<tr>
<td>25</td>
<td>1/946</td>
<td>1/1062</td>
</tr>
<tr>
<td>30</td>
<td>1/626</td>
<td>1/703</td>
</tr>
<tr>
<td>31</td>
<td>1/543</td>
<td>1/610</td>
</tr>
<tr>
<td>32</td>
<td>1/461</td>
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<td>34</td>
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<tr>
<td>35</td>
<td>1/249</td>
<td>1/280</td>
</tr>
<tr>
<td>36</td>
<td>1/196</td>
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</tr>
<tr>
<td>37</td>
<td>1/152</td>
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<tr>
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<td>1/21</td>
<td>1/24</td>
</tr>
<tr>
<td>45</td>
<td>1/16</td>
<td>1/18</td>
</tr>
</tbody>
</table>

FIRST-TRIMESTER SCREENING

Nuchal Translucency
NT is elevated in fetuses with Down’s syndrome at 10 3/7 to 13 6/7 weeks (crown-rump length [CRL] about 36–86 mm). The NT measurement is obtained in a mid-sagittal plane with the neck of the fetus in a neutral position. The amnion should be enlarged >75% of the screen. The measurement is obtained from the inner to inner aspect of the NT, and multiples of the medians are used to calculate the Down’s risk via computer software. An increased NT is >70% sensitive for trisomy 21, trisomy 18, and trisomy 13 (Table 5.2) [1]. Rigorous training, certification, and on-going quality control are necessary to achieve the detection rates published in the literature (www.fetalmedicine.com; www.ntgr.org). The optimal time to perform NT for Down’s screening is about 11 weeks. Compared with a Down’s screening policy of amniocentesis for age ≥35 years of age and ultrasound at 18 weeks, a policy of NT screening is associated with similar numbers of Down’s neonates born (a nonsignificant 37.5% decrease) and a significant 82% decrease in invasive tests [3]. One explanation for the small nonsignificant difference in Down’s neonates born alive may be that NT screening certainly identifies better Down’s fetuses, but the majority of these identified Down’s fetuses are those that would have miscarried without intervention.

Biochemistry
The maternal serum analytes measured are B-HCG and PAPP-A. B-HCG normally decreases in pregnancy, but is increased in fetuses affected with trisomy 21. Free B-HCG performs better than total HCG as an independent marker, but there does not appear to be a clinically significant difference in sensitivity when either is combined with NT and PAPPA-A for FTS [4]. PAPP-A normally increases in pregnancy, but is decreased in fetuses affected with trisomy 21. HCG discrimination is greatest at 13 weeks, while PAPP-A’s is greatest at 10 weeks, making 11 weeks the optimal time for first-trimester analyte screening.

First Trimester Screening
FTS consists of measurement of the NT combined with maternal serum screening (PAPP-A and B-HCG). Over 20 studies
in over 200,000 women have been performed to assess the sensitivity of this screening test, making this the best studied screening test in pregnancy [4–7]. The GA for FTS is about 10 3/7 to 13 6/7 weeks (about 73–98 days; CRL about 36–86 mm). The usual cutoff risk is 1:270. The detection rate (sensitivity) is about 84–87% (95% CI 80%–90%). The best detection rate is achieved at 11 weeks.

FTS should be offered only if the following criteria can be met [4–7]:

1. Appropriate training and ongoing quality monitoring programs are in place for both ultrasound (NT) and laboratory assays of analytes.
2. Sufficient information and resources are available to provide comprehensive counseling to women regarding the different options and limitations of these tests.
3. Access to an appropriate diagnostic test (i.e., CVS) is available when screening results are positive.

The Maternal-Fetal Medicine Foundation (www.mfmf.org) and the Fetal Medicine Foundation (www.fetalmedicine.com) both provide NT education and quality review programs. There is sufficient evidence to support implementing FTS for Down’s syndrome provided the above three requirements are met. Almost 85% of women in the United States present for care within 12 weeks and can be offered FTS. FTS can provide high detection, early reassurance, more time / diagnostic options, earlier completion of aneuploidy screening.

Compared with management using STS, management using FTS is associated also with a significant reduction in induction for post-term pregnancy because of better dating with first-trimester ultrasound [8].

**Nasal Bone**

Over 12 studies in over 18,000 women demonstrated that nasal bone when imaged at 11–14 weeks is absent in approximately 70% of Down’s fetuses, and in only 1.5% of unaffected fetuses. When added to FTS (NT, PAPP-A, and B-HCG), it can increase the detection rate to about 95%, decreasing the FPR to 2%. Possibly given to the difficulty of this exam, these data have not been confirmed in all studies. Abnormal ductus venosus Doppler flow and tricuspid regurgitation have also been found to be >70% sensitive for trisomy 21, but there is insufficient prospective data for any increase in accuracy over FTS alone.

**SECOND-TRIMESTER SCREENING**

Maternal analyte screening had the first reported association with Down syndrome with low MSAFP (multiples of the median [MoM] 0.75), followed by the association of high hCG (MoM 2.3) and a low unconjugated estriol (MoM 0.7) to form the “triple screen.” The detection rate for women under 35 with a triple screen ranges between 57% and 74%, with a constant 5% FPR [3,5]. For women above 35 (using similar cut-off values), the sensitivity increases to 87%, but the FPR also balloons to 25% [9]. Inhibin was added to analyte screening (quadruple screen), but, since levels correlate somewhat with hCG, it is not an independent predictor like the other markers, and the increase in detection is more limited (7–11 percentage points). Approximately 70–81% of cases are detected in the majority of studies, holding the FPR at 5% [4,6]. This screening test can be performed between 15 and 22 weeks, with best results obtained at 16–18 weeks. Values need to be adjusted, as needed, for diabetes, obesity, and other factors.

Other combinations of markers have been assessed; however, with the addition of extra markers, the potential benefit versus the cost must be balanced. With each additional marker, costs to society reach into the millions secondary to the numbers of pregnancies tested each year. The relative cost and value of raising the sensitivity or lowering the false-positive rate a few percentage points is an ongoing debate.

**COMBINING BOTH FIRST- AND SECOND-TRIMESTER TESTS**

Combined screening programs in the first trimester (using both ultrasound assessments of the NT as well as maternal analytes) and the second trimester (using maternal analytes) have been described. Patterns of testing include sequential testing (results given after each test) and integrated testing (delaying reporting until both tests have been completed).

**Integrative Screening**

Performance of screening tests at different times during pregnancy with a single result provided to the patient only after all tests have been completed. A protocol for integrated screening for Down syndrome is based upon tests performed during the first and second trimester (NT, PAPP-A, MSAFP, HCG, estriol, and inhibin). Mathematical models calculated that >85% of affected pregnancies would be detected with a FPR of only 0.9%. The FASTER trial (First- And Second-Trimester Evaluation of Risk) performed integrated screening in 33,557 women (84 with Down syndrome) [6]. Cut-off values for the different tests varied (first-trimester combined test cutoff 1:150, second-trimester “quad screen”, 1:300). The authors report a sensitivity of 86% with first-trimester screening (FPR 5%), 85% with second-trimester (FPR high at 8.5%), and, when combined, a 94% detection of Down syndrome cases. If results are revealed after second-trimester screening, the FPR is only 4.9%, with the best sensitivity. If NT is not available, an “integrated serum screening test” has a detection rate of 85% with 3.9% FPR [4]. Disadvantages of integrative screening include the lack of early diagnosis, the physical and psychological ramifications created if an abnormality is found and the woman opts for termination (compared with the first trimester), the increase in costs (compared with either FTS or STS), the perception of “hiding” abnormal results, as well as the limitations it places on multiple gestations if discordant karyotypes are found.

**Sequential Screening**

It involves performance of different screening tests at different times during pregnancy with results provided to the patient after each test. There are three approaches to sequential testing: independent, stepwise, and contingent.

**Independent**

This approach involves the independent interpretation of FTS and STS. While the sensitivity is as exemplary with this approach as with integrative screening (94–95%), combining screening tests and revealing the results after each increases the chance for false-positive results. The FASTER trial’s [6] FPR was 10.8% with sequential independent screening, far too high for population-based usage. As a high FPR means higher loss rates due to more invasive testing, independent sequential screening is the least efficient risk assessment strategy, and should NOT be used.
**Step-Wise**

Both FTS and STS (usually quadruple screen or QS) are performed, with results revealed after FTS: If FTS risk is above a certain cutoff, invasive testing (i.e., CVS) is offered; if FTS is below a certain cutoff, STS is recommended with a final risk revealed at that point. In the FASTER trial, such an approach (low cutoff 1:150; high cutoff 1:300) had a detection rate of 95% with an FPR of 4.9% [6]. The advantages of this approach are a very high detection rate (as with integrative screening), with the option of early results in first trimester for the highest risk women. This is currently the most common way to perform screening for aneuploidy in women presenting in the first trimester in centers with adequate expertise and facilities. With FTS and such low FPRs, there has also been a decrease in the number of women requesting invasive prenatal diagnosis.

**Contingent**

Both FTS and STS (usually QS) are performed, with results revealed after FTS: If FTS risk is above a certain cutoff (e.g., 1/150), invasive testing (i.e., CVS) is offered; if FTS is below a certain cutoff (e.g., 1/300), no further screening is necessary; if FTS is in between, STS is recommended with a final risk revealed at that point. Careful determination of risk cutoffs is necessary. This strategy has not been studied prospectively.

**CELL-FREE FETAL DNA SCREENING**

The discovery of cell free fetal DNA (placental DNA) in maternal plasma opened up new possibilities for noninvasive maternal prenatal screening for fetal chromosomal aneuploidy. The methods utilized for detecting fetal aneuploidy differ based on whether amplified regions throughout the genome, chromosome specific regions or single nucleotide polymorphisms are the targets for sequencing are used [10–14]. In the high-risk population—those who have undergone prenatal diagnosis—the sensitivity of cfDNA for trisomy 21 is greater than 99%, for trisomy 18 greater than 99% and for trisomy 13 greater than 90% [15–19].

In a mixed population of low and high risk women the sensitivity of cfDNA for trisomy 21 was reported between 99% and 100%, for trisomy 18 between 90% and 99% and for trisomy 13 greater than 99% [19–21].

These studies have all shown that fetal fraction increases with GA and that the fetal fraction is decreased in obese patient. These studies also seem to say that an equivocal result or non-result on cfDNA should be cause for concern because this raises this possibility of aneuploidy.

The Society for Maternal-Fetal Medicine regarding prenatal aneuploidy screening using cell-free DNA made the following recommendations [22]:

**Indications:**

1. Maternal age ≥ 35 years at delivery
2. Fetal ultrasound findings that indicate an increased risk for aneuploidy, specifically for trisomies 21, 18, 13
3. History of a previous pregnancy with a trisomy detectable by cfDNA screening
4. Positive screen results for aneuploidy
5. Parental balanced translocation with increased risk of fetal trisomy 13 or 21

After failed cfDNA test, genetic counseling should be performed that includes offering diagnostic testing [15].

**“GENETIC” ULTRASOUND SCREENING FOR DOWN’S SYNDROME**

An ultrasound of the fetus performed at 18–24 weeks is associated with several important benefits (see Chapter 4). One of the benefits is the antenatal detection of anomalies. The identification of a fetus with an issue allows for directed counseling and optimization of antepartum, intrapartum, and postpartum care. Whether routine or targeted anatomic assessment is being performed, it should be done by experienced centers with ongoing quality assessment to increase detection of anomalies and limit false positive results.

The in-utero diagnosis of Down’s syndrome can be suspected when anomalies or physical features that occur more frequently in Down’s syndrome than in the general population are noted on an ultrasound examination. Certain of these major structural congenital anomalies, such as atrioventricular canal or duodenal atresia, strongly suggest the possibility of Down syndrome and are independent indications to offer invasive testing. Although, when present, there is a high risk of trisomy 21, these anomalies have low sensitivity and, thus, are not useful in screening. For example, when duodenal atresia is identified, there is approximately a 40% risk of Down’s syndrome, yet it is seen in only 8% of affected fetuses. About 50% of Down’s fetuses have congenital heart defects.

Physical characteristics that are not structural anomalies but occur more commonly in fetuses with Down’s syndrome are called **markers**. By comparing the prevalence of markers in Down’s syndrome fetuses to their prevalence in the normal population, a likelihood ratio (LR) can be calculated which can be used to modify risk. This is the basis for ultrasound screening for Down’s syndrome. In order for a marker to be useful for Down’s syndrome screening, it should be sensitive (i.e., present in a high proportion of Down’s syndrome pregnancies), specific (i.e., not commonly seen in normal fetuses), easily imaged in standard sonographic examination, and present early enough in the second trimester that diagnostic testing can be performed so that results are available when pregnancy termination remains an option. A list of presently available markers and LRs are seen in Tables 5.4 and 5.5, respectively [23–25]. Markers commonly sought to assess the risk of Down syndrome are discussed in the following sections.

<table>
<thead>
<tr>
<th>Table 5.4 Selected Ultrasound Findings Associated with Down’s Syndrome</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Major anomalies</strong></td>
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<tr>
<td>Congenital heart defects</td>
</tr>
<tr>
<td>Duodenal atresia</td>
</tr>
<tr>
<td><strong>Major markers</strong></td>
</tr>
<tr>
<td>Increased nuchal thickness</td>
</tr>
<tr>
<td>Hyperechoic bowel</td>
</tr>
<tr>
<td>Shortened humerus</td>
</tr>
<tr>
<td>Shortened femur</td>
</tr>
<tr>
<td>Echogenic intracardiac focus</td>
</tr>
<tr>
<td>Renal pyelectasis</td>
</tr>
<tr>
<td><strong>Minor markers</strong></td>
</tr>
<tr>
<td>Shortened or absent nasal bone</td>
</tr>
<tr>
<td>Foot length</td>
</tr>
<tr>
<td>“Sandal gap” of the foot</td>
</tr>
<tr>
<td>Widened ischial spine angle</td>
</tr>
<tr>
<td>Hypoplasia of the mid-phalynx of the fifth digit</td>
</tr>
<tr>
<td>Brachycephaly</td>
</tr>
</tbody>
</table>

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Increased Nuchal Fold
About 35% of Down’s syndrome fetuses, but only 0.7% of normal fetuses, have a nuchal skin fold measurement ≥6 mm (some studies use ≥5 mm). When an increased nuchal fold is an isolated finding, the LR is strong at 10–17. Thus, the presence of an increased nuchal fold alone is usually an indication to offer invasive testing.

Increased Echogenicity of the Fetal Bowel
When brighter than the surrounding bone, this marker has a Down’s syndrome LR of 3.0 to 6.7. This finding can also be seen with fetal cystic fibrosis, congenital cytomegalovirus (CMV) infection, swallowed bloody amniotic fluid, and severe fetal growth restriction (FGR).

Short Humerus, and in a Lesser Degree Short Femur
These markers in the second trimester are associated with Down’s syndrome, relative to the length expected from their biparietal diameter. This can be used to identify at-risk pregnancies by calculating a ratio of observed to expected (O/E) femur/humerus length based on the fetus’ biparietal diameter (BPD). An O/E ratio for femur length of <0.91 has a reported LR of 1.5–2.7 when present as an isolated finding. A short humerus is more strongly related to Down’s syndrome, with reported LRs ranging from 4.1 to 7.5.

Pyelectasis
This marker is defined as a renal anteroposterior (AP) diameter of ≥4–5mm, and has a LR that ranges from 1.1 to 1.9 as an isolated marker. This has been found by some not to be significantly more frequent in Down’s syndrome pregnancies than in normals (low specificity) [25].

Echogenic Intracardiac Foci
This marker occurs in up to 5% of normal pregnancies and in approximately 13–18% of Down syndrome gestations. The LR for Down’s syndrome when an echogenic focus is present as an isolated marker has ranged from 1.0 to 2.8. This has been found by most investigators not to be significantly more frequent in Down syndrome pregnancies than in normals (low specificity) [12]. The risk does not seem to vary if the focus is in the right or left ventricle or if it is unilateral or bilateral, but may be affected by ethnicity.

Other markers described include a hypoplastic fifth middle phalanx of the hand, short ears, a sandal gap between the first and second toes, an abnormal iliac wing angle, an altered foot to femur ratio, short or absent nasal bone, and others. These markers are inconsistently used because of the time and expertise required to obtain them. Mild ventriculomegaly (10–15 mm) can be an indication for invasive prenatal diagnosis, since it is associated with 1%–2% risk of aneuploidy if isolated. If the karyotype is normal, mild ventriculomegaly is still associated with about 8% structural anomalies, 3% perinatal death, and 10%–20% abnormal neurodevelopment.

Except for major anomalies and increased nuchal thickness, isolated ‘genetic ultrasound’ markers should in general not be used as the sole indication for invasive testing. As with other screening modalities, “genetic” ultrasound can be used to alter the a priori risk in either direction. A positive LR can be used to increase estimated risk. The magnitude of the increase depends upon the marker(s) or anomalies seen. While most of the clinical prospective data justifying this approach have come from a baseline age-related risk, some have advocated using these LRs to adjust whichever baseline risk, even that derived by other screening tests (e.g., FTS and/or QS, or even integrative or consecutive approaches). A benign second-trimester scan having none of the known markers and no anomalies has been suggested to have a LR of 0.4–0.5, assuming the image quality is satisfactory when the “genetic ultrasound” is normal. It is doubtful that the same sensitivity can be achieved in every center.

Ultrasound Screening for Other Chromosomal Abnormalities
Fetal aneuploidy other than Down’s syndrome can be suspected based on ultrasound findings [26]. The rates reported are usually in high-risk populations, and may overestimate the strength of the association when such findings are noted on a screening examination.

Trisomy 18
FTS and STS with MSAFP, HCG, unconjugated estriol and inhibin (QS) have a high detection rate for trisomy 18. Second-trimester ultrasound also has a high detection rate for trisomy 18.

Choroid plexus cysts (CPCs) have a very weak association with trisomy 18, and should not be the sole indication for invasive testing if isolated. The presence of CPCs CPC should be an indication for a detailed second-trimester ultrasound for trisomy 18 major anomalies, such as cardiac, central nervous system (CNS), hands defects, etc.

Positive Screening for Aneuploidy but Normal Karyotype
NT
A NT above the 95% percentile for GA, and especially ≥3.5 mm at 10 3/7–13 6/7 weeks is associated with an increased risk of other anomalies and syndromes, with the risk directly proportional to the increase in NT [27] (Table 5.6). The list of anomalies is long [14], and a detailed second-trimester ultrasound is recommended. The incidence of cardiac anomalies is ≥3.7% for NT ≥3.5 mm, so that a fetal cardiac ultrasound by experienced operator is recommended.
First Trimester PAPP-A and B-HCG
Low PAPP-A in FTS in the presence of a normal karyotype is associated with several adverse pregnancy outcomes, including fetal loss, PTB, and FGR. Low free HCG is associated with fetal loss. There are no randomized trials assessing any type of intervention or treatment for patients with abnormal serum markers [28].

Second Trimester Screening
High MSAFP is associated with NTDs, as well as abdominal wall defects and several other fetal abnormalities. High MSAFP, negative acetylcholinesterase (AChE) and normal ultrasound can be associated with congenital nephrosis or other syndromes, or normal pregnancy. Unexplained high MSAFP is associated with mild increases in the incidence of preeclampsia, abruptio, placental ischemia, preterm birth, fetal demise, low birth weight, and sudden infant death syndrome (SIDS). No trials have assessed specific management to prevent these complications [28].

Low unconjugated estriol is associated with steroid sulfatase deficiency, Smith–Lemli–Opitz or other conditions when very low, usually <0.3 MoM [28].

MSAFP Screening for NTD
Elevated (usually ≥2.5 MoM) MSAFP between 14 and 21 weeks is associated with a ≥90%–95% sensitivity for NTDs (false negative rate 5%). Given that ultrasound is also ≥95% sensitive for NTDs, the routine use of MSAFP screening is most important for pregnancies that will not have a detailed second-trimester ultrasound.

Screening for Aneuploidies in Twins
NT is accurate in estimating Down’s risk in dizygotic twins, using each NT separately for each fetus. In monochorionic twins, the average NT is the most effective screening method. Detection rates comparable to singletons can be achieved. Detection rates of FTS or STS tests are usually lower than in singletons, with higher rates of false positive and false negative results. Chorionicity does not seem to affect serum analytes in FTS or STS (see Chapter 44 in Maternal-Fetal Evidence Based Guidelines).

DIAGNOSTIC TESTS

CVS and Amniocentesis
Selected common indications for invasive prenatal diagnosis of fetal aneuploidy are listed in Table 5.7. Both CVS and amniocentesis have been performed for many years and can fairly safely diagnose a karyotypic or genetic abnormality.

Both procedures have been studied extensively. Differences in technique, as well as timing of the procedure, affect loss rates. To fairly compare procedure-induced loss rates between the two procedures, adjustments must be made for the higher background frequency of pregnancy loss earlier in gestation.

Second Trimester Amniocentesis
One study in a low-risk population (n = 4606) with a background pregnancy loss of around 2% found that a second-trimester amniocentesis increases total pregnancy loss by another 1% compared with no amniocentesis (relative risk [RR] 1.41; 95% confidence interval [CI] 0.99–2.00) [29]. Compared with no amniocentesis, second-trimester amniocentesis is associated with a 0.8% increase in spontaneous miscarriage (2.1% versus 1.3%; RR 1.60; 95% CI 1.03–2.52), but similar incidence of perinatal deaths (0.4% vs. 0.7%) [29,30]. In non-RCT data, the procedure-related risk of miscarriage following amniocentesis has been reported to be about 1/1000 [31].

There is insufficient data to assess the effect of PCR testing (fluorescent in situ hybridization [FISH]). In a small trial, reporting karyotype in 3 days with PCR did not affect maternal anxiety level compared with about 3 weeks later in Chinese women with an abnormal screening test for Down’s syndrome [32]. Another study compared median trait- and state-anxiety scores and found no difference between the two groups [33]. Therefore, there is insufficient evidence that, while waiting for the full karyotype following amniocentesis, issuing results from a rapid analysis reduces maternal anxiety. The limited evidence from the two trials does not help resolve the dilemma about whether full karyotyping should be abandoned in favor of limited rapid testing for women undergoing Down’s syndrome screening. This choice rests on clinical arguments and cost-effectiveness rather than impact on anxiety [34].

Early amniocentesis
Early amniocentesis (<15 weeks) is not a safe early alternative to second-trimester amniocentesis, because it is associated with increased pregnancy loss (7.6% versus 5.9%; RR 1.29; 95% CI 1.03–1.61), and higher incidence of talipes (1.8% versus 0.2%) compared with CVS (RR 4.61; 95% CI 1.82–11.66) [30,35].

Table 5.6 Risk of Chromosome Abnormalities and (If Normal Karyotype) of Fetal Death or Anomalies According to Nuchal Translucency

<table>
<thead>
<tr>
<th>NT</th>
<th>Chromosomal defects (%)</th>
<th>Normal karyotype</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;95th centile</td>
<td>0.2</td>
<td>1.3</td>
</tr>
<tr>
<td>95th–99th centiles</td>
<td>3.7</td>
<td>1.3</td>
</tr>
<tr>
<td>3.5–4.4</td>
<td>21.1</td>
<td>2.7</td>
</tr>
<tr>
<td>4.5–5.4</td>
<td>33</td>
<td>3.4</td>
</tr>
<tr>
<td>5.5–6.4</td>
<td>50</td>
<td>10</td>
</tr>
<tr>
<td>≥6.5</td>
<td>65</td>
<td>19</td>
</tr>
</tbody>
</table>

Abbreviation: NT, nuchal translucency.

Table 5.7 Selected Common Indications for Invasive Prenatal Diagnosis

- Abnormal first or second-trimester aneuploidy screen
- Abnormal ultrasound findings
- Parental concern/anxiety
CVS
CVS before 10 weeks is associated with an unacceptably high incidence of limb deficiencies, and should not be performed. The optimal time for CVS is 10–12 weeks, with trials of safety performed only in this GA. TA CVS can also be performed after 12 weeks (called usually placental biopsy) in cases in which placental karyotype is needed, but there are no trials on this “late” CVS.

Compared with second-trimester amniocentesis, CVS in general (TA and TC combined) is associated with a slight increased incidence of pregnancy losses and a slight increased incidence of spontaneous miscarriages [30].

Compared with second trimester amniocentesis, TC CVS is associated with a higher risk of pregnancy loss (14.5% versus 11%) and higher (12.9% versus 9.4%) risk in spontaneous miscarriage, although the results are quite heterogeneous and certainly operator-and experience-dependent [30].

Compared with second-trimester amniocentesis, TA CVS is associated with similar risk of pregnancy loss (6.3% versus 7.0%) and spontaneous miscarriage (3.0% versus 3.9%) in one study [30].

A systemic review of procedure related losses for CVS and amniocentesis revealed pooled loss rates for CVS at 14 days, 30 days and prior to 24 weeks of 0.7%, 1.3%, and 1.3% respectively. The pooled loss rates for amniocentesis for the same time period were 0.6%, 0.8%, and 0.9% [37]. In non-RCT data, the procedure-related risk of miscarriage following CVS has been reported to be about 2/1,000 [31]. Since CVS and amniocentesis are performed at different time periods, it is difficult to compare these procedures due to the higher background loss rate for the time period when CVS is performed.

The learning curve for TA and TC CVS has been estimated to exceed 400 cases, with post-procedure loss rates for operators having performed less than 100 cases being two to three times higher when compared with more experienced operators. The importance of operator experience cannot be overemphasized, particularly for route of CVS, with TC CVS requiring more experience.

TC vs. TA CVS

Compared with TA CVS, TC CVS is associated with similar pregnancy loss rates (9.0% vs. 7.4%) and similar spontaneous miscarriages (79% vs. 4.5%) [30]. The results related to comparative pregnancy loss between TA and TC CVS are inconclusive, with significant heterogeneity between studies [30].

TC CVS technical instrument

There is some evidence to support the use of small forceps compared with cannulas for TC chorionic villus sampling. When different types of cannulae are compared, Portex cannula is more likely to result in an inadequate sample and a difficult or painful procedure when compared with either the silver or aluminum cannula respectively. The evidence is not strong enough to support change in practice for clinicians who have become familiar with aspiration cannulae, and no recent studies have been performed [38].

Microarrays and Other Genetic Tests

Advances in technology have demonstrated many new avenues for noninvasive diagnostic testing in uterus (i.e., fetal cells from maternal circulation or cervical sampling, free fetal DNA in the maternal circulation, etc.).

Array comparative genomic hybridization (aCGH) is a new technique (based on invasive prenatal samples such as CVS and amniocentesis results) which has found clinical use. It can query the entire human genome for copy number changes such as aneuploidy, deletions, duplications, and unbalanced translocations. Unlike traditional cytogenetics which requires dividing cells, aCGH does not [39–44]. Conventional karyotype remains the principal cytogenetic tool for prenatal diagnosis, but the indications for aCGH are

- Abnormal ultrasound findings with normal karyotype
- Intrauterine fetal demise with congenital anomalies and culture failure with conventional karyotype [45]

In a high risk population referred for prenatal diagnosis, chromosomal microarray analysis (CMA) was compared with conventional karyotyping. The indications for prenatal diagnosis were ultrasound abnormalities, advanced maternal age or a positive result on prenatal screening. Microarray was similar to conventional karyotyping in detecting common chromosomal aneuploidy. In addition microarray was able to detect clinically significant aneuploidies not detected by karyotype [46]. Another study demonstrated that microarray was more likely to provide genetic results after stillbirth when compared with conventional karyotype [47]. When structural abnormalities are detected by prenatal ultrasound, chromosomal microarray can identify clinically significant chromosomal abnormalities in approximately 6% of the fetuses that have a normal karyotype [46]. For this reason, CMA should be recommended as the primary test (replacing conventional karyotype) to patients undergoing prenatal diagnosis in whom a structural abnormality is detected by ultrasound.

COMMON KARYOTYPE ABNORMALITIES

Trisomy 21 (Down’s Syndrome)

Historic Notes

First complete description in 1846 by Seguin. Report by Down in 1866 established the name of the syndrome. In 1959 LeJeune and Jacobs independently described that Down syndrome was caused by trisomy 21.

Definition

Down syndrome is trisomy 21, or the presence of an extra chromosome number 21, either as three number 21s, or as a translocation between 21 and another chromosome, usually an acrocentric in a Robertsonian translocation.

Epidemiology/Incidence

About 1 in 800 live births. This is the most common trisomy at birth. Incidence increases with increasing maternal age.

Embryology

The Down’s syndrome critical region on chromosome 21 is being studied extensively to identify the genes involved in the Down’s syndrome phenotype. However, this region is not one small isolated spot, but most likely several areas on chromosome 21 that are not necessarily side by side.
• Biometry may reveal symmetric FGR by the third trimester
• Several “soft markers,” such as short humerus and femur, hydrops or some hydropic changes (pleural effusion, ascites, etc.)
• Several “soft markers,” such as short humerus and femur, echogenic bowel, renal pyelectasis, cardiac (usually left ventricular) echogenic focus, short middle phalanx 5th digit, “sandal foot,” iliac crest >90° angle, short ear length
• Biometry may reveal symmetric FGR by the third trimester
• Amniotic fluid: Polyhydramnios (if gastrointestinal [GI] obstruction or macroglossia present)
• Placenta: Normal

Ultrasound after 14 weeks only detects ~50% of fetuses with Down syndrome. CPCs do not increase the risk. Screening provides the mother and family with a risk for Down's syndrome, but true diagnosis can only be achieved with CVS or amniocentesis.

Diagnosis
CVS or amniocentesis achieve the diagnosis by a study of the fetal chromosomes, which reveal trisomy 21. In the neonate, usually peripheral blood is cultured and karyotyped.

Counseling
The major abnormalities are increased risk of FGR, congenital heart defects, fetal and postnatal death, and developmental delay, with average IQ 50–75. Congenital heart defects are major contributors to mortality.

Work-Up/Investigations and Consultations
A fetal echocardiogram is recommended. Depending on the lesions detected, specific pediatric subspecialty consultation can be offered. Genetic counseling can be offered as well. Care in a tertiary care center is indicated if there are significant associated anomalies, or if they cannot be ruled-out adequately.

Fetal Intervention
None available.

Termination Issues
Termination can be offered as sole intervention as regulated by local law (usually legal <24 weeks)

Fetal Monitoring/Testing
No specific trials. Nonstress tests (NSTs) weekly at ≥32 weeks can be offered. Nonreassuring fetal heart rate (NRFHR) is common.

Delivery/Anesthesia
Mode and management of delivery should not be affected by the diagnosis of Down syndrome. NRFHT is common.

Neonatology Management

Resuscitation
Providing life support as needed as in any other infant is generally appropriate.

Transport
Indicated if counseling, general care, and/or major anomalies cannot be assessed and treated adequately at the birth institution.

Testing and Confirmation
Karyotype is usually confirmed by blood lymphocyte culture.

Nursery Management
Neonatal echocardiogram, and physical exam to assess any anomaly. Surgery may need to be scheduled for GI or cardiac anomalies. Down's syndrome presents with a wide variety of features and characteristics. There is a wide range of intellectual disability and developmental delay noted among children
with Down's syndrome. There is a great deal of variability in the presence of other anomalies such as CHD, GI, and hematological problems in these children. Hypothyroidism occurs in a high percentage of individuals with DS and should be monitored closely for their lifetime. Early intervention and specialized help with education and home rearing has improved the outcome in children with Down's syndrome. Many young adults with Down syndrome move into community living arrangements and work regular jobs or in sheltered workshops.

Future Pregnancy Preconception Counseling
With full trisomy 21, the recurrence risk is empirically 1% or the age related risk as a woman gets older. With Robertsonian translocation, parental chromosomes should be checked, with genetic counseling regarding specific future risks. There are other rare translocations leading to DS. One is a Robertsonian translocation between two chromosomes 21, t(21;21); this has a 100% risk for DS when transmitted by a carrier parent. Also rare is a non-Robertsonian translocation formed by the union of two 21s such that the translocation forms a mirror image of the normal 21. There is some literature that suggest in some families where there have been recurrent trisomies, a relationship exists with their MTHFR status. This has not been proven in large studies.

Helpful Websites
General: http://www.ds-health.com/
Health care guidelines for care of individual with DS: http://www.denison.edu/collaborations/dsq/health96.html

Trisomy 18 (Edward Syndrome)

Historic Notes
Trisomy 18 was independently described by Edwards et al. and Smith et al. in 1960.

Definition
Edward syndrome is trisomy 18, or the presence of an extra chromosome number 18.

Epidemiology/Incidence
Incidence of 1 in 6600 live births in the United States and the United Kingdom. This is the second most common trisomy at birth. Incidence increases with increasing maternal age.

Embryology
Extra number 18 affects development of all organs.

Genetics/Inheritance/Recurrence
Extra chromosome number 18 is usually (95%) secondary to de novo meiotic nondisjunction associated with advanced maternal age. In approximately 90% of cases, the extra chromosome is maternal in origin, with meiosis II errors occurring twice as frequently as meiosis I errors. Other human trisomies have a higher frequency of nondisjunction in maternal meiosis I. Approximately 80% of nondisjunctions occurs in females. Mosaicism occurs in approximately 10% and is due to postzygotic nondisjunction or anaphase lag. The causes of meiotic and mitotic nondisjunctions are unknown. Translocations may also result in trisomy or partial trisomy 18 with varying phenotype due to monosomy of another chromosome and variable size of piece of chromosome number 18 involved. The smallest extra region necessary for expression of serious anomalies of trisomy 18 appears to be 18q11–q12.

Teratology
None.

Classification
Trisomy 18 (95%), mosaic trisomy 18, and variable partial trisomy 18 related to translocations.

Risk Factors/Associations
Advanced maternal age and translocation carriers have increased risk. Recurrence risk approximately 1% for full trisomy 18.

Pregnancy Management
Screening
NT has a sensitivity of >80%; FTS has a sensitivity of >90%, second-trimester multiple marker screening is typically low alpha-fetoprotein (AFP), low HCG, low estriol with a sensitivity of about >80%. Accurate ultrasound is usually >90% sensitive for trisomy 18.

Ultrasound Findings in Fetus
- Thickened NT at 11–14 weeks (80%) (at times cystic hygroma ~15%)
- Thickened nuchal fold (≥6 mm) at 16–23 weeks
- CHD (90%)
- Omphalocele (25%)
- NTDs (20%)
- Clenched hands with overlapping fingers
- Clubbed or rocker-bottom feet
- CPCs (25%) (most commonly isolated and seen in normal fetuses; karyotyping probably not indicated if isolated)
- Enlarged (>1 cm) cisterna magna
- Single umbilical artery
- Micrognathia
- Cleft lip or palate
- Hydrops or some hydropic changes (pleural effusion, ascites, etc.)
- Biometry may reveal FGR by the third trimester
  - Amniotic fluid: Polyhydramnios (25%)
  - Placenta: Normal
  - Biometry/measurement data: Symmetric FGR (>50%) microcephaly in third trimester
  - When detectable: At 11–14 weeks if increased NT is detected

Diagnosis
CVS or amniocentesis

Counseling [48,49]:
Approximately 95% of conceptuses with trisomy 18 die in embryonic or fetal life. Five to ten percent of affected children born alive survive beyond the first year of life. In utero, there are decreased fetal movements. Clinical findings the parents should be informed about include severe psychomotor and growth delay, microcephaly, microphthalmia, malformed ears, micrognathia or retrognathia, microstomia, distinctively clenched fingers, rocker-bottom feet and other congenital malformations. CHD occurs in 90%, with ventricular septal defect (VSD) and poly-valvular heart disease (pulmonary and aortic valve defects) common. Renal anomalies, GI and brain malformations are common. Classical dermatoglyphics with digital arch patterns on finger and toe tips and distal palmar triradius with hypoplastic finger tips and small nails. Central apnea is
Patau syndrome is trisomy 13, or the presence of an extra chromosome number 13.

**Definition**
Patau syndrome is trisomy 13, or the presence of an extra chromosome number 13.

**Prenatal Diagnosis and Screening for Aneuploidy**

A frequent cause of death, along with cardiac, CNS and renal malformations.

If diagnosed prenatally, recommend discussion with parents about how to proceed in labor and delivery allowing “nature to take its course” without monitoring, or level of intervention desired by parents including the extent of resuscitation after delivery. Indication for C-section for fetal indications may be futile. Parents need to be counseled that some children with trisomy 18 do survive and require lifelong complete care, but never achieve any independence. Few milestones are reached. There is an increased incidence Wilms tumor in trisomy 18 children who survive. Cardiac surgery is controversial. In first weeks it may be considered a heroic measure, but if the child is surviving it may make life more comfortable (comfort care). Apnea is a common cause of death, and can happen at home; there is the need to understand “nobody’s fault” if this happens.

Work-up/Investigations and Consultations Required

A fetal echocardiogram is recommended. Genetic counseling can be offered. Neonatal consultation is extremely important, to help the couple decide regarding neonatal management; usually just comfort care for the baby and psychological support for the parents is most appropriate. Fetal intervention: None available.

**Termination Issues**
Termination can be offered as sole intervention as regulated by local law (usually legal <24 weeks).

**Antepartum Testing**
As NRFHT is very common, and prognosis poor, fetal testing is not recommended. Many pregnancies continue without spontaneous labor until post-term (>42 weeks).

**Delivery/Anesthesia**
Fetal heart monitoring is usually declined, and not indicated. Every attempt should be made to maximize the chances of vaginal delivery to minimize maternal morbidity given frequently fatal neonatal prognosis. Cesarean delivery for fetal indications is not recommended and should be discussed.

**Neonatology Management**
Resuscitation
Comfort care only. Allow parents to grieve appropriately. Providing life support is usually not appropriate.

Transport
Not indicated.

Testing and confirmation
Karyotype is usually confirmed by blood lymphocyte culture.

Future Pregnancy Preconception Counseling
Test parents if due to translocation.

**Helpful Websites**
http://www.emedicine.com/ped/topic652.htm

**Trisomy 18**

**Historic Notes**
Patau first identified in laboratory in 1960 noting three of the group 13–15 chromosomes.

**Definition**
Patau syndrome is trisomy 13, or the presence of an extra chromosome number 13.

**Epidemiology/Incidence**
1 in 10,000 live births. Incidence increases with increasing maternal age. Approximately 1% of all first-trimester spontaneous losses are due to trisomy 13.

**Embryology**
Extra chromosome number 13 affects development of all organs.

**Genetics/Inheritance**
Extra number 13 chromosome resulting in full trisomy 13 (80% of cases). This is due to maternal nondisjunction usually in meiosis I. About 15% of cases are due to translocation, mostly Robertsonian translocation t(13q14). In 5% translocation is familial with recurrence risk of 5% and risk of spontaneous abortion (SAB) of 20%. The other cases are due to mosaicism (5%) with trisomy 13 and a normal cell line. Mosaicism cases may have milder phenotype.

**Teratology**
None.

**Classification**
Trisomy 13, mosaic trisomy 13 and translocation trisomy or partial trisomy 13.

**Risk Factors/Associations**
Advanced maternal age. Individuals who are carriers of balanced Robertsonian translocations involving number 13 have an increased risk.

**Pregnancy Management**

**Screening**
First-trimester ultrasound (with NT). First or second-trimester multiple marker screening are not sensitive and clinically useful for detecting trisomy 13. Accurate ultrasound is usually 90% sensitive for trisomy 13.

**Ultrasound Findings in Fetus**
Thickened NT at 11–14 weeks (>70%) (at times cystic hygroma ~20%)
- Thickened nuchal fold (26 mm) at 16–23 weeks
- CHD (80%) (atrial septal defect (ASD) and VSD most common, but also often complex CHD)
- Holoprosencephaly (40%)
- Cleft lip and palate (45%)
- Hypotelorism/microphthalmia
- Polydactyly
- Rocker-bottom feet
- Omphalocele (10%)
- Polycystic kidneys (30%)
- Enlarged (>1 cm) cisterna magna (15%)
- NTDs
- Hydrops or some hydroptic changes (pleural effusion, ascites, etc.)
- Biometry may reveal symmetric FGR by the third trimester
  - Amniotic fluid: Polyhydramnios or oligohydramnios
  - Placenta: Normal
  - Biometry/measurement data: Symmetric FGR (50%)
  - When detectable: At 11–14 weeks if increased NT is detected

**Diagnosis**

CVS or amniocentesis achieve the diagnosis by a study of the fetal chromosomes, which reveals trisomy 13. In the neonate, usually peripheral blood is cultured and karyotyped.
Counseling

Most trisomy 13 conceptions result in SABs. Median survival is fewer than 3 days. Mean life expectancy is 130 days with most dying in first month of life, 95% die within 6 months. Apnea is a common cause of death, and can happen at home; there is the need to understand “nobody’s fault” if this happens. Family needs to be prepared for intense care needs and possible sudden death. Most common causes of death are cardiopulmonary arrest, 69%; CHD, 13%; and pneumonia, 4% [48,49].

Some infants do survive and those need complete care and achieve few milestones. Survival depends on associated medical problems. Survivors with trisomy 13 have severe intellectual disability and developmental delays. For survivors there are specific growth charts available for monitoring growth. Children with trisomy 13 are irritable, do not achieve milestones beyond smiling and most need to be fed by tube.

If diagnosed prenatally, recommend discussion with parents about how to proceed, usually allowing “nature to take its course” given the grim prognosis. Parents need to be counseled that some children with trisomy 13 do survive and require life-long complete care, never achieve any independence.

Work-up/Investigations and Consultations

A fetal echocardiogram is recommended. Genetic counseling can be offered. Neonatal consultation is extremely important, to help the couple decide regarding neonatal management; usually just comfort care for the baby and psychological support for the parents is most appropriate.

Fetal intervention
None available.

Termination Issues

Termination can be offered as sole intervention as regulated by local law (usually legal <24 weeks).

Antepartum Testing

As NRFHT is very common, and prognosis poor, fetal testing is not recommended. Many pregnancies continue without spontaneous labor until post-term (>42 weeks).

Delivery/Anesthesia

Fetal heart monitoring is usually declined, and not indicated. Every attempt should be made to maximize the chances of vaginal delivery to minimize maternal morbidity given almost universally fatal neonatal prognosis.

Cesarean delivery for fetal indications is not recommended and should be discussed.

Neonatology Management

Resuscitation Comfort care only. Allow parents to grieve appropriately. Providing life support is usually not appropriate.

Transport
Not indicated.

Testing and confirmation

Karyotype is usually confirmed by blood lymphocyte culture.

Long-term care

Feeding issues, gastrostomy; irritability; chronic infections, aspiration pneumonia; heart failure; frequent hospitalizations; seizures; blindness and hearing loss; few milestones achieved (smile, laugh); parental stress.

Future Pregnancy Preconception Counseling

With full trisomy 13, the recurrence risk is empirically 1% or the age related risk as a woman gets older. With Robertsonian translocation, parental chromosomes should be checked, with genetic counseling regarding specific future risks. There are other rare translocations leading to trisomy 13. Rare translocation of t(13q13q) the risk of recurrence or SAB is 100%.

Helpful Website
http://www.emedicine.com/ped/topic1745.htm

Turner Syndrome

Historic Notes

In 1938 Turner described the combination of sexual infantilism, webbed neck and cubitus valgus. Ford showed in 1959 that this combination of findings was associated with a missing X chromosome.

Definition

Turner syndrome is the presence of single X chromosome, or any karyotype with Xp missing such as isochromosome Xq, ring X or deletion Xp. Also called 45X0 or 45X syndrome.

Epidemiology/Incidence

1/2500 female births (1/5000 total births). Approximately 98%–99% of Turner fetuses are spontaneously aborted; about 20% of all SABs are due to Turner syndrome.

Embryology

Lymphedema usually due to congenital hypoplasia of lymphatic channels.

Genetics/Inheritance

The presence of single X chromosome, or any karyotype with Xp missing such as isochromosome Xq, ring X or deletion Xp. The presence of a single X chromosome results from chromosomal nondisjunction. Mosaicism is common (40%) and may include a 46,XY karyotype associated with ambiguous genitalia. Since features of Turner syndrome are seen in other syndromes, karyotype is essential to make the diagnosis. Chromosome studies on more than one tissue may be needed to detect mosaicism. Not associated with advanced maternal age.

Teratology
None.

Classification

45,X in 50%. 46,X,i(Xq) in 17%, mosaicism in 40%.

Risk Factors/Associations

Not associated with advanced maternal age. Differentiate from Noonan syndrome by karyotype which is normal in Noonan syndrome.

Pregnancy Management

Screening First-trimester screen with NT measurement. Biochemical screening is usually not sensitive enough for clinical use.

Ultrasound findings in fetus

- Cystic hygroma
- Thickened nuchal fold (≥ 6mm) at 16–23 weeks;
- CHD (20%) (usually left side: coarctation, aortic stenosis, bicuspid aortic valve, left hypoplastic heart)
• Renal anomalies (60%)
• Hydrops or some hydropic changes (pleural effusion, ascites, etc.)
• Amniotic fluid: Occasionally oligohydramnios
• Placenta: Normal
• Biometry/measurement data: Usually normal
• When detectable: At 10+ weeks if cystic hygroma detected

Diagnosis

CVS or amniocentesis achieve the diagnosis with a study of the fetal chromosomes, which reveals 45,X or missing Xp. In the neonate, usually peripheral blood is cultured and karyotyped.

Counseling

45,X conceptions frequently (>95%) end in SAB. The presence of a cystic hygroma with the diagnosis of Turner syndrome is >99% fatal. If cystic hygroma is not present or resolves, and fetus is still alive >20 weeks, many survive until birth. Female infants with Turner syndrome have excess nuchal skin and edema of the hands and feet (80%) due to lymphedema. CHD, if present, usually most affects prognosis, requires surgery and long-term care. In childhood short stature is apparent. Teenagers have delayed puberty and primary amenorrhea (>90%), with infertility (>99%). Other clinical findings include shield-shaped chest with widely spaced nipples, low posterior hair line with webbing or shortness of neck, renal anomalies (60%), cubitus valgus, short 4th metacarpal, narrow, hyper convex and deep set nails, hearing loss and thyroid dysfunction. Some girls with Turner syndrome have learning difficulties including difficulty with math and reading maps related to a deficit in spatial ability. Intelligence and verbal skills are usually within the normal range. Mosaicism with a normal female cell line may result in a milder phenotype and spontaneous puberty with fertility but often early menopause. If there is a deleted X but the X-inactive specific transcript (XIST) locus is intact, normal random X-inactivation may occur and the phenotype may be milder. If XIST is not present in a small X chromosome marker the phenotype may be more severe. In mosaic 45X/46,XY individuals clitoral enlargement may be present and virilization may occur. In these cases there is an increased risk of gonadoblastoma and the gonad should be removed. Psychological impact of short stature, infertility, and learning difficulties needs to be discussed.

Work-up/Investigations and Consultations

A fetal echocardiogram is recommended. Depending on the lesions detected, specific pediatric subspecialty (in particular for cardiac anomalies) consultation can be offered. Genetic counseling can be offered as well. Care in a tertiary care center is indicated if there are significant associated anomalies, or if they cannot be ruled-out adequately. Endocrine follow-up is indicated.

Fetal intervention

None available.

Termination issues

Termination can be offered as sole intervention as regulated by local law (usually legal <24 weeks).

Fetal monitoring/testing

No specific trials. NSTs weekly at ≥32 weeks can be offered.

Delivery/anesthesia

Mode and management of delivery should not be affected by the diagnosis of Turner syndrome.

Neonatology Management

Resuscitation

Providing life support as needed as in any other infant is generally appropriate.

Transport

Indicated if counseling, general care, and/or major anomalies cannot be assessed and treated adequately at the birth institution.

Testing and confirmation

Karyotype is usually confirmed by blood lymphocyte culture.

Nursery Management

Neonatal echocardiogram, renal ultrasound, and physical exam are indicated to assess any anomaly. Surgery may need to be scheduled for cardiac anomalies. Early intervention and specialized help with education has improved the outcome in children with Turner syndrome.

Long-Term Care

Thyroid studies annually; hearing test if otitis and not done before; speech evaluation, if needed; blood pressure checks routinely (hypertension a complication); annual echocardiogram to measure aortic root; annual urinalysis and culture if renal anomaly; use Turner growth curve after 2 years old; monitor diet (calories and calcium); ophthalmology follow-up as indicated; psychological support; individualized education plan (IEP) at school if indicated; refer to endocrinologist in infancy, discuss growth hormone (GH) and hormone replacement therapy (HRT): GH treatment can improve growth and influence a girl’s final adult height. HRT helps the girl with Turner syndrome develop the physical changes of puberty. In vitro fertilization can make it possible for some women with Turner syndrome to become pregnant using a donor egg. It is important to discuss at what age to inform the child of her diagnosis and its implications.

Future Pregnancy Preconception Counseling

Recurrence risk in 45,X is not increased over population risk. There is an increased risk if associated with a translocation.

Klinefelter Syndrome

Historic Notes

In 1942, Dr. Harry Klinefelter described males who had enlarged breasts, sparse facial and body hair, small testes, and azoospermia. By the late 1950s these findings were associated initially with an extra Barr body and later the extra X chromosome was identified with the karyotype 47,XXY.

Diagnosis/Definition

Chromosome study 47,XXY. There are no specific phenotypic features to identify Klinefelter syndrome in an infant.

Epidemiology/Incidence

1 in 500 to 1 in 1000 male births.

Genetics/Inheritance/Recurrence/Future Prevention

Advanced maternal age slightly increases the risk for the XXY. Recent studies have shown that half the time, the extra chromosome comes from the father.

Risk Factors/Associations

Advanced maternal age.
Screening
No phenotypic features noted prenatally.

Clinical Features
Occasional breast enlargement, lack of facial and body hair, and a female-type body configuration. Small testes. Taller than others in their family. Delayed speech occurs in >50%. Poor gross motor coordination is present in ~27%. School difficulties are relatively common and many boys with 47,XXY need assistance at school. Many are shy and somewhat passive and easy babies to care for. The average IQ is 90, verbal IQ higher than performance IQ. XXY boys enter puberty normally, without any delay of physical maturity. But as puberty progresses, they fail to keep pace with other males. Most XXY boys benefit from receiving an injection of testosterone every 2 weeks, beginning at puberty.

Counseling
Regular injections of the male hormone testosterone, beginning at puberty, can promote strength and facial hair growth as well as bring about a more muscular body type. Psychological support and therapy can help with self-esteem issues and interaction with peers. Depression also may be a problem in adults.

Boys with 47, XXY have a slightly increased risk of autoimmune disorders such as type I (insulin dependent) diabetes, autoimmune thyroiditis, and lupus erythematosus. XXY males with enlarged breasts have the same risk of breast cancer as do women—roughly 50 times the risk of XY males. XXY males who do not receive testosterone injections may have an increased risk of developing osteoporosis in later life.

It is unnecessary to share this diagnosis outside of medical providers as diagnosis may be misunderstood.

Rare/Related Variations include the XY/XXY mosaic who may have enough normally functioning cells in the testes to allow them to father children. Males with two or even three additional X chromosomes have also been reported in the medical literature. In these individuals, the classic features of Klinefelter syndrome may be exaggerated, with low IQ or moderate to severe mental retardation also occurring. Testosterone injections may not be appropriate for all of them.

Helpful Website
http://www.nichd.nih.gov/publications/pubs/klinefelter.htm#xwhat

47,XXX
Diagnosis/Definition
Karyotype shows 47,XXX.

Epidemiology/Incidence
1 in 1000 newborn females.

Genetics/Inheritance/Recurrence
Sporadic, increased by advanced maternal age.

Risk Factors/Associations
Advanced maternal age.

Screening
No identifying physical features. Must have karyotype to make diagnosis.

Clinical Features
Girls with 47,XXX are usually tall. Pubertal development is usually normal and fertility is probably normal but there may be an increased incidence of offspring with chromosome abnormalities.

Counseling
Girls with 47,XXX are shy and may demonstrate immaturity. The support of a loving and understanding family can improve the outcome for these girls. Many have learning difficulties (math and reading) but they are not intellectually disabled. The IQ of a girl with 47,XXX may be a few points lower than that of her siblings. Those diagnosed on amniocentesis or CVS with normal ultrasound, where indication is AMA, have better prognosis than those diagnosed postnatally because a problem has been noted.

SELECTED MICRODELETIONS AND DUPLICATIONS

DiGeorge Syndrome (22q11.2 Deletion Syndrome)

Historic Notes
DiGeorge syndrome, described in 1965 and velo-cardiofacial syndrome described in 1978 are different manifestations of the same deletion of chromosome 22q11.2.

Diagnosis/Definition
FISH study reveals an interstitial deletion of chromosome 22 p11.2. Can be detected on microarray.

Epidemiology/Incidence
1 in 2000–4000 births.

Embryology
Defects occur in the 3rd and 4th pharyngeal pouches, which later develop into the thymus and parathyroid glands. Developmental abnormalities may also occur in the 4th branchial arch.

Genetics/Inheritance/Recurrence/Future Prevention
Autosomal dominant inheritance. In 6% of affected individuals one of the parents is affected. Expression is very variable so both parents of an affected child should be tested for the deletion.

Risk Factors/Associations
Other conditions have been noted to be associated with de 22q11.2 including Conotruncal Anomaly Face syndrome (CAFS) (Japan) and sometimes Opitz G/BBB syndrome, CHARGE Association and Cayler-Cardiofacial syndrome.

Screening
All women with a fetus or neonate with a diagnosis of congenital heart defect, especially if conotruncal, can be offered testing (usually FISH) for this deletion. Testing is available from amniocytes or chorionic villi.

Clinical Features
Characteristic facies, cardiovascular defects in 85%, most are VSD. Cleft of secondary palate, may be submucous cleft or velo-pharyngeal incompetence, nasal reflux in infants, transient neonatal hypocalcemia, hypotonia, immune system dysfunction, postnatal growth delay, developmental delay,
learning disability and psychological problems especially in adolescence, hypernasal speech.

**Counseling (Prognosis, Complications, Pregnancy Considerations)**

Clinical features should be reviewed. Early death due to congenital heart defects before 6 months of age in 8%. Early intervention for speech and motor delays. Special education for older children. Chance of psychiatric disorders in 10%. Very variable phenotype, not predictable from laboratory result.

### 5p-(Cri-Du-Chat Syndrome)

**Historic Notes**

In 1963, Lejeune et al. described a syndrome of multiple congenital anomalies, developmental delay, microcephaly, dysmorphic features, and a high-pitched, cat-like cry in infants with deletion of a B group chromosome (Bp-), later identified as 5p-.

**Diagnosis/Definition**

5p-Syndrome is characterized at birth by a high pitched cat-like cry, low birth weight, poor muscle tone, microcephaly. The cry is caused by abnormal laryngeal development. The cry disappears by age 2 years in about one-third of children with 5p-.

A karyotype is needed for the diagnosis. The size of the deletion of the short arm of chromosome 5 is variable and a very small deletion may be missed using conventional G-banding. High resolution studies may be needed or a FISH study using a specific probe for the small deleted area of 5p that is essential for this diagnosis. Microarray would identify this deletion.

**Epidemiology/Incidence**

Estimated prevalence is about 1 in 50,000 live births. Up to 1% of profoundly retarded individuals have 5p-.

**Genetics/Inheritance/Recurrence/Future Prevention**

May be sporadic (80%–85%) if both parents have normal chromosomes with a recurrence risk of less than 1%. In rare cases gonadal mosaicism in one parent may result in a recurrence. If one parent carries a balanced translocation (10%–15%) involving 5p, the recurrence risk is substantially higher.

Most cases have a terminal deletion of 5p. The cat-like cry maps to 5p15.3 and the Cri-du-chat critical region is 5p15.2, which is associated with all the clinical features of the syndrome. The deletion is paternal in origin in 80% of cases.

Affected females are fertile and have a 50% of passing on the deletion to their offspring although none is documented to have reproduced.

**Risk Factors/Associations**

Increased risk if translocation carrier involving 5p.

**Screening**

Amniocentesis, CVS, ultrasound.

**Counseling**

Early feeding problems are common because of swallowing difficulties; poor suck with resultant failure to thrive. Death occurs in 6%–8% of the overall population with Cri-du-chat due to pneumonia, aspiration pneumonia, and congenital heart defects. Survival to adulthood is possible. Children who are raised at home with early intervention and schooling do better than those described in the early literature. Almost all individuals with 5p- have significant cognitive, speech, and motor delays (IQ rarely above 35). Many children can develop some language and motor skills. They may also become independent in self-care skills. Physical features include microcephaly, growth retardation, hypertelorism, epicanthal folds, down-slanting palpebral fissures, round face with full cheeks, flat nasal bridge, down-turned corners of mouth, micrognathia, low-set ears, and variable cardiac defects. Renal anomalies have been described as have cleft lip and palate, talipes equinovarus and gut malrotation. Treatment is symptomatic.

**Helpful Websites**

http://www.emedicine.com/ped/topic504.htm

**REFERENCES**


34. Mujezinovic F, Prosnik A, Alfirevic Z. Different communication strategies for disclosing results of diagnostic prenatal testing. *Cochrane Database of Systematic Reviews*. 2010;11. [Meta-Analysis, 2 RCTs, n = 286]


Carrier screening for inherited genetic conditions

Lorraine Dugoff

KEY POINTS

• The primary goals of carrier screening are to identify individuals and couples who are carriers of genetic mutations that place them at increased risk of having a child with a serious medical disorder and provide them with information to optimize pregnancy outcomes based on their personal values and preferences.

• Most inherited genetic conditions are autosomal recessive. Carriers for these conditions are usually asymptomatic, have no significant family history and are unaware of their carrier status.

• Patients with a positive family history of a genetic condition should be referred for genetic counseling to review the family history, provide accurate information regarding risk, offer the appropriate genetic testing, and discuss reproductive options.

• Information regarding genetic carrier screening should be provided to every pregnant woman.

• Preconception carrier screening is preferred to prenatal carrier screening as it enables couples to consider the most complete range of reproductive options. Information regarding the risk of having an affected child may influence a couple's decision to conceive or to consider the use of donor gametes, preimplantation genetic diagnosis, or prenatal genetic testing.

• Professional practice guidelines currently recommend offering targeted carrier screening for individual conditions based on race or ethnicity, condition severity, carrier frequency, prevalence, detection rates, and residual risk. This approach is limited due to inaccurate or unavailable information regarding ancestry, increased intermixing between different races/ethnicities and the presence of carriers for genetic conditions in the nontargeted populations.

• Technological advances have made it possible to perform expanded carrier screening which involves screening for mutations associated with multiple genetic diseases simultaneously irrespective of ancestry.

• While many practitioners are currently offering expanded carrier screening, professional practice guidelines and recommendations are needed regarding patient and provider education and the conditions that should be included on the screening panels. Future research in expanded carrier screening is needed to assess the impact of this approach on reproductive outcomes.

• Appropriate counseling regarding prognosis, possible complications, long-term issues, and follow-up should be provided to every couple with a pre- or postnatal diagnosis of aneuploidy or other genetic disorders.

• Cystic fibrosis (CF) screening should be offered to all women of reproductive age.

• Individuals of African, southeast Asian, and Mediterranean ancestry should have screening for hemoglobinopathies with a CBC in combination with a hemoglobin electrophoresis.

• Prenatal testing for fragile X syndrome should be offered to known carriers of the fragile X premutation or full mutation. Women with a family history of fragile X-related disorders, unexplained intellectual disability or developmental delay, autism, or premature ovarian insufficiency are candidates for genetic counseling and fragile X premutation carrier screening.

• Preconception and prenatal screening for spinal muscular atrophy (SMA) should be offered to individuals with a family history. Pan-ethnic screening is controversial.

• Carrier screening for Tay–Sachs disease (TSD) (enzyme and DNA), Canavan disease, CF, and familial dysautonomia should be offered to Ashkenazi Jewish (one Jewish grandparent) individuals before conception or during early pregnancy. The availability of genetic carrier screening for mucolipidosis IV, Niemann–Pick disease type A, Fanconi anemia group C, Bloom syndrome, and Gaucher disease should be discussed.

BACKGROUND

Ultrasound (see Chapter 4) as well as prenatal diagnosis and screening for aneuploidy (see Chapter 5) have been reviewed in previous chapters. Carrier screening for inherited genetic conditions is an important component of preconception and prenatal care. There are no trials to assess the downstream effects of any intervention for genetic screening and testing in pregnancy, but clearly there are significant effects. For example, the incidence of TSD in the Ashkenazi Jewish population has decreased by 90% since screening was initiated.

GENETIC COUNSELING

All individuals and couples should have a basic screen for family history of genetic disorders, with a pedigree to at least the second prior generation (three-generation pedigree). Questions should also involve history of birth defects, stillbirth, intellectual disability, developmental delay, recurrent spontaneous pregnancy loss and history of a previous fetus or child who was affected by a genetic disorder. Information regarding genetic carrier screening should be provided to every pregnant woman. Those patients belonging to an ethnic group at increased risk for a recessive condition (e.g., sickle cell—African-American; TSD and others—Jewish; TSD in Irish, Cajun, and French Canadian; α-thalassemia—southeast Asian; and β-thalassemia—Mediterranean) should be offered specific screening (see Chapters 14 and 15 in Maternal-Fetal Evidence...
Genetic carrier screening should be performed on potential gamete donors. Providers may offer an expanded carrier screening panel. Individuals offered expanded carrier screening should have appropriate pretest counseling. It is optimal to offer carrier screening preconception so that women and their partners have the option to consider how much genetic information they would like to have before starting a family. Carrier screening results in the preconception period may influence a couple’s decision to conceive or to consider the use of donor gametes, preimplantation genetic diagnosis, or prenatal genetic testing.

Women with a specific indication for genetic testing should be referred for genetic counseling and a discussion of options available for prenatal diagnosis. Responsibilities of clinicians (e.g., ob-gyns, family medicine specialists, and nurse midwives) caring for pregnant women regarding genetic screening are shown in Table 6.1 [1]. Possible indications for genetic consultation for preconception and prenatal patients are shown in Table 6.2 [2]. Possible indications for genetic consultation for adult patients are shown in Table 6.3 [2].

### COMMON GENETIC DISORDERS AND CARRIER SCREENING GUIDELINES

#### Cystic Fibrosis

CF is an autosomal recessive disorder. The most common mutation is DF508, but >1700 other mutations have been described [3]. The mutation leads to faulty chloride transport, increased sweat chloride levels, and increased thick mucus in lungs, pancreas, biliary tree, and intestines. This is the most common life-limiting genetic disorder in Caucasians (Table 6.4) [3].

CF screening should be offered to all women of reproductive age [3–6]. It is generally more cost effective and practical to initially perform CF carrier screening for the patient only [3]. Nonetheless, screening can be concurrent or sequential, and both strategies are acceptable alternatives.

Concurrent screening:
- Both partners tested simultaneously
- Both partners’ results revealed
- Assesses couple’s risk
- Identifies couples at risk more rapidly
- More precise

Sequential screening:
- Initial screening of one partner
- Other partner tested if first partner is positive
- Low-risk racial/ethnic groups
- Other partner not available

**Interpretation of Results**

- Both partners negative (−/−)—prenatal diagnostic testing not indicated
- One partner carrier (−), one not screened:
  - Intermediate risk—prenatal diagnostic testing not indicated
- One partner carrier (+), one carrier (−):
  - Prenatal diagnostic testing not recommended
- One partner carrier (+), one untested:
  - Partner should be tested if possible genetic counseling
  - Availability/limitations of prenatal testing

#### Table 6.1 Responsibilities of Clinicians Caring for Pregnant Women Regarding Genetic Screening

1. Clinicians should be able to identify patients within their practices who are candidates for genetic testing and should maintain competence in the face of increasing genetic knowledge.
2. Obstetrician–gynecologists should recognize that geneticists and genetic counselors are an important part of the healthcare team and should consult with them and refer as needed.
3. Discussions with patients about the importance of genetic information for their kindred, as well as a recommendation that information be shared with potentially affected family members as appropriate, should be a standard part of genetic counseling.
4. Obstetrician–gynecologists should be aware that genetic information has the potential to lead to discrimination in the workplace and to affect an individual’s insurability adversely. In addition to including this information in counseling materials, physicians should recognize that their obligation to professionalism includes a mandate to prevent discrimination.

**Source:** Modified from American College of Obstetricians and Gynecologists, Obstet Gynecol, 111, 1495–1502, 2008.

#### Table 6.2 Possible Indications for Genetic Consultation for Preconception and Prenatal Patients

<table>
<thead>
<tr>
<th>Either Member of the Couple with</th>
<th>Reason to Consider Consultation</th>
</tr>
</thead>
<tbody>
<tr>
<td>A positive carrier screening test for a genetic condition such as cystic fibrosis</td>
<td>Discuss additional testing strategies and inheritance, testing of partner and siblings</td>
</tr>
<tr>
<td>A personal history of stillbirths, previous child with hydrops, recurrent pregnancy losses, or a child with sudden infant death syndrome (SIDS)</td>
<td>Rule out a chromosomal, metabolic, or syndromic diagnosis associated with an unexplained neonatal death or SIDS</td>
</tr>
<tr>
<td>A progressive neurologic condition known to be genetically determined such as progressive ataxia</td>
<td>Discuss a potential diagnosis, the differential diagnosis, inheritance, and testing options</td>
</tr>
<tr>
<td>A statin-induced myopathy</td>
<td>Discuss a potential mitochondrial disorder, inheritance, and testing options</td>
</tr>
<tr>
<td>A family or personal history of: A birth defect such as cleft lip</td>
<td>Reason to consider consultation</td>
</tr>
<tr>
<td>A chromosomal abnormality such as translocation</td>
<td>Discuss recurrence risks and testing options</td>
</tr>
<tr>
<td>Significant hearing or vision loss thought to be genetically determined</td>
<td>Discuss risks to the fetus and testing options</td>
</tr>
<tr>
<td>Intellectual disability, developmental delay, or autism</td>
<td>Discuss risks to the fetus and testing options</td>
</tr>
</tbody>
</table>

**Source:** Modified from Pletcher BA et al., Genet Med, 9, 385–389, 2007.
Both partners carrier (+):
- 25% Chance of having an affected offspring
- Genetic counseling
- Prenatal diagnosis offered (chorionic villus sampling [CVS], amniocentesis)
- Counseling regarding continuation versus termination of pregnancy for affected pregnancies

Risk of affected offspring depends also on prevalence of carrier status in the specific ethnic group (Table 6.4).

If a patient or the patient's partner has a family history of CF, medical record review should be performed to identify of cystic fibrosis transmembrane conductance regulator (CFTR) mutation analysis in the affected family member is available. If a woman's reproductive partner has apparently isolated congenital bilateral absence of the vas deferens, the couple should be referred for a genetics consultation to discuss potential clinical implications and mutation analysis.

**Hemoglobinopathies**

Hemoglobinopathies are a diverse group of inherited single-gene disorders that result from variations in the structure and/or function of hemoglobin. The most common hemoglobinopathies, sickle cell disease, β-thalassemia and α-thalassemia, are all autosomal recessive conditions. The American College of Obstetricians and Gynecologists (ACOG) recommends carrier screening for individuals of African, southeast Asian, and Mediterranean ancestry. A complete blood count (CBC) in combination with a hemoglobin electrophoresis is recommended for screening individuals of African ancestry. A CBC with red cell indices is the initial recommended screening test for individuals of southeast Asian, and Mediterranean ancestry. Individuals with a low mean

---

**Table 6.3** Possible Indications for Genetic Consultation for Adult Patients

<table>
<thead>
<tr>
<th>Personal History of</th>
<th>Reason to Consider Consultation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abnormal sexual maturation or delayed puberty</td>
<td>Rule out an intersex condition, chromosomal abnormality, or syndromic diagnosis</td>
</tr>
<tr>
<td>Recurrent pregnancy losses (more than two)</td>
<td>Rule out a chromosomal rearrangement such as a balanced translocation with karyotype</td>
</tr>
<tr>
<td>Tall or short stature for genetic background</td>
<td>Rule out a skeletal dysplasia, chromosomal or syndromic diagnosis</td>
</tr>
<tr>
<td>One or more birth defects</td>
<td>Rule out a chromosomal or syndromic diagnosis, and provide genetic counseling</td>
</tr>
<tr>
<td>Six or more cafe-au-lait macules &gt;1.5 cm in diameter</td>
<td>Rule out neurofibromatosis Type 1</td>
</tr>
<tr>
<td>Statin-induced myopathy</td>
<td>Rule out a mitochondrial disorder</td>
</tr>
<tr>
<td>A cancer or cancers known to be associated with specific genes or mutations in the context of a compelling family history; young age at onset, bilateral lesions, and familial clustering of related tumors</td>
<td>Rule out a mutation in a causative or contributory gene; discuss surveillance, treatment, testing options, and inheritance</td>
</tr>
<tr>
<td>Cardiovascular problems known to be associated with genetic factors such as cardiomyopathy</td>
<td>Rule out a syndrome (Marfan syndrome); discuss surveillance, treatment, testing options, and inheritance</td>
</tr>
<tr>
<td>Suspected genetic disorder affecting connective tissue</td>
<td>Confirm or rule out suspected diagnosis; discuss treatment, testing options, and inheritance</td>
</tr>
<tr>
<td>Hematologic condition associated with excessive bleeding or excessive clotting</td>
<td>Rule out a syndrome (Stickler syndrome); discuss testing options, if applicable, and inheritance</td>
</tr>
<tr>
<td>Progressive neurologic condition known to be genetically determined such as unexplained myopathy</td>
<td>Rule out a syndrome or nonsyndromic genetic form of hearing loss; discuss surveillance, testing options, and inheritance</td>
</tr>
<tr>
<td>Visual loss known to be associated with genetic factors such as retinitis pigmentosa</td>
<td>Confirm the diagnosis, discuss prognosis, medical management, and inheritance</td>
</tr>
<tr>
<td>Early-onset hearing loss</td>
<td>Rule out a genetic condition associated with this history, discuss diagnosis, inheritance, recurrence risks, and identify syndromes, when possible</td>
</tr>
<tr>
<td>Recognized genetic disorder including a chromosomal or single-gene disorder</td>
<td>Rule out the diagnosis, genetic counseling, and inheritance</td>
</tr>
<tr>
<td>Mental illness such as schizophrenia</td>
<td>Rule out a genetic condition associated with this history, discuss diagnosis, inheritance, recurrence risks, and identify syndromes, when possible</td>
</tr>
<tr>
<td>A close relative with a sudden, unexplained death, particularly at a young age</td>
<td>Rule out the diagnosis, genetic counseling, and inheritance</td>
</tr>
</tbody>
</table>


**Table 6.4** Incidence and Carrier Risk for CF Based on Race or Ethnicity

<table>
<thead>
<tr>
<th>Racial or Ethnic Group</th>
<th>Detection Rate (%)</th>
<th>Estimated Carrier Risk Before Test</th>
<th>Estimated Carrier Risk After a (−) Test</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ashkenazi Jewish</td>
<td>94</td>
<td>1/24</td>
<td>~1/380</td>
</tr>
<tr>
<td>Non-Hispanic white</td>
<td>88</td>
<td>1/25</td>
<td>~1/200</td>
</tr>
<tr>
<td>Hispanic white</td>
<td>72</td>
<td>1/58</td>
<td>~1/200</td>
</tr>
<tr>
<td>African-American</td>
<td>64</td>
<td>1/61</td>
<td>~1/170</td>
</tr>
<tr>
<td>Asian-American</td>
<td>49</td>
<td>1/94</td>
<td>~1/180</td>
</tr>
</tbody>
</table>


**Abbreviation:** CF, cystic fibrosis.
corpuscular volume (MCV) and normal iron status should have a hemoglobin electrophoresis. Southeast Asians with a normal hemoglobin electrophoresis should have molecular testing to rule out α-globin gene deletions characteristic of α-thalassemia. Couples determined to be at increased risk for having a child with sickle cell disease or thalassemia should be referred to a genetic counselor [7]. For additional information regarding hemoglobinopathies, refer to Chapters 14 and 15 in Maternal-Fetal Medicine Evidence Based Guidelines.

Fragile X Syndrome

Diagnosis/Definition
Most common inherited cause of intellectual disability; caused by expansion of a repetition of the cytosine–guanine–guanine (CGG) trinucleotide segment of the fragile X mental retardation 1 (FMR-1) gene that is located on the X chromosome at Xq27.3.

The diagnosis is based on molecular genetic analysis. Most laboratories use a combination of two approaches: (1) Southern blot analysis to measure the degree of methylation and (2) polymerase chain reaction (PCR) to discriminate small differences in the intermediate and premutation sizes. Fragile X syndrome occurs when expansion of the CGG repeat occurs.

Epidemiology/Incidence
Both males and females can be affected. The prevalence is 1 in 3600 males and 1 in 4000–6000 females.

Genetics/Inheritance/Recurrence
The FMR-1 gene is typically comprised of a limited number of CGG repeats. This region is usually stable, relatively small (<45 repeats), and passes from one generation to the next. However, during meiosis in oocytes, the region can expand, reaching either intermediate (45–54 repeats) or premutation (55–200 repeats) lengths. As repeat size increases, stability decreases and further increases in the number of repeats become likely. Once the premutation reaches expansion to >90 repeats, the likelihood of expansion to a full mutation is at least 80% (Table 6.5) [8].

Only women with a premutation FMR-1 can pass the full mutation to their offspring. Fathers with premutations usually pass the gene in a stable fashion to all of their daughters and none of their sons and will have affected grandsons. Premutation male and female carriers are at risk for fragile X-associated tremor/ataxia syndrome (FXTAS) usually after age 50 years [9].

Table 6.5  Fragile X: Risk of Full Mutation (and Therefore Affected Child) Based on Number of Maternal CGG Repeats

<table>
<thead>
<tr>
<th>No. Maternal CGG Repeats</th>
<th>% Risk Expansion to Full Mutation</th>
</tr>
</thead>
<tbody>
<tr>
<td>55–59</td>
<td>4</td>
</tr>
<tr>
<td>60–69</td>
<td>5</td>
</tr>
<tr>
<td>70–79</td>
<td>31</td>
</tr>
<tr>
<td>80–89</td>
<td>58</td>
</tr>
<tr>
<td>90–99</td>
<td>80</td>
</tr>
<tr>
<td>100–200</td>
<td>98</td>
</tr>
</tbody>
</table>

Abbreviation: CGG, cytosine–guanine–guanine.

Spinal Muscular Atrophy

Diagnosis/Definition
SMA refers to a group of diseases that affect the motor neurons of the spinal cord and brainstem, which are responsible for supplying electrical and chemical signals to muscle cells. Without proper signals, muscle cells do not function properly and thus become much smaller (atrophy). This leads to muscle weakness. Individuals affected with SMA have progressive muscle degeneration and weakness, eventually leading to death.

Incidence
The incidence is 1/10,000 infants.

Genetics/Classification
There are several forms of SMA, depending on the age of onset and the severity of the disease. Two genes, SMN1 and SMN2, have been linked to SMA types I, II, III, and IV. Type I is the most severe form of SMA and characterized by muscle weakness present from birth, often manifested by difficulties with breathing and swallowing, and death usually by age 2–3 years. Type II has onset of muscle weakness after 6 months of age and can obtain some early physical milestones like sitting without support. Type III is a milder form of SMA, with onset of symptoms after 10 months of age. Individuals with Type III SMA often achieve the ability to walk but may have frequent...
SMA is most often caused by a deletion of a segment of DNA, in exon 7 and exon 8, in the SMN1 gene located on chromosome 5. Rarely, SMA is caused by a point mutation in the SMN1 gene.

**Screening**

The American College of Medical Genetics and Genomics (ACMG) recommends pan-ethnic screening for SMA while the ACOG only recommends screening when a family history is present [10,11]. Approximately 1 in 50 individuals is a carrier for SMA.

Carrier testing for SMA measures the number of copies of the deleted segment in the SMN1 gene. A noncarrier is expected to have two copies present (no deletion), while a carrier will have only one copy present (a deletion of one copy). SMA carriers most commonly have one functional SMN1 gene on one chromosome and an SMN1 gene deletion on the other chromosome. Carriers can also have two functional SMN1 gene copies on one chromosome (in cis) and none on the other chromosome (the ‘2 + 0’ genotype), or one chromosome with a nonfunctional SMN1 gene with a point mutation or a microdeletion. Since carrier testing relies on a PCR-based gene-dose approach to determine SMN1 copy number, this method will not identify carriers with the ‘2 + 0’ genotype. Carrier testing will also not identify point mutations. Approximately 90% of SMA carriers in the Ashkenazi Jewish population can be identified with this PCR-based testing method. A systematic review and meta-analysis including 169,647 SMA carrier tests in 14 studies reported an 87%–95% detection rate of SMA carrier testing in a non-Black population and a 71% sensitivity in the Black population due to a relatively high incidence of the ‘2 + 0’ genotype in the black population. The positive predictive value of carrier testing when one copy of the SMN1 gene is identified is 100%, regardless of the population tested. There is a 5%–13% false-negative rate in non-Black individuals with a two-copy SMN1 result while there is a 29% false-negative rate in Black individuals with this result. Individuals with three SMN1 copies are almost always noncarriers [12].

**ASHKENAZI JEWISH ANCESTRY GENETIC SCREENING**

**Who to Screen**

The family history of individuals considering pregnancy, or who are already pregnant, should be included when performing carrier screening in individuals of Ashkenazi Jewish descent. The ACOG recommends that carrier screening for TSD, Canavan disease, CF, and familial dysautonomia should be offered to Ashkenazi Jewish individuals before conception or during early pregnancy so that a couple has an opportunity to consider prenatal diagnostic testing options. If the woman is already pregnant, it may be necessary to screen both partners simultaneously so that the results are obtained in a timely fashion to ensure that prenatal diagnostic testing is an option. ACOG acknowledges that carrier screening is available also for mucolipidosis IV, Niemann–Pick disease type A, Fanconi anemia group C, Bloom syndrome, and Gaucher disease. The ACMG recommends screening for these five additional disorders, for a total of nine [14]. ACMG provides criteria for adding new diseases to the panel of Jewish genetic diseases based on significant burden of the disorder, carrier rate >1/100 and/or test sensitivity >90%. Patient education materials can be made available so that interested patients can make an informed decision about having additional screening tests. Some patients may benefit from genetic counseling.

There are currently 19 diseases that are being made available for screening on Jewish genetic disease panels. In addition to those mentioned above, the following diseases are included in the new panels: maple syrup urine disease, glycogen storage disease type 1a, dihydrolipoamide dehydrogenase (DLD) deficiency, familial hyperinsulinism, Joubert syndrome, nemaline myopathy, SMA, Usher syndrome type IF, Usher syndrome type III, and Walker–Warburg syndrome [15].

**How to Screen**

All the disorders on the Ashkenazi Jewish genetic disease panel are autosomal recessive. A carrier is unaffected but if two carriers of a mutation in the same gene have a child together, they have a 25% risk with each pregnancy of having an affected child (Figure 6.1).

When only one partner is of Ashkenazi Jewish descent, that individual should be screened first. If it is determined that this individual is a carrier, the other partner should be offered screening for that disorder. However, the couple should be informed that the carrier frequency and

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**Figure 6.1** Autosomal recessive inheritance: where both parents are carriers, there is a one in four chance a child will be free of disease and not a carrier, a one in two chance a child will be a carrier, and a one in four chance a child will have the disease.
the detection rate in non-Jewish individuals are unknown for many of these disorders. Therefore, it is difficult to accurately predict the couple’s risk of having a child with the disorder, and gene sequencing may be necessary to determine whether a non-Jewish partner is a carrier. Referral to a genetic counselor is recommended in these cases.

Individuals with a positive family history of one of these disorders should be offered carrier screening for the specific disorder (and others for this group) and may benefit from genetic counseling.

When both partners are carriers of one of these disorders, they should be referred for genetic counseling and offered prenatal diagnosis. Carrier couples should be informed of the disease manifestations, range of severity, and available treatment options. Prenatal diagnosis by DNA-based testing can be performed on cells obtained by CVS and amniocentesis.

When an individual is found to be a carrier, his or her relatives are at risk for carrying the same mutation. The patient should be encouraged to inform his or her relatives of the risk and the availability of carrier screening. The provider should not contact these relatives because there is no provider-patient relationship with the relatives, and confidentiality must be maintained [13].

SELECT DISORDERS
Carrier frequencies apply to the Ashkenazi Jewish population [15].

**Tay–Sachs Disease**
Carrier frequency is 1 in 30 in the Ashkenazi Jewish population. The carrier frequency is also increased in individuals of French Canadian and Cajun descent.

*Clinical Features*
The most well-known of these diseases. An apparently healthy child begins to lose skills around 4–6 months of age, and there is a progressive neurological decline leading to blindness, seizures, and unresponsiveness. Death usually occurs by the age of 4–6 years.

*Testing*
Carrier testing should be performed using DNA-based testing and hexosaminidase enzymatic activity testing [15]. The enzyme assay detects all carriers, regardless of ethnicity. DNA-based testing detects 94% of carriers and hexosaminidase A enzyme testing detects 98% of carriers in the Ashkenazi Jewish population [13]. Enzyme testing in pregnant women and women taking oral contraceptives should be performed using leukocyte testing as serum testing is associated with an increased false-positive rate in these populations.

**Familial Dysautonomia**
Carrier frequency is 1/32 with disease incidence of 1/3700.

*Clinical Features*
Causes the autonomic and sensory nervous system to malfunction, affecting the regulation of body temperature, blood pressure, and stress response, and causes decreased sensitivity to pain. Frequent pneumonia and poor growth may occur. Survival into adulthood is possible but with careful medical care.

**Canavan Disease**
Carrier frequency is 1 in 40 with a disease incidence of 1 in 3000 to 1 in 6000.

*Clinical Features*
Appear normal at birth; then progressive loss of skills, hypotonia, macrocephaly, and spasticity. Most die within first year of life but may survive to second decade.

*Testing*
Carrier testing detects 98% carriers.

**Bloom Syndrome**
Carrier frequency is 1/100 with a disease incidence of 1/48,000.

*Clinical Features*
Short stature, a sun-sensitive skin rash, an increased susceptibility to infections and higher incidence of leukemia and other cancers, infertility, and immunodeficiency.

*Testing*
Carrier testing detects 95%–97% of carriers.

**Fanconi Anemia Type C**
Carrier frequency is 1/89 with disease incidence of 1/32,000.

*Clinical Features*
Associated with short stature, bone marrow failure, and a predisposition to leukemia and other cancers. Some children have limb, heart, or kidney abnormalities and learning difficulties.

*Treatment*
Pancytopenia can be treated with stem cell transplantation.

*Testing*
Carrier testing detects 99% of carriers.

**Gaucher Disease**
Carrier rate is 1/15 with disease incidence of 1/900.

*Clinical Features*
A variable condition both in age of onset and symptoms. It may present with a painful, enlarged spleen, anemia, and low white blood cell count. Bone deterioration is a major cause of pain and disability.

*Treatment*
Treatment is available.

**Mucolipidosis Type IV**
Carrier frequency is 1/127.
Clinical Features
A progressive neurological disorder with variable symptoms beginning in infancy. Characteristics include muscle weakness, mental retardation, and eye problems—both corneal clouding and retinal abnormalities develop.

Treatment
No treatment is available [15].

Carrier Testing
95% detection rate.

Glycogen Storage Disease, Type 1a
Carrier frequency is 1 in 64.

Clinical Features
A metabolic disorder that causes poor blood sugar maintenance with sudden drops in blood sugar, growth failure, enlarged liver, and anemia.

Treatment
Disease management involves lifelong diet modification.

Carrier Testing
95% detection rate.

Maple Syrup Urine Disease
Carrier frequency is 1 in 97.

Clinical Features
A variable disorder of amino acid metabolism. Named for the characteristic maple syrup smell of urine in those with the disorder.

Treatment
With careful dietary control, normal growth and development are possible. If untreated, it can lead to poor feeding, lethargy, seizures, and coma.

Carrier Testing
95% detection rate.

Niemann–Pick Disease Type A
Carrier frequency is 1 in 90.

Clinical Features
A progressive neurodegenerative disease characterized by hepatosplenomegaly, leading to death by age 2–4 years.

Carrier Testing
95% detection rate.

DLD Deficiency
Carrier frequency is 1 in 107.

Clinical Features
Presents in early infancy with poor feeding, frequent episodes of vomiting, lethargy, and developmental delay. Affected individuals develop seizures, enlarged liver, blindness, and ultimately suffer an early death.

Carrier Testing
>95% detection rate.

Familial Hyperinsulinism
Carrier frequency is 1 in 68.

Clinical Features
Characterized by hypoglycemia that can vary from mild to severe. It can be present in the immediate newborn period through the first year of life with symptoms such as seizures, poor muscle tone, poor feeding, and sleep disorders.

Carrier Testing
90% detection rate.

Joubert Syndrome
Carrier frequency is 1 in 92.

Clinical Features
Characterized by structural mid- and hindbrain malformations. Affected individuals have mild to severe motor delays, developmental delay, decreased muscle tone, abnormal eye movements, and characteristic facial features.

Carrier Testing
Detection rate is unknown (not enough data).

Nemaline Myopathy
Carrier frequency is 1 in 168.

Clinical Features
Most common congenital myopathy. Infants are born with hypotonia and usually have problems with breathing and feeding. Later some skeletal problems may arise, such as scoliosis. In general, the weakness does not worsen during life but development is delayed.

Carrier Testing
95% detection rate.

Usher Syndrome Type 1F
Carrier frequency is 1 in 147.

Clinical Features
Characterized by profound hearing loss, which is present at birth, and adolescent-onset retinitis pigmentosa, a disorder that significantly impairs vision.

Carrier Testing
≥75% detection rate.

Usher Syndrome Type III
Carrier frequency is 1 in 120.

Clinical Features
Causes progressive hearing loss and vision loss. Hearing is often normal at birth with progressive hearing loss typically beginning during childhood or early adolescence. Often leads to blindness by adulthood.

Carrier Testing
95% detection rate.

Walker–Warburg Syndrome
The carrier frequency in the Ashkenazi population for one Ashkenazi founder mutation is 1 in 149.
Clinical Features
It is a muscle, eye, and brain syndrome. It is a severe condition that presents with muscle weakness, feeding difficulties, seizures, blindness, brain anomalies, and delayed development. Life expectancy is below 3 years.

Carrier Testing
Detection rate is unknown (not enough data).

EXPANDED CARRIER SCREENING
Expanded carrier screening involves simultaneous screening for multiple genetic conditions regardless of a patient’s race/ethnicity. Technological advances have made it possible to efficiently screen for a large number of conditions simultaneously for the same cost as one or two single-gene carrier tests. Expanded carrier screening has a number of advantages compared with the targeted screening approach. It can overcome the limitations associated with the targeted ethnicity-based screening approach including inaccurate and unavailable information regarding ancestry, increased intermixing between different races/ethnicities and the presence of carriers for genetic conditions in the nontargeted populations. Data from expanded carrier screening on 23,453 individuals from practices in the United States demonstrated that diseases currently recommended for screening only in certain populations, such as Canavan disease and sickle cell disease, are widely distributed outside of their targeted populations. Twenty-four percent of individuals were identified as carriers of at least one mutation [16]. Future research in expanded carrier screening is needed to assess the impact of this approach on reproductive outcomes. A 2012 survey of ACOG Fellows indicated that 15% of the responders offered expanded carrier screening to all of their patients and 52% provided this screening upon patient request [17].

Practice guidelines from national societies are needed to recommend conditions that should be included on expanded carrier panels and provide guidance to practitioners regarding pretest counseling. Since expanded carrier screens can include over 100 conditions, it is not feasible to counsel patients regarding the specific conditions on the panel. Table 6.6 lists points to include in the pretest counseling and consent process. If a patient and her partner both test positive as carriers for the same autosomal recessive genetic condition, they should be referred to a certified genetics professional to discuss the implications and reproductive options [19,20].

DIRECT-TO-CONSUMER GENETIC TESTING
Direct-to-consumer (DTC) genetic testing differs from traditional genetic testing in that consumers order tests and receive test results without an independent medical provider serving as an intermediary. Some DTC companies offer genetic counseling (generally by telephone), whereas others do not. These tests are typically advertised and sold over the Internet. DTC is permitted in about half the states in the United States and is subject to little oversight at the federal level. Internationally, several countries have issued reports cautioning against its use, and several European countries have banned or are considering banning it entirely. The American Society of Human Genetics has issued guidelines for transparency, provider education, and test and laboratory quality on DTC genetic testing [18]. The ACMG suggests the following regarding DTC genetic testing (www.acmg.net):

- A knowledgeable professional should be involved in the process of ordering and interpreting a genetic test.
- The consumer should be fully informed regarding what the test can and cannot say about his or her health.
- The scientific evidence on which a test is based should be clearly stated.
- The clinical testing laboratory must be accredited by Clinical Laboratory Improvement Amendments (CLIA), the State, and/or other applicable accrediting agencies.
- Privacy concerns must be addressed.

Table 6.6 Discussion Points for Pretest Counseling and Consent Process for Expanded Carrier Screening

1. Carrier screening is voluntary. Patients can choose to participate or decline.
2. Genetic testing results are confidential and protected in employment and health insurance by the Genetic Information Non-Discrimination Act of 2008.
3. Expanded carrier screening panels include conditions that may vary in severity. The panels include many conditions that are associated with significant adverse outcomes including decreased life expectancy, cognitive impairment and the need for medical and/or surgical intervention.
4. Accurate knowledge of paternity is necessary to provide accurate information regarding risk. DNA (or blood) from the biological father is necessary in order to provide risk assessments for autosomal recessive conditions.
5. There is a residual risk for having an affected offspring even if both partners screen negative. This risk may vary depending on ethnicity, the specific condition and the laboratory performing the testing.
6. It is common to identify carriers for one or more conditions when using expanded screening panels. In most cases, being a carrier has no significant medical consequences for the individual. If two partners are carriers of different autosomal recessive conditions, offspring are not likely to be affected.
7. It is possible that carrier screening will determine that an individual has two pathogenic variants for a condition and thus has an autosomal recessive condition that might affect their health. Expanded carrier screening panels that include autosomal dominant and X-linked conditions may detect individuals affected with one of these conditions. In these situations, individuals should be referred for genetic counseling and appropriate medical management.

REFERENCES


Before labor and first stage of labor

Serena Xodo

KEY POINTS

• Before labor
  • Prediction of spontaneous labor is imprecise, with best evidence for transvaginal ultrasound (TVU) cervical length (CL) having good predictive accuracy for spontaneous labor within 7 days.
  • If nonvertex presentation, perform cesarean delivery (CD) at 39 0/7–39 6/7 weeks. For timing and indications of other planned cesarean or vaginal deliveries, see Chapters 13 and 56 in Maternal-Fetal Evidence Based Guidelines.
  • If woman is candidate for vaginal delivery and ≥41 weeks, start induction (see Chapter 21).
  • Maternal antenatal training to prepare for labor and delivery (L&D) is associated with arriving to L&D ward more often in active labor, less visits to the labor suite, and using less epidural analgesia.
  • X-ray pelvimetry should not be performed as it is associated with no benefits, and it increases incidence of CDs. There is insufficient data regarding MRI or clinical pelvimetry.
  • There is limited evidence to assess the safety and efficacy of planned home birth for low-risk pregnant women. Compared with planned hospital birth, planned home birth is associated with a higher risk of neonatal deaths (0.2% vs. 0.09%), low Apgar scores, and neonatal seizures. The hospital is the safest setting for L&D.
  • A birth center in the hospital is a safe location for birth. The trend for a 67% higher perinatal mortality should be weighted during counseling against the significant 4% increase in spontaneous vaginal delivery (SVD) and 96% higher satisfaction.
  • Most women (those without risk factors) should be offered midwife-led models of care, as midwife-led care is associated with a lower incidence of preterm birth, regional analgesia, instrumental vaginal birth, and fetal/neonatal death, and with shorter labor and a higher incidence of spontaneous vaginal birth. So women should be encouraged to ask for this option.
  • Training of traditional birth assistants in middle- and low-income countries is associated with a trend for less maternal mortality and significantly less perinatal mortality.
  • Delayed hospital admission until active labor (regular painful contractions and cervical dilatation >3 cm) is associated with less time in the labor ward, less intrapartum oxytocics, and less analgesia.
  • Admission tests such as fetal heart rate (FHR) tracing and amniotic fluid assessment have not been associated with any benefit.
  • Routine enema is not recommended.
  • Perineal shaving is not recommended.
  • Vaginal chlorhexidine irrigation is not recommended.
  • Universal prenatal maternal screening with anovaginal specimen at 35–37 weeks and intrapartum antibiotic treatment are the most efficacious of the current strategies for prevention of neonatal early-onset group B streptococcus (GBS) disease.
  • All women should have continuous, one-on-one support throughout labor and birth (e.g., doula), as it is associated with less intrapartum analgesia, cesarean birth, operative birth, and dissatisfaction with the childbirth experiences, and more spontaneous vaginal birth.
  • There is insufficient evidence for providing nutritional recommendations for women in labor. There is little justification for the restriction of fluids and food in labor for women at low risk of complications.
  • Intravenous (IV) fluids at 250 mL/hour are associated with shorter duration of labor and less cesarean deliveries compared with 125 mL/hour.
  • Upright positions (either standing, sitting, kneeling, or walking around) in the first stage of labor reduce the length of labor by approximately over 1 hour and are associated with less epidural analgesia. Since walking does not seem to have a beneficial or detrimental effect on L&D, women can choose freely to walk or lay in bed, preferably upright, during labor, whichever is more comfortable for them.
  • Water immersion during the first stage of labor reduces the use of analgesia and by about 30 minutes the duration of the first stage of labor, without adverse maternal or neonatal outcomes.
  • Routine early (or even late) amniotomy cannot be recommended as part of standard labor management and care.
  • The use of the partogram cannot be recommended as a routine intervention in labor.
  • There is insufficient evidence to recommend any particular frequency of vaginal cervical examinations in labor. Most studies, including those with active management, perform cervical examinations every 2 hours in active labor, but the risk of chorioamnionitis increases with increasing number of examinations.
  • For women making slow progress in the first stage of spontaneous labor, the use of oxytocin augmentation is associated with a reduction in the time to delivery of approximately 2 hours.
  • The individual interventions that are part of active management of labor should be studied separately, and only those that are beneficial (e.g., support by doula) implemented.
  • Dystocia cannot be diagnosed unless rupture of membranes (ROM) has occurred, and adequate oxytocin to
achieve at least three to five adequate contractions per hour has been instituted. Dystocia also cannot be diagnosed reliably before the first stage of labor has entered the active phase, which has been defined, especially in nulliparous with epidurals in place, as at least 6 cm of cervical dilatation. Before performing a CD for active phase labor arrest, labor should be arrested for a minimum of 4 hours (if uterine activity is greater than 200 Montevideo units as documented with intrauterine pressure catheter [IUPC]) or 6 hours (if greater than 200 Montevideo units could not be sustained).

INTRODUCTION
For management of induction, meconium, oligo/polyhydramnios, intrapartum monitoring (including amnioinfusion for variables), operative vaginal delivery, shoulder dystocia, trial of labor after cesarean (TOLAC), intrauterine growth restriction (IUGR), macrosomia, abnormal third stage, etc., see appropriate distinct guidelines in this book and its companion, Maternal-Fetal Evidence Based Guidelines. This chapter discusses prediction of the onset of spontaneous labor, and especially interventions before labor and in the first stage of labor that can influence L&D outcomes.

PREDICTION OF THE ONSET OF SPONTANEOUS LABOR
Weather Effects
There is no strong association between changes in barometric pressure and onset of labor [1].

Time of Day
Diurnal rhythms seem to show a higher rate of starting labor in the evening and night hours [2].

Cervical Status
Both the Bishop score and CL measure usually by TVU have been evaluated for their predictive accuracy for the onset of spontaneous labor. TVU CL is associated with very good prediction of onset of spontaneous labor: for example, the chance of labor within 7 days is about 10% for a woman with a TVU CL of 40–45 mm at about 37–39 weeks, and about 90% if TVU CL is about 10 mm at about 37–39 weeks [3].

BEFORE LABOR
Antenatal Classes
Antenatal classes to improve the birth process have been studied in at least three randomized controlled trials (RCTs).

Compared with no such training, 9 hours of antenatal training to prepare for L&D are associated with arriving to L&D ward more often in active labor (relative risk [RR] 1.43, 95% confidence interval [CI] 1.26–1.65) and using less epidural analgesia (RR 0.84, 95% CI 0.73–0.97) [4].

Compared with no such education, antenatal education focusing on natural childbirth preparation with training in breathing and relaxation techniques is not associated with any effects on maternal or perinatal outcomes, including similar incidences of epidural analgesia, childbirth, or parental stress, in nulliparous women and their partners [5].

In a small RCT, specific antenatal education program is associated with a reduction in the mean number of visits to the labor suite before the onset of labor [6]. It is unclear whether this results in fewer women being sent home because they are not in labor.

Fear of Childbirth
Fear of childbirth is associated with a 47 minutes longer duration of labor, compared with no fear of childbirth [7]. Intensive counseling in women with fear of childbirth reduces anxiety and concerns related to pregnancy and birth, and is associated with shorter labors [8].

Nonvertex Presentation
If nonvertex presentation is detected, CD at 39 0/7–39 6/7 weeks is recommended (see Chapter 24). For timing and indications of other planned cesarean or vaginal deliveries, see Chapters 13, and 56 in Maternal-Fetal Evidence Based Guidelines.

Late-Term Gestation
Induction advised at ≥41 weeks (see Chapter 27).

Pelvimetry
There is insufficient data regarding MRI or clinical pelvimetry.

Interventions to Expedite the Onset of Labor
For acupuncture, breast stimulation, castor oil, enemas and baths, homeopathy, sexual intercourse, as well as other non-pharmacologic and pharmacologic techniques to expedite the onset of labor, or for induction, see Chapter 21.

SITE OF LABOR MANAGEMENT
Planned Home Birth
There is limited evidence to assess the safety and efficacy of planned home birth for low-risk pregnant women as there is only one RCT with only 11 women ever published [10,11]. In healthy low-risk women, compared with planned hospital birth, planned home birth is associated with a higher risk of neonatal deaths (0.2% vs. 0.09%) in a meta-analysis of nonrandomized studies including over 500,000 births [12]. In a large (>79,000 women with singletons, term, cephalic, non-anomalous pregnancies) retrospective cohort study, planned out-of-hospital births were associated with a higher rate or perinatal deaths (3.9 vs. 1.8 deaths per 1000 deliveries; OR 1.52, 95% CI 0.51–2.54) than planned in-hospital deliveries, as well as more neonatal seizures [13]. Data from U.S. births showed a significantly increased risk of neonatal deaths, low Apgar scores, and neonatal seizures associated with home births or births in free standing birth centers, compared with hospital births [14,15]. Some of these studies included free-standing birth centers with home births. This means, therefore, that the hospital is the safest setting for L&D [16]. A home birth service ought to be backed up by a modern hospital system. There are diverging opinions even in Western countries, with about 30% of Dutch births occurring at home, versus <1% of U.S. births. In the United States, about 75% of home births are delivered by noncertified midwives. In the Netherlands, travel time of >20 minutes from home to
hospital is associated with increased risk of perinatal mortality and other adverse outcomes [17]. Women with risk factors for abnormal outcome should deliver in a hospital setting. All women should be aware of possible maternal and fetal risks, including severe morbidity and mortality, associated with L&D even in low-risk women, and should be aware of the absence of intensive care and operative capabilities in the home setting [10]. Inference from results of “home-like” versus conventional ward setting (see below) should warn against home birth.

Ethical experts agree that hospital birth is safer. Some medical ethics experts emphasize that physicians should counsel strongly against home birth [18]. Some other medical ethics experts argue instead that, given the increased risk of perinatal mortality is about 1–2/1000 in home compared with hospital birth, women should be counseled on these absolute numbers and allowed a choice based on autonomy [19].

**Freestanding Birth Center**

There are no RCTs of freestanding birth centers, and therefore the evidence is insufficient to recommend this setting [20]. For non-RCT evidence, see above, under “Planned home births.”

**Planned Hospital Birth: Alternative Setting (Home-Like, e.g., Hospital Birth Center) versus Conventional Hospital Ward Setting**

Several RCTs have evaluated the option of “alternative setting” versus conventional hospital ward setting for L&D. For alternative setting, most trials included care by midwives in a location in the hospital, close to the regular L&D ward, that did not look like a usual L&D setting. There are no RCTs of freestanding birth centers or Snoezelen rooms. Some RCTs randomized women in labor, while others at the beginning of pregnancy. So continuity of care was usually higher in the alternative setting group. Usually about 40%–50% of patients randomized to alternative setting (often about 60% for nulliparous women, 20%–30% for multiparous women) need to be moved in labor to the conventional L&D ward.

Compared with conventional hospital ward, allocation to an alternative hospital setting increased the likelihood of the following: no intrapartum analgesia/anesthesia (RR 1.18, 95% CI 1.05–1.33); SVD (RR 1.03, 95% CI 1.01–1.05); breastfeeding at 6–8 weeks (RR 1.04, 95% CI 1.02–1.06); and very positive views of care (RR 1.96, 95% CI 1.78–2.15). Allocation to an alternative setting decreased the likelihood of epidural analgesia (RR 0.80, 95% CI 0.74–0.87); oxytocin augmentation of labor (RR 0.87, 95% CI 0.67–0.88); and episiotomy (RR 0.83, 95% CI 0.77–0.90) [13]. There was no apparent effect on maternal morbidity and mortality, serious perinatal morbidity/mortality (RR 1.17, 95% CI 0.51–2.67), perinatal mortality (RR 1.67, 95% CI 0.93–3.00), other adverse neonatal outcomes, or postpartum hemorrhage. The 4% increase in SVD may be secondary to less epidural anesthesia, which may in turn be secondary to less availability in home-like settings, and/or to less intrapartum monitoring. The trend for a 67% higher perinatal mortality should be weighted against the significant 4% increase in SVD and 96% higher satisfaction, during counseling. Episiotomy should be avoided in general (see Chapter 8). No firm conclusions can be drawn regarding the effects of variations in staffing, organizational models, or architectural characteristics of the alternative settings [20]. A birth center in the hospital is a safe location for birth [16].

**PROVIDER Midwife-Led Care**

In many parts of the world, midwives are the primary providers of care for childbearing women. Elsewhere it may be medical doctors or family physicians who have the main responsibility for care, or the responsibility may be shared. The underpinning philosophy of midwife-led care is normality, continuity of care, and being cared for by a known and trusted midwife during labor. There is an emphasis on the natural ability of women to experience birth with minimum intervention [21].

The effect of midwife-lead care compared with physician-led care or to other provider-led care has been evaluated mostly for the whole pregnancy, including together both antepartum care and care during L&D (see also Chapter 2). Therefore, it is difficult to assess the effect of midwife-led care just on L&D.

A systematic review compared the midwife-led continuity model versus other models, including 15 RCTs, for a total of 17,674 women. Women who midwife-led continuity models of care were less likely to experience regional analgesia (RR 0.85, 95% CI 0.78–0.92), instrumental vaginal birth (RR 0.90, 95% CI 0.83–0.97), preterm birth less than 37 weeks (RR 0.76, 95% CI 0.64–0.91), and less overall fetal/neonatal death (RR 0.84, 95% CI 0.71–0.99) [21]. Women who had midwife-led continuity models of care were more likely to experience spontaneous vaginal birth (RR 1.05, 95% CI 1.03–1.07). There were no differences between groups for cesarean births or intact perineum. In addition, women assisted by midwives were less likely to receive some medical interventions, such as amniotomy (RR 0.80, 95% CI 0.66–0.98), episiotomy (RR 0.84, 95% CI 0.77–0.92), and intrapartum analgesia/anesthesia (RR 1.21, 95% CI 1.16–1.37). Women who had midwife-led continuity models of care were less likely to experience longer mean length of labor (hours) (mean difference [MD] 0.50, 95% CI 0.27–0.74), to be attended at birth by a known midwife (RR 7.04, 95% CI 4.48–11.08), and had less fetal loss/neonatal death before 24 weeks (RR 0.81, 95% CI 0.67–0.98). There were no differences between groups for fetal loss or neonatal death more than or equal to 24 weeks, induction of labor, antenatal hospitalization, antepartum hemorrhage, augmentation/artificial oxytocin during labor, opiate analgesia, perineal laceration requiring suturing, postpartum hemorrhage, breast-feeding initiation, low birth weight infant, 5-minute Apgar score ≤ 7, neonatal convulsions, admission of infant to special care or neonatal intensive care unit(s), or mean length of neonatal hospital stay (days). The majority of included studies reported a higher rate of maternal satisfaction in midwife-led continuity models of care [21] (see also section “Continuous Support in Labor”).

**Training of Birth Assistants**

Training of traditional birth assistants in middle- and low-income countries is associated with a trend for less maternal mortality and significantly less perinatal mortality [22]. There are no trials in high-income countries.

**Teamwork Training**

There is insufficient evidence to assess the effects of training and teamwork education for L&D personnel. Quality and safety initiatives are probably beneficial but have not been sufficiently studied in RCTs. Compared with no such training, a standardized teamwork training curriculum based on crew resource management that emphasized communication and team structure was not associated with significant effect on adverse maternal and perinatal outcomes in one RCT [23].
ADMISSION
Delayed versus Early Hospital Admission
Labor assessment programs, which aim to delay hospital admission until active labor, may benefit women with term pregnancies. Active labor was defined as regular painful contractions and cervical dilatation $\geq 3$ cm. In one RCT, compared with direct admission to hospital, delayed admission until active labor is associated with less time in the labor ward, less intrapartum oxytocics, and less analgesia [24]. Women in the labor assessment and delayed admission group report higher levels of control during labor. CD rates are similar, with a nonsignificant $30\%$ decrease. A $30\%$–$40\%$ decrease in CD has been reported in retrospective studies with delayed versus direct admission. There is insufficient evidence (a larger trial needed) to assess true effects on rate of CD and other important measures of maternal and neonatal outcome. Potential risks of delayed admission include unplanned out-of-hospital births and the potentially harmful effects of withholding caregiver support and attention to women in early or latent phase labor.

In another RCT, compared with no use of algorithm, use of an algorithm by midwives to assist in their diagnosis of active labor (painful, regular contractions with at least one of the following: $3$ cm dilated, ROM, or “show”) was associated with more women being discharged after their first labor ward assessment, and no effect of oxytocin augmentation and other medical interventions in labor [25].

Suggested criteria for admission based on these studies are a cervix of at least $3$–$4$ cm dilatation and regular painful contractions. Pregnant women should be informed of these data during prenatal care.

Fetal Assessment Tests upon Admission
FHR Tracing for 20 Minutes
Women allocated to admission cardiotocography (CTG) have an increase in incidence of cesarean section than women allocated to intermittent auscultation (RR 1.20, 95% CI 1.00–1.44). There is no significant difference in instrumental vaginal birth (RR 1.10, 95% CI 0.95–1.27) and fetal and neonatal deaths (RR 1.01, 95% CI 0.30–3.47) [26].

Amniotic Fluid Volume Index
Obtaining an amniotic fluid volume index (AFI) in early labor is associated with a higher incidence of CD, and similar neonatal outcomes, compared with no AFI [27].

Neither a $2 \times 1$ pocket (abnormal in 8%) nor an AFI (abnormal in 25%) upon admission for labor identifies a pregnancy at risk for adverse outcome such as non reassuring fetal heart rate tracing (NRFHT) or CD for NRFHT [28].

Other Tests
There is insufficient evidence to support the use of vibro-acoustic stimulation or Doppler ultrasound as fetal admission tests, as there are no RCTs on these interventions.

Enemas
Enemas do not have a significant beneficial effect on infection rates such as perineal wound infection or other neonatal infections and women’s satisfaction. Compared with no enema, enema in labor is associated with no significant differences for infection rates in puerperal women (RR 0.66, 95% CI 0.42–1.04) and no significant differences in neonatal umbilical infection rates (RR 3.16, 95% CI 0.50–19.82). No significant differences are found in the incidence of neonatal lower or upper respiratory tract infections [29].

One RCT described labor to be significantly shorter (50 minutes) with enema versus no enema. A second RCT found labor to be significantly longer (112 minutes) with enema compared with no enema. No significant differences in the duration of labor were found in the third RCT that scored as having a low risk of bias and was adjusted for parity. One RCT that researched women’s views found no significant differences in satisfaction between groups. The routine use of enamens during labor should be discouraged [29]. This intervention (enema) generates discomfort in women and increases the costs of delivery.

Perineal Shaving
There is no support for routine perineal shaving (shaving with a razor) for women prior to or in labor. In a very old trial, 389 women were alternately allocated to receive either skin preparation and perineal shaving or clipping of vulvar hair only. In the second old trial, which included 150 participants, perineal shaving was compared with the cutting of long hairs for procedures only. In the third trial, 500 women were randomly allocated to shaving of perineal area or cutting of perineal hair. Compared with no shaving, shaving was associated with a similar incidence of maternal febrile morbidity (RR 1.14, 95% CI 0.73–1.76), perineal wound infection (RR 1.47, 95% CI 0.80–2.70), and perineal wound dehiscence (RR 0.33, 95% CI 0.01–8.00). In the smaller trial, fewer women who had not been shaved had Gram-negative bacterial colonization compared with women who had been shaved (OR 0.43, 95% CI 0.20–0.92). There were no differences in maternal satisfaction immediately after birth [30]. The potential for complications (redness, multiple superficial scratches and burning and itching of the vulva, and embarrassment and discomfort afterward when the hair grows back), which often occur later, suggests that shaving should not be part of routine clinical practice. The first two trials are old (1922 and 1965) and included the clipping of long hairs in their control groups to aid in operative procedures, which is itself usually unnecessary and can lead to complications.

Morphine Sleep
Morphine sleep, also called therapeutic rest, given usually in the latent phase of the first stage of labor, has never been evaluated with an RCT [31].

FIRST STAGE
Chlorhexidine
There is no evidence to support the use of vaginal chlorhexidine by either irrigation or vaginal wipes during labor in order to prevent maternal and neonatal infections. The effect on the incidence of postpartum endometritis is not statistically significant (RR 0.83; 95% CI 0.61–1.13) [32]. Chlorhexidine solution is safe, not expensive, and vaginal irrigation is easy to perform, but apparently not beneficial.

GBS Prophylaxis
In the United States and some European countries universal prenatal maternal GBS screening is performed, with anovaginal specimen collected at 35–37 weeks and antibiotic (amoxicillin first line) treatment administered intrapartum to GBS colonized women [33]. Women with GBS bacteriuria in the
current pregnancy or who had a prior infant with GBS sepsis are candidates for intrapartum antibiotics prophylaxis and should be the only two groups not screened. Intrapartum treatment for chorioamnionitis is recommended regardless of GBS maternal status (see Chapter 22 and Chapter 37 in Maternal-Fetal Evidence Based Guidelines). Other European countries use a risk-based strategy, which means to offer intrapartum antibiotic treatment to all women with recognized risk factors: previous infant affected by GBS sepsis, GBS bacteriuria during current pregnancy, preterm labor < 37 weeks, prelabour ROM ≥ 18 hours and/or fever in labor ≥ 38°C. Support for the universal screening strategy is based on a large retrospective study, which concluded that this policy was more than 50% effective in preventing early GBS sepsis [33]. However, no RCTs are available on this issue. There is no intervention 0.90, 95% CI 0.85–0.96), or a baby with a low 5-minute Apgar score (RR 0.69, 95% CI 0.50–0.95). There is no apparent impact on other intrapartum interventions, maternal or neonatal complications, or on breast-feeding. Continuous support is most effective when provided by a woman who is neither part of the hospital staff nor the woman’s social network, and in settings in which epidural analgesia is not routinely available. No conclusions could be drawn about the timing of onset of continuous support. Hospitals should permit and encourage women to have a companion of their choice during labor and birth, and administrators and policymakers should ensure that these services are available to every pregnant woman [34] (see also the section “Midwife-Led Care”).

It may be possible to increase access to one-to-one continuous labor support worldwide by encouraging women to invite a family member or friend to commit to being present at the birth and assuming this role. The mother selects her doula during pregnancy; they establish a relationship (which is likely to involve the woman’s partner, if any) and discuss the mother’s and partner’s preferences and concerns before labor. The doula brings her experience and training (often to the level of certification) to the labor support role during childbirth, and the mother and doula frequently have telephone and/or face-to-face contact in the early postpartum period. Other models of support, for which there are little or no data, include support by a female family member and support by the husband/partner [34].

Continuous Support in Labor

Definition

For support, it is generally intended emotional support (continuous presence, reassurance, and praise), information about labor progress and advice regarding coping techniques, comfort measures (comforting touch, massage, warm baths/showers, promoting adequate fluid intake and output), and advocacy (helping the woman articulate her wishes to others) [34].

Mechanism of Action

Anxiety during labor is associated with high levels of the stress hormone epinephrine in the blood, which may in turn lead to abnormal FHR patterns in labor, decreased uterine contractility, a longer active labor phase with regular well-established contractions, and low Apgar scores. One example of possible mechanisms of action for support to reduce complications of L&D is decreased anxiety [34]. This in turn can lead to a beneficial “chain-reaction”: for example, if continuous support leads to reduced use of epidural analgesia, then several complications associated with regional anesthesia (see Chapter 11) can be prevented.

Types of Support

Family member or friend (not part of hospital staff) or hospital-based (part of hospital staff). Doula (a Greek word for “handmaiden”) is a support person with the sole job of providing support to the laboring woman. They are usually not part of the hospital staff. This member of the caregiver team may also be called a labor companion, birth companion, labor support specialist, labor assistant, or birth assistant.

Effectiveness

Continuous support during labor has clinically meaningful benefits for women and infants and no known harm. All women should have support throughout labor and birth. Women allocated to continuous support are more likely to have a spontaneous vaginal birth (RR 1.08, 95% CI 1.04–1.12) and less likely to have intrapartum analgesia (RR 0.90, 95% CI 0.84–0.96) or to report dissatisfaction (RR 0.69, 95% CI 0.59–0.79). In addition, their labors are about 1-hour shorter (MD –0.58 hours, 95% CI –0.86 to –0.30) (MD –0.58 hours, 95% CI 0.59–0.79). In their labors are about 1-hour shorter (MD –0.58 hours, 95% CI –0.86 to –0.30) (MD –0.58 hours, 95% CI 0.59–0.79). In two studies looked at water only versus giving women specific fluids and foods; and two studies looked at water only versus giving women carbohydrate drinks.

When comparing any restriction of fluids and food versus women given some nutrition in labor (three RCTs), the meta-analysis was dominated by one study undertaken in a highly medicalized environment. There were no statistically significant differences identified in the following: cesarean section (RR 0.89, 95% CI 0.63–1.25), operative vaginal births (RR 0.98, 95% CI 0.88–1.10), and Apgar scores < 7 at 5 minutes (RR 1.43, 95% CI 0.77–2.68), nor in any of the other outcomes assessed. Women’s views were not assessed. The pooled data were insufficient to assess the incidence of Mendelson syndrome (bronchopulmonary reaction following aspiration of gastric contents during anesthesia), an extremely rare outcome [36].

Aromatherapy

There is insufficient evidence to assess the effects of aromatherapy in labor. Compared with no such intervention, administration of selected essential oils during labor is not associated with significant effects on CD, instrumental delivery, or use of oxytocin. While maternal pain perception in nulliparous women was decreased in one RCT on this intervention, this RCT was also not blind [35].

Nutrition in Labor

Since the evidence shows no benefits or harms, there is little justification for the restriction of fluids and food in labor for women at low risk of complications. All studies looked at women in active labor and at low risk of potentially requiring a general anesthetic. One study looked at complete restriction versus giving women the freedom to eat and drink at will; two studies looked at water only versus giving women specific fluids and foods; and two studies looked at water only versus giving women carbohydrate drinks.

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Oral carbohydrate intake, equivalent to about 10 teaspoons of sugar, in labor is not associated with any effect on labor outcomes, like length of labor or mode of delivery [37]. Some umbilical cord studies revealed lactate transport to the fetal circulation with potential (but not observed) fetal academia [38,39].
Maternal Position
Upright and mobile positions (either standing, sitting, kneeling, or walking around) in the first stage of labor reduce the length of labor and do not seem to be associated with increased intervention or negative effects on mothers’ and babies’ well-being. Compared with recumbent positions, the first stage of labor is approximately 1 hour and 22 minutes shorter for women randomized to upright and mobile positions. Women randomized to upright positions are less likely to have epidural analgesia (RR 0.81, 95% CI 0.66–0.99), and cesarean section (RR 0.71, 95% CI 0.54–0.94) [46]. There are no differences between groups for other outcomes including length of the second stage of labor, or other outcomes related to the well-being of mothers and babies, except for a lower incidence of neonatal intensive care unit admission in the upright group. For women who had epidural analgesia there were no differences between those randomized to upright and recumbent positions for any of the outcomes examined in the review. Little information on maternal satisfaction was collected, and none of the studies compared different upright or recumbent positions. A woman semireclining or lying down on the side or back during the first stage of labor may be more convenient for staff and can make it easier to monitor progress and check the baby. Fetal monitoring, epidurals for pain relief, and use of IV infusions also limit movement. Women should be encouraged to take up whatever position they find most comfortable in the first stage of labor [46].

Ambulation
Compared with remaining in bed, walking in labor is associated with similar length of first stage of labor, use of oxytocin, use of analgesia, need for forceps vaginal delivery, or CD, and also similar neonatal outcomes in women at term with cephalic presentation starting at 3–5 cm of dilatation [47] or in other groups of women [46,48–51]. Since walking does not seem to have a beneficial or detrimental effect on L&D, women can choose freely to walk or stay upright (see the section “Maternal Position”) in bed during labor, whichever is more comfortable for them. See also the section “Maternal Position,” as upright position and mobility have been associated with shorter labors.

Immersion in Water
Compared with controls (labor not in water), water immersion during the first stage of labor reduces by 10% the use of epidural/spinal analgesia (RR 0.90, 95% CI 0.82–0.99) and by about 30 minutes the duration of the first stage of labor. There is no difference in other outcomes: assisted vaginal deliveries (RR 0.86, 95% CI 0.71–1.05), cesarean sections (RR 1.21, 95% CI 0.87–1.68), use of oxytocin infusion (RR 0.64, 95% CI 0.32–1.28), perineal trauma or maternal infection, Apgar score <7 at 5 minutes (RR 1.58, 95% CI 0.63–3.93), neonatal unit admissions (RR 1.06, 95% CI 0.71–1.57), or neonatal infection rates (RR 2.00, 95% CI 0.50–7.94) [52]. There is limited information for other outcomes related to water use during the first and second stages of labor, due to intervention and outcome variability. There is no evidence of increased adverse effects to the fetus/neonate or woman from laboring in water. Laboring in water is usually linked to midwifery care, which is associated with its own benefits (see the section “Midwife-Led Care”). There are no trials evaluating different baths/pools, immersion in water during pregnancy, or during the third stage of labor. For water birth and immersion in water in the second stage, see Chapter 8.

No studies looked specifically at women at increased risk of complications; hence, there is no evidence to support restrictions in this group of women [36].

Ice chips to moisten the mouth and sips of clear liquids are the only oral intake recommended by U.S. authorities [40]. Some experts also allow sport drinks, yogurt, or sherbet. In the Netherlands women in labor are allowed to eat and drink. The reason for avoiding solid food is risk of aspiration (“Mendelson syndrome”), which is rare (about 15/10,000 CDs) [41]. When there is increased gastric volume, there is increased risk of vomiting and therefore aspiration. Airway precautions in L&D are paramount to avoid aspiration.

Acid Prophylaxis Drugs
There is no good evidence to support the routine administration of acid prophylaxis drugs in normal labor to prevent gastric aspiration and its consequences [42]. Giving such drugs to women once a decision to give general anesthesia is made is discussed in Chapter 11.

IV Fluids
Two RCTs compared IV fluids (up to 250 mL/hour + oral intake) versus oral intake only. There was a reduction in the duration of labor in IV fluids + oral intake of 29 minutes (MD –28.86, 95% CI –47.41 to –10.30, two studies, 241 women). There was no statistically significant reduction in the number of cesarean deliveries. There was no evidence of a statistically significant difference in admission to the neonatal unit and in low Apgar score at five minutes [43].

Four RCTs compared 250 mL/hour with 125 mL/hour IV fluids in labor in women with restricted oral intake. There was a significant reduction of over one and a half hour in the duration of labor in women who received 250 mL/hour (MD 105.61 minutes, 95% CI 53.19–158.02 (P < 0.0001). There was a significant reduction in the cesarean section rate in women receiving a higher rate of IV fluid infusion (RR 1.56, 95% CI 1.10–2.21; (P = 0.01) [36].

Three RCTs compared women who received 250 mL/hour versus 125 mL/hour of IV fluids with free oral fluids in both groups. Women receiving 250 mL/hour had shorter labors than those receiving 125 mL/hour (MD 23.87 minutes, 95% CI 3.72–44.02, 256 women). There was no statistically significant reduction in the number of cesarean deliveries in the 250 mL IV fluid group (RR 1.00, 95% CI 0.54–1.87). In one study the number of assisted vaginal deliveries was lower in the group receiving 125 mL/hour (RR 0.47, 95% CI 0.27–0.81).

The benefits are substantiated by the fact that several trials in nonpregnant adults demonstrate that increased fluid intake improves exercise performance.

Compared with normal saline without dextrose, normal saline with 5% dextrose, both given IV at 125 mL/hour in early labor (3–5 cm), is associated with a shorter (by about 70 minutes) duration of labor from initiation of IV infusion to delivery. The incidence of labor lasting >12 hours was decreased from 22% to about 10% [44]. Five percent dextrose has also been associated with less umbilical cord acidemia compared with lactated Ringer’s solution [45].

In summary, IV fluids at 250 mL/hour are associated with shorter duration of labor and less cesarean deliveries compared with 125 mL/hour and should be preferred. There is insufficient evidence for which type of IV fluids to use, with possibly some benefit for those containing 5% dextrose.
Stripping in Labor
There are no trials to evaluate stripping during spontaneous labor (see also Chapter 21 for stripping before labor).

Early Artificial Rupture of Membranes (AROM) (aka Amniotomy)
Compared with no amniotomy, amniotomy is not associated with significant differences in length of the first stage of labor (MD –20.43 minutes, 95% CI –95.93–55.06), cesarean section (RR 1.27, 95% CI 0.99–1.63), maternal satisfaction with childbirth experience, or low Apgar score <7 at 5 minutes (RR 0.53, 95% CI 0.28–1.00). Other maternal and perinatal outcomes are also not different. There is no consistency between RCTs regarding the timing of amniotomy during labor in terms of cervical dilatation. An association between early amniotomy and CD for NRRTHT is noted in one large trial [53]. Therefore, routine early (or even late) amniotomy cannot be recommended as part of standard labor management and care. Women should be counselled regarding these results, and make an informed decision regarding the option of this intervention [53]. Early amniotomy may be possibly reserved for women with slow labor progress.

Use of Partogram
A partogram is a preprinted form, the aim of which is to provide a pictorial overview of labor to plot progress in labor and to alert health professionals to any problems with the mother or baby. The general intervention with the partogram is early use of oxytocin as soon as the cervical dilatation falls to the right of the partogram, usually on the 2-hour cervical examinations.

The use of partogram cannot be recommended as a routine intervention in labor. Compared with no partogram, the use of the partogram is not associated with significant effects on cesarean section (RR 0.64, 95% CI 0.24–1.70), instrumental vaginal delivery (RR 1.00, 95% CI 0.85–1.17), or Apgar score <7 at 5 minutes (RR 0.77, 95% CI 0.29–2.06), in two RCTs including 1590 women [54].

Compared with a 4-hour action line, women in the 2-hour action line group are more likely to require oxytocin augmentation (RR 1.14, 95% CI 1.05–1.22). When the 3- and 4-hour action lines are compared, cesarean section rate is lowest in the 4-hour action line group (RR 1.70, 95% CI 1.07–2.70) [53–56]. When a partogram with a latent phase (composite) and one without (modified) were compared, the cesarean section rate was lower in the partogram without a latent phase [54].

Frequency of Cervical Examinations
Only one RCT assessed different frequencies of cervical examination in labor. Comparing two-hourly with four-hourly vaginal examinations in labor, there was no difference in length of labour (MD in minutes –6.00, 95% CI –88.70 to 76.70; one RCT, n = 109). There were no data on maternal or neonatal infections requiring antibiotics, and women's overall views of labor. The study did show that significantly fewer women reported that vaginal examination was very uncomfortable compared with rectal examinations (RR 0.42, 95% CI 0.25–0.70). There were no differences in augmentation, cesarean section, spontaneous vaginal birth, operative vaginal birth, perinatal mortality, and admission to neonatal intensive care.

In summary, cervical assessment should be done by cervical and not rectal examination, as per patient's safety and comfort. There is insufficient evidence to recommend any particular frequency of vaginal cervical examinations in labor. Most studies, including those regarding active management, perform cervical examinations every 2 hours in labor. The risk of chorioamnionitis though increases with increasing number of examinations [58].

Oxytocin Augmentation
There are no trials to evaluate the timing and dosing of oxytocin in labor in women making normal progress in labor.

For women making slow progress in the first stage of spontaneous labor, treatment with oxytocin as compared with no treatment or delayed oxytocin treatment does not result in any discernible difference in the number of cesarean sections performed, in one meta-analysis. In addition there are no detectable adverse effects for mother or baby [59]. The meta-analysis included eight appropriate studies, for a total of 1338 women with low risk singleton pregnancies at term in the active stage of labor. IV oxytocin versus placebo or no treatment (three trials; 138 women) showed no difference as for cesarean section rates; only three small trials were considered so that the comparison was clearly underpowered to make any firm conclusions. Early use of IV oxytocin versus delayed use (five trials; 1200 women), however, was much larger and also showed no effect on cesarean section rates. The early use of oxytocin did significantly increase uterine hyperstimulation with FHR changes; however, this did not translate into serious neonatal morbidity or perinatal death. The early use of oxytocin, as opposed to its delayed use, did significantly shorten the time to delivery by approximately two hours, which might be important to some women.

In a different meta-analysis, early oxytocin was associated with a 9% significant increase in the probability of SVD, but also a tripling of the rate of hyperstimulation, without apparent perinatal consequences [60].

In summary, oxytocin augmentation in women making slow progress in the first stage seems to be associated with a 2-hour shortening of labor, with apparent effect on mode of delivery or other maternal and perinatal outcomes.

There is insufficient evidence to assess how to best use (i.e., dosing issues) oxytocin for augmentation of labor. There are four variables to assess: (1) type of dilution, (2) initial dose, (3) incremental increase, and (4) maximum dose [61]. In a meta-analysis, a higher initial dose (e.g., 2 mU/minute) of oxytocin and incremental dose (e.g., 4 mU/minute or more) was associated with a significant reduction in length of labor reported from one trial (MD –3.50 hours, 95% CI –6.38 to –0.62, one RCT, n = 40). There was a decrease in rate of cesarean section (RR 0.62, 95% CI 0.44–0.86) and an increase in the rate of spontaneous vaginal birth (RR 1.35, 95% CI 1.13–1.62). There were no significant differences for neonatal mortality, hyperstimulation, chorioamnionitis, epidural analgesia; or neonatal outcomes of Apgar scores, umbilical cord pH, or admission to special care baby unit. Many outcomes were not evaluated, such as perinatal mortality, women's satisfaction, instrumental vaginal...
birth, uterine rupture, postpartum hemorrhage, abnormal cardiotocography, women’s pyrexia, dystocia, and neonatal neurological morbidity [62]. Based mostly on physiologic studies, an oxytocin dilution of 10 mU/mL, and initial dose of 2 mU/minute (12 mL/hour), incremental increase of 2 mU (12 mL) every 30–45 minutes until adequate labor, and maximum dose of 16 mU/minute (up to 24 mU) have been proposed [61].

Stopping oxytocin, instead of continuing it, once the active phase of labor is reached in women in spontaneous labor is not associated in maternal or perinatal benefits. Stopping oxytocin is associated with longer (by about 30 minutes) time to delivery [63].

For use of oxytocin in induction, see Chapter 21.

Active Management of Labor

Active management of labor was originally devised to shorten labor, and therefore prevent prolonged labor. Its components have varied somewhat in the literature but generally include antenatal classes, admission not before premature rupture of membranes (PROM) or 2 cm dilatation and full effacement (active labor), early amniotomy, support by doula, use of partogram, and vaginal examinations every 2 hours, with oxytocin started for rate of progress off the partogram or < 1 cm/hour. Oxytocin rate is started at 4–6 mU/minute, increased by 4–6 mU every 15 minutes to reach contractions every 2–3 minutes (but not more than 7/15 minutes) or 40 mU/minutes. Early amniotomy and early use of high-dose oxytocin are the two most characteristic interventions of active management of labor.

Compared with “routine” care (no active management), active management of labor is associated with a trend for a slightly lower incidence of CD (RR 0.88, 95% CI 0.77–1.01). However, in one study there were a large number of postrandomization exclusions. On excluding this study, CD rates in the active management group were statistically significantly lower than in the routine care group (RR 0.77, 95% CI 0.63–0.94). More women in the active management group had labors lasting less than 12 hours, but there was wide variation in length of labor within and between trials. The reduced duration of labor is about 50–100 minutes, mostly in the first stage. There were no differences between groups in use of analgesia, rates of assisted vaginal deliveries, or maternal or neonatal complications. Only one RCT examined maternal satisfaction; the majority of women (over 75%) in both groups were very satisfied with care [64–68].

The shorter labor is probably due to the early amniotomy (see the section “Early Artificial Rupture of Membranes (AROM) [aka Amniotomy]”). The effects on incidence of CD may be due to the fact that some aspects of active management (e.g., support by doula) decrease CD rate, but some others (e.g., early amniotomy) may increase it. It is recommended that the individual interventions that are part of active management of labor should be studied separately, and only those that are beneficial (e.g., support by doula) implemented.

Use of Continuous versus Intermittent Monitoring, Amnioinfusion for Variables, Scalp Sampling, etc.
See Chapter 10.

Bladder Catheterization

There are no trials to evaluate the necessity, timing, and frequency of bladder catheterization in labor per se.

Use of Ultrasound During Labor

There is growing evidence, including RCTs, to assess the effect of using ultrasound during L&D. There are several potential uses, including as an aid to the diagnosis of malposition, dystocia, etc. [69]. Performing routine ultrasound to assess fetal head position in singleton pregnancies in labor ≥ 28 cm, compared with digital vaginal assessment alone, is associated with an increase in operative vaginal delivery, with no other effects on maternal or perinatal outcomes [70].

Use of Intrauterine Pressure Catheter

The IUPC can measure more objectively than external tocometry the intensity of uterine contractions. It necessitates ROM. Intensity is usually calculated by Montevideo units, that is, sum of peak pressures above baseline of all contractions in 10 minutes.

Several RCTs have assessed the effect of IUPC on labor outcomes. In the largest RCT, compared with no IUPC, IUPC use is associated with no effect in rate of operative deliveries, maternal or fetal infection, or other maternal or perinatal recorded outcome [71]. In the meta-analysis, the neonatal outcome was not statistically different between groups: Apgar score < 7 at 5 minutes (RR 1.78, 95% CI 0.83–3.83); umbilical artery pH < 7.15 (RR 1.31, 95% CI 0.95–1.79); admission to the neonatal intensive care unit (RR 0.34, 95% CI 0.07–1.67); and more than 48 hours hospitalization (RR 0.92, 95% CI 0.71–1.20). The pooled risk for instrumental delivery (including cesarean section, vacuum and forceps extraction) was not statistically significantly different (RR 1.05, 95% CI 0.91–1.21). Hyperstimulation was similar between groups (RR 1.21, 95% CI 0.78–1.88). No serious complications were reported in the trials and no neonatal or maternal deaths occurred [72]. See also section “Criteria for Diagnosis of Failure to Progress in First Stage.”

CRITERIA FOR DIAGNOSIS OF FAILURE TO PROGRESS IN FIRST STAGE

According to studies now >60 years old in women without regional anesthesia by Friedman, the active phase began at 2.5 cm and ended at complete dilatation, with an average duration of 4.6–4.9 hours, a mean rate of cervical dilation of 3 cm/hour and the slowest acceptable rate (95th percentile) of 1.2 cm/hour [73,74]. More recently investigators found that the active phase labor in nulliparous women lasts longer than previously thought. According to a systematic review, which included 18 studies (for a total of 7009 women), the active phase of labor among healthy, low risk, nulliparous women at term with a spontaneous labor onset, lasts 6 hours on average, and the average cervical dilation rate is 1.2 cm/hour. This means that rates of cervical dilatation during active labor from 4 cm forward are much slower than those reported by Friedman. According to recent evidence it seems that the slowest acceptable rate of cervical dilation is 0.6 cm/hour. In term women in labor necessitating oxytocin and with epidural, the fifth percentile rate for dilatation is about 0.5 cm/hour for both nulliparous and parous women [75]. In term women with epidural, labor may take more than 6 hours to progress from 4 to 5 cm, and more than 3 hours to progress from 5 to 6 cm of dilation [76]. Moreover, cervical dilation during active phase
labor is often conceptualized linearly, however recent evidence suggest that cervical dilation rates accelerate throughout the majority of labor. So, in common practice, the likelihood of accelerative intervention is much greater in earlier active labor. It must be kept in mind that progression in the earlier part of “active labor” will typically be slower than 0.6 cm/hour, while progression in more advanced active labor will typically be more rapid [77].

Abnormal progression of labor, including terms such as dystocia, dysfunctional labor, failure to progress, cephalopelvic disproportion, and others, is the most common problem in labor, and the reason for the majority of CDs [78]. Risk factors for dystocia are, among others, as follows: obesity, induction, Bishop <5 at start of labor, station higher than –2, persistent occiput posterior, macrosomia, epidural anesthesia, etc. There are no RCTs on interventions for asynclitism [79] (see also Chapter 8). For malposition, see Chapter 24. Dystocia cannot be diagnosed unless ROM has occurred, and adequate oxytocin to achieve at least three to five adequate contractions per hour has been instituted. Dystocia also cannot be diagnosed reliably before the first stage of labor has entered the active phase, which has been defined, especially in nulliparous patients, reliably before the first stage of labor has entered the active phase of labor [80]. Before performing a CD for active phase labor arrest, labor should be arrested for a minimum of 4 hours (if uterine activity is greater than 200 Montevideo units as documented with IUPC) or 6 hours (if greater than 200 Montevideo units could not be sustained) [80,81]. These data are not from an RCT, and there was a significant higher risk of shoulder dystocia among parturient who had arrest for 4 hours or more. Vaginal birth after cesarean (VBAC) and disorders were not included in this study. Please see Chapter 8 for additional evidence on dystocia.

In women at term with singleton gestations and requiring oxytocin by obstetrician because of “dystocia” at 4–6 cm, meperidine 100 mg IV does not affect operative delivery rates and worsens neonatal outcomes compared with placebo [82] (see also Chapter 8).

REFERENCES


54. Lavender T, Hart A, Smyth RM. Effect of partogram use on outcomes for women in spontaneous labour at term. *Cochrane Database Syst Rev*. 2013;7:CD005461. [Meta-analysis: 6 RCTs, n = 7706; of which 2 RCTs partogram vs no partogram; 4 RCTs assessed different partogram designs]


Second stage of labor

Alexis Gimovsky

KEY POINTS

- Prophylactic intrapartum maternal oxygen should not be used in the second stage of normal labor, since it is associated with more frequent low cord blood pH values (<7.20) than the control group.

- Prophylactic intrapartum betamimetics should not be used in the second stage of normal labor, since their usage is associated with an increase in forceps deliveries.

- Women should be encouraged to give birth in the position they find most comfortable, which is usually upright. Use of any upright or lateral position, compared with supine or lithotomy positions, is associated, in women without epidural analgesia, with reduction in duration of second stage of labor, reduction in assisted deliveries, reduction in episiotomies, increase in second-degree perineal tears, increase in estimated blood loss >500 mL, reduction in reporting of severe pain during second stage of labor and reduction in abnormal fetal heart rate patterns. In women with epidural anesthesia, the evidence is limited and insufficient to make a recommendation.

- There is insufficient evidence to recommend for or against maternal stirrup use in the second stage of labor.

- There is insufficient evidence to recommend water immersion in second stage of labor, and the risks have not been adequately assessed.

- In women at term with epidural analgesia and a singleton, cephalic fetus, delayed pushing (waiting 1–3 hours or until "urge to push") is associated with similar rate of operative vaginal delivery and cesarean delivery (CD), and a higher incidence of spontaneous vaginal delivery compared with early (immediate upon entering second stage) pushing. Additionally, the duration of the second stage is longer by about 60 minutes, the duration of pushing is similar, as are all other studied maternal and perinatal outcomes. Careful monitoring of mother and fetus is necessary to allow labor to continue safely.

- In women without epidural anesthesia, compared with encouraging a woman’s own urge to push (open glottis), pushing using the Valsalva maneuver (closed glottis) is associated with shorter (by 19 minutes) duration of labor, similar incidences of operative vaginal deliveries, need for perineal repair, postpartum hemorrhage, and similar neonatal outcomes. Labor attendant should counsel women in labor regarding these data and support the parturient in her own choice of pushing technique.

- Coaching during pushing in the second stage of labor may shorten the second stage of labor, but lack of available coaching support does not adversely affect labor outcomes.

- Use of a dental support device/mouthguard has been associated with a shorter second stage of labor and reduction in the number of failures to descend requiring operative intervention.

- Both manual and belt fundal pressure to aid in vaginal delivery have not been associated with effect on maternal and perinatal outcomes, except for decreased maternal satisfaction.

- Perineal massage and stretching of the perineum with a water-soluble lubricant in the second stage of labor is associated with similar rates of intact perineum compared with the control group. The incidence of third-degree lacerations is decreased.

- Use of perineal warm packs in the second stage of labor is associated with a reduction in third- and fourth-degree tears, as well as a reduction in pain on postpartum days 1 and 2.

- Routine use of the Ritgen’s maneuver does not appear to be associated with any benefits.

- “Hands-poised” is preferred to the “hands-on” method, since they are associated with similar incidences of perineal and vaginal tears, but the hands-on method is associated with a higher incidence of episiotomies.

- Routine episiotomy should not be performed during the second stage of labor, as restricting episiotomy use is associated with less posterior perineal trauma, less suturing, and fewer healing complications.

- Nulliparous women with epidurals should be allowed to labor for at least one additional hour after prolonged (3 hours) second stage of labor, since this results in a lower chance of CD (by 55%) without increasing maternal or neonatal morbidities. Apart from this trial, there is insufficient evidence to determine exactly when the second stage is considered to be prolonged and associated with enough complications as to justify either operative vaginal delivery or CD.

PROPHYLACTIC INTERVENTIONS

Maternal Oxygen

Prophylactic intrapartum maternal oxygen in the second stage of normal labor is associated with more frequent low (<7.20) cord blood pH values than the control group. There are no other statistically significant differences between the groups. There is a tendency toward reduced cord arterial blood oxygen content and oxygen saturation in mothers treated with oxygen compared with controls. Therefore, routine maternal oxygenation throughout the second stage should not be performed. Short-term oxygenation may be beneficial and long-term oxygenation harmful.

Prophylactic Tocolysis

There is no evidence to support the prophylactic use of betamimetics to prevent nonreassuring fetal heart rate tracing (NRFHT) during the second stage of labor. Compared with placebo, prophylactic betamimetic therapy is associated with
an increase in forceps deliveries. The trial protocol required forceps to be used if the second stage of labor exceeded 30 minutes in both groups. There are no clear effects on postpartum hemorrhage, neonatal irritability, feeding slowness, umbilical arterial pH values, or Apgar scores at 2 minutes [2]. Therefore, routine prophylactic tocolysis should not be performed in the second stage of labor.

Prophylactic Positioning to Prevent Shoulder Dystocia
See Chapter 25.

**MANAGEMENT**

**Maternal Position**

There are several benefits for upright posture: sitting (obstetric chair/stool), semirecumbent (trunk tilted backward 30° to the vertical), kneeling, squatting (unaided or using squatting bars) and squatting (aided with Birth cushion) during the second stage of labor.

Use of any **upright or lateral position**, compared with supine or lithotomy positions, is associated, in women without epidural analgesia, with a small (4 minutes) reduction in duration of second stage of labor, a 20% reduction in assisted deliveries, a 17% reduction in episiotomies, a 23% increase in second-degree perineal tears, a 63% increase in estimated blood loss >500 mL, a 23% reduction in reporting of severe pain during second stage of labor, and a 69% reduction in abnormal fetal heart rate patterns [3]. Use of the birth stool showed no effect and results with the birth chair were variable. Estimation of blood loss in the upright group may be influenced by the fact that blood loss in the birth chair is collected in a receptacle. Physiological advantages for upright labor may include lessened risk of aortocaval compression, improved acid–base outcomes in the neonates, stronger and more efficient uterine contractions, improved alignment of the fetus for passage through the pelvis (“drive angle”) and larger anteroposterior and transverse pelvic outlet diameters, resulting in an increase in the total outlet area in the squatting and kneeling positions [3].

In women with epidural anesthesia, the evidence is limited and insufficient to make a recommendation. In a small randomized controlled trial (RCT), compared with a supported sitting position, lateral position was associated with a lower chance of an operative vaginal delivery [4]. Kneeling and sitting upright are associated with similar duration of second stage and other outcomes, except for a more favorable maternal experience and less pain associated with kneeling [5]. In a Cochrane review of five RCTs, it was concluded that there was no significant difference in operative vaginal delivery, CD, lacerations or neonatal outcomes when comparing women with analgesia during labor in the upright or recumbent positions [6].

In summary, **women** without epidural anesthesia **should be encouraged to give birth in the upright position**, which is also the position they usually find most comfortable [3]. The evidence is insufficient for a recommendation on women with epidural anesthesia.

**Maternal Stirrup Use**

Routine use of stirrups during labor has had limited study insufficient to make a recommendation. In one small RCT, women who delivered in bed with stirrups versus those who delivered without stirrups had similar perineal lacerations, incidence of prolonged second stage, operative vaginal delivery and CD incidence [7].

**Epidural or Other Anesthesia**

See Chapter 11.

**Water Immersion**

Although there is some evidence that water immersion in the first stage of labor may reduce the need for epidural/spinal anesthesia and the length of first stage of labor (see Chapter 7), immersion during the second stage has been insufficiently studied and may be best avoided given lack of sufficient safety data. Of the three trials that compared water immersion during the second stage with no immersion, only one trial showed a significantly higher level of satisfaction with the birth experience (relative risk [RR] 0.24, 95% confidence interval [CI] 0.07–0.70). There are reports of neonatal aspirations from birth (end of second stage) in water [8,9]. American College of Obstetricians and Gynecologists (ACOG) Committee Opinion concludes that, as safety has not been well established, water immersion in the second stage of labor should be considered experimental [10]. In summary, there is insufficient evidence to recommend water immersion in second stage of labor, and risks have not been adequately assessed.

**Pushing**

**Delayed versus Early Pushing**

In women at term with epidural analgesia and a singleton, cephalic fetus, delayed pushing (waiting 1–3 hours or until urge to push) is associated with a similar rate of operative vaginal delivery and of CD, and a higher incidence of spontaneous vaginal delivery compared with early (immediately upon entering second stage) pushing (RR 1.22, 95% CI 1.05–1.42) [11]. The duration of the second stage is **longer by about 60 minutes**, the duration of pushing is similar, as are all other studied maternal outcomes. The neonatal outcomes are also similar, including incidence of admission to neonatal intensive care unit (NICU). In a meta-analysis of 12 RCTs, similar results were noted. Second stage is longer in the delayed pushing group by 57 minutes, while there is a **shortened duration of active pushing** (by 22 minutes). Delayed pushing increases spontaneous vaginal delivery rate compared to immediate pushing, while operative vaginal delivery rates are not different [12]. The longer duration of second stage with delayed pushing has not been associated with detrimental effects on the fetus, but careful monitoring of both mother and fetus is necessary to allow labor to continue safely (see also section “Criteria for Diagnosis of Prolonged Second Stage”). There is no evidence available regarding how long to delay pushing. The 2012 International Federation of Gynecology and Obstetrics (FIGO) guideline for management of the second stage of labor recommends allowing nulliparous women to wait up to four hours before pushing and multiparous women up to one hour before pushing [13].

**Pushing Method: Valsalva versus Spontaneous**

Most women spontaneously choose to Valsalva during the second stage of labor. In women without epidural anesthesia, compared with encouraging a woman’s own urge to push (open glottis), pushing using the Valsalva maneuver (closed glottis: taking a deep breath, holding it and pushing for as long and hard as possible, two to three times during each contraction) is associated with shorter (by 19 minutes) duration of labor, similar incidences of operative vaginal deliveries, need for perineal repair, postpartum hemorrhage, and
neonatal outcomes [14]. Urodynamics 3 months after delivery are worse in the closed glottis group, but long-term outcome has not been studied [14]. Labor attendants should counsel women in labor regarding these data and support the parturient in her own choice of pushing technique.

Coached Pushing
Although coached pushing confers the benefit of a slightly shorter second stage, coached maternal pushing confers no other advantages and withholding such coaching is not detrimental to maternal or fetal outcomes [15].

Use of a Dental Support Device
Wearing a dental support device (mouthguard) is associated with a shorter second stage (19 minutes) in one RCT. In addition, less operative intervention was used in the dental support group. Further research into optimizing maternal expulsive efforts is needed to evaluate the overall benefit of dental devices [16].

Fundal Pressure
In the second stage of labor, fundal pressure has been evaluated either as just manual pressure or with an obstetric belt wrapped around the woman's abdomen above the level of the uterine fundus.

Compared with no manual fundal pressure, manual fundal pressure (Kristeller maneuver) concomitant with each contraction while patient has the urge to push during the second stage is not associated with any significant changes in duration of labor or other maternal and perinatal outcomes in one RCT [17].

The fundal pressure belt inflates with each contraction to a maximum of 200 mmHg for 30 seconds. Compared with no belt, the inflatable obstetric belt is associated with similar incidence of spontaneous vaginal delivery in nuliparous women with singleton term pregnancies and an epidural at term. All other maternal and neonatal outcomes are similar, but women with no belt have greater satisfaction [18].

In summary, neither manual nor belt fundal pressures to aid in vaginal delivery have been associated with effect on maternal and perinatal outcomes, except for decreased maternal satisfaction.

Perineal Massage
Perineal massage has been evaluated for decrease in perineal lacerations. Perineal massage has not been associated with complications. For perineal massage during pregnancy and before labor, see Chapter 2. Perineal massage and stretching of the perineum with a water-soluble lubricant in the second stage of labor are associated with similar rates of intact perineum compared with the control group. The incidence of third-degree lacerations is decreased [19]. In another RCT, perineal massage with lubricant was associated with similar incidence of genital tract trauma compared to no massage or to warm compresses [20]. A meta-analysis favored perineal massage versus hands off for a reduction in third or fourth degree lacerations (RR 0.52, 95% CI 0.29–0.94) [21]. The most recent double-blinded RCT comparing perineal oil with liquid wax (jojoba oil) showed no significant difference in perineal lacerations [22].

Perineal Warm Packs
Although no difference was seen in minor perineal trauma or requirement for suturing, application of perineal warm packs during the second stage of labor was associated with less severe (third- or fourth-degree) tears than was standard management. In addition, pain scores were less on postpartum days 1 and 2 in the intervention group. After 3 months, a trend toward decreased symptoms of urinary incontinence was seen in the intervention group [23]. In another RCT, warm compresses were associated with similar incidence of genital tract trauma compared to the presence or absence of a “no touch” technique or to perineal massage [20]. The decrease in third or fourth degree lacerations with perineal warm packs in the second stage of labor is also supported by a Cochrane review (RR 0.48, 95% CI 0.28–0.84) [21].

Manual Rotation
Persistent fetal occiput posterior position is a risk factor for prolonged labor and higher rate of CD. Malposition includes usually occiput transverse or posterior positions. These are often associated with asynclitism, defined often as the “oblique malpresentation of the fetal head in labor” [24]. There is insufficient evidence (no trials) to evaluate the efficacy of manual rotation in labor. In a retrospective cohort study of women with a fetus in persistent occiput posterior or transverse position in the second stage of labor, manual rotation was associated with lower rates of CD, perineal lacerations, postpartum hemorrhage and chorioamnionitis, but with a higher rate of cervical lacerations [25]. Another prospective study (but not randomized) of singletons with occiput posterior position reported an increase in fetuses delivered in occiput anterior position (93% vs. 15%) and in spontaneous vaginal delivery (77% vs. 27%) compared with no manual rotation, using historic controls [26] (see also Chapter 24).

Ritgen’s Maneuver
Ritgen’s maneuver (originally described by Ritgen in 1855) involves reaching for the fetal chin when it reaches the plane between the maternal anus and the coccyx and pulling it anteriorly, while using the fingers of the other hand on the fetal occiput to control speed of delivery and keep flexion of the fetal neck. Its aim is usually to protect the perineum from lacerations. Compared with no Ritgen’s maneuver, Ritgen’s maneuver performed for delivery of fetal head during uterine contractions is not associated with any effect on the incidence of perineal tears or other reported maternal or perinatal outcomes. The length of second stage and several perinatal outcomes were not reported in this RCT [27]. In summary, Ritgen's maneuver does not appear to be associated with any benefits.

“Hands-On” versus “Hands-Poised”
The hands-on method (also described by Ritgen) involves employing pressure on the infant's head upon crowning and supporting the perineum with the other hand. The aim is to protect against lacerations. In the hands-poised method, the fetal head and perineum are not touched or supported by the delivering personnel. These two methods are associated with similar incidences of perineal and vaginal tears, but the hands-on method is associated with higher incidence of episiotomies (RR 0.69, 95% CI 0.50–0.96) [21,28]. A policy of hands-poised has also been supported by a quasi-randomized study, reporting less third-degree tears compared with hands-on [29]. In a meta-analysis of five RCTs, the hands-on method does not protect against obstetric and anal sphincter injuries (RR 1.03, 95% CI 0.32–3.36) [30]. Therefore, the hands-on method should not be routinely employed in labor.
Episiotomy

Routine episiotomy should not be performed, as restrictive episiotomy policies have a number of benefits compared with routine episiotomy policies. In the most recent meta-analysis of RCTs, 75% of women had episiotomies in the routine episiotomy group, while the rate in the restrictive episiotomy group was 28%. Compared with routine use, restrictive episiotomy is associated with less severe perineal trauma (RR 0.67, 95% CI 0.49–0.91), less suturing (RR 0.71, 95% CI 0.61–0.81), and fewer healing complications (RR 0.69, 95% CI 0.56–0.85) [22].

Restrictive episiotomy is associated with more anterior perineal trauma (RR 1.84, 95% CI 1.61–2.10). There was no difference in severe vaginal/perineal trauma (RR 0.92, 95% CI 0.72–1.18), dyspareunia (RR 1.02, 95% CI 0.90–1.16), urinary incontinence (RR 0.98, 95% CI 0.79–1.20), or several pain measures. Results for restrictive versus routine mediolateral versus midline episiotomy were similar to the overall comparison [31]. There is insufficient evidence to evaluate whether these findings are strictly for use of episiotomy, such as in operative vaginal delivery, preterm delivery, breech delivery, predicted macrosomia or presumed imminent tears. Episiotomy should be avoided if at all possible; but, if used, it is unknown which episiotomy technique (mediolateral or midline) provides the best outcome.

CRITERIA FOR DIAGNOSIS OF PROLONGED SECOND STAGE

There is insufficient evidence to determine when the second stage is prolonged, and associated with enough complications with continuing labor so as to justify either operative vaginal delivery or CD. Some have proposed the criteria in Table 8.1 for the diagnosis of prolonged second stage (or criteria for dystocia or failure to progress), but these are not based on level 1 evidence [32–35]. Clinically, the start of the second stage is imprecise and begins when the subjectively timed cervical examination reveals complete (10 cm) cervical dilation. It is also important to realize that, based on data above, women often do not begin to push until 1 hour or more after complete dilation has been ascertained.

American guidelines discuss the concepts of passive and active second stage [36]. Passive second stage is defined as full dilation of the cervix without voluntary or involuntary pushing. Active second stage is defined as when the fetus is visible or once pushing has started with or without contractions. These guidelines suggest that the passive phase in a nulliparous woman be up to 2 hours regardless of anesthesia. In a multiparous woman, passive phase is suggested to be 1 hour without an epidural and 2 hours with an epidural. The active phase of the second stage of labor is suggested in nulliparous women to have a time limit of 1 hour without an epidural and 2 hours with an epidural. In a multiparous woman, active phase is suggested as 1 hour regardless of anesthesia [36]. The suggestions regarding length of the active second stage are not supported by level 1 data, and we allow longer second stages (see Table 8.1).

Several non-level-1 studies have compared maternal and perinatal outcomes between women with shorter versus “prolonged” second stage. In a review of all studies up to 2004, a strong association between prolonged second stage and operative vaginal delivery was noted [37]. The definition of prolonged second stage was inconsistent across studies. In addition, significant associations with maternal outcomes such as postpartum hemorrhage, infection, and severe obstetric lacerations were reported, but methods varied widely. From other data, urinary incontinence may also be increased with prolonged second stage [34]. Anal incontinence does not seem to be affected [38]. No clear associations between prolonged second stage and adverse neonatal outcomes were reported [26]. The length of the second stage is not associated with poor neonatal outcome as long as reassuring fetal testing is present. A recent large retrospective study suggested that neonatal sepsis (odds ratio [OR] 2.08, 95% CI 1.60–2.70), asphyxia (OR 2.39, 95% CI 1.28–4.27), and mortality (OR 5.92, 95% CI 1.43–24.51) were increased in nulliparous women with prolonged second stage of labor [39]. Nevertheless, these data are limited by their retrospective nature.

The problem with the evidence above is that these maternal detriments of prolonged second stage occur when these women are compared with women without prolonged second stage. It is evident that a planned CD before labor might decrease some of these complications (e.g., bleeding, infection, lacerations, and incontinence). The clinical issue is different, though. Once a woman has prolonged second stage, should she be delivered operatively or should she continue labor? CD performed after prolonged second stage has been associated with longer surgery time, increased postoperative fevers, maternal intraoperative trauma including higher risk of extensions of the uterine incision and higher composite maternal morbidity, but similar perinatal outcomes, compared with CD performed before prolonged second stage [40].

In a recent RCT, extending the second stage of labor beyond 3 hours by at least one additional hour in nulliparous women with epidural anesthesia decreased the CD incidence by 55% without incurring any additional increase in neonatal or perinatal morbidity (RR 0.45, 95% CI 0.22–0.93) [41]. Operative intervention is not warranted based solely on the time elapsed in the second stage. If there are no signs of infection (maternal or fetal), no maternal exhaustion and reassuring fetal testing, labor can be allowed to continue beyond current limits (Table 8.1) as long as some progress has been made. Nevertheless, if contractions are adequate, the chance of vaginal delivery decreases progressively after 3–5 hours of pushing in the second stage [42].

Mandatory second opinion is associated with 22 fewer intrapartum CDs per 1000 deliveries, without affecting maternal or perinatal outcomes [43] (see Chapter 13).

Table 8.1 Proposed Criteria for Prolonged Second Stage

<table>
<thead>
<tr>
<th>Maternal Characteristics</th>
<th>Suggested Upper Limit of Length of Second Stage (Hours)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nulliparous, epidural</td>
<td>4</td>
</tr>
<tr>
<td>Nulliparous, no epidural</td>
<td>3</td>
</tr>
<tr>
<td>Multiparous, epidural</td>
<td>3</td>
</tr>
<tr>
<td>Multiparous, no epidural</td>
<td>2</td>
</tr>
</tbody>
</table>


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PART 1—NORMAL THIRD STAGE OF LABOR

Key Points

- **Oxytocin** is the prophylactic uterotonic of choice at delivery as it reduces blood loss and has fewer side effects compared with other agents such as ergot alkaloids and prostaglandins, including misoprostol. Oxytocin 10 IU intramuscular (IM), preferably to 5 IU IM, or 20 IU (10–40 IU) intravenous (IV) in 500 cc normal saline (NS) or lactated Ringer’s (LR) infused over 1 hour should be routinely administered following vaginal delivery, either after delivery of the anterior shoulder or immediately after delivery of the neonate and before delivery of the placenta.

- **Active management of the third stage of labor (AMTSL)**, including routine, prophylactic use of an oxytocic agent, early cord clamping (ECC) and cutting, and cord traction, shortens the third stage and reduces blood loss and postpartum hemorrhage (PPH), compared with expectant management.

- **Tranexamic acid (TA)**, (in addition to oxytocin administered after delivery), before skin incision at cesarean section could be given, as current data suggest it can significantly decrease blood loss >1000 cc. It reduces the risk of blood loss >400 or >500 cc but not >1000 cc after vaginal delivery.

- **Carbetocin**, (not currently available in the United States) has been shown to reduce the need for therapeutic uterotonics compared with oxytocin in women who underwent cesarean section, but not for vaginal delivery. Carbetocin is associated with less blood loss and fewer side effects when used prophylactically compared with syntometrine.

- For cesarean, misoprostol combined with oxytocin appears to be more effective than oxytocin alone in reducing intraoperative and postoperative hemorrhage.

- **Delayed cord clamping (DCC)** in preterm neonates by 30–120 (maximum 180) seconds is associated with fewer transfusions and lower risk of intraventricular hemorrhage (IVH).

- DCC in term neonates is associated with higher hematocrit and lower risk of iron deficiency at less than a year of age, but with an increased risk of hyperbilirubinemia requiring phototherapy.

- Vaginal and perineal lacerations should be repaired with one continuous absorbable synthetic suture, including continuous subcuticular skin repair.

- Rectal nonsteroidal anti-inflammatory drugs (NSAIDs), topical anesthetics, and therapeutic ultrasound can each decrease perineal pain and need for additional pain therapy.

Definitions

Third stage of labor is defined as the interval between delivery of the neonate and expulsion of the placenta. Mean length of time of the third stage is about 6 minutes. An interval of greater than 30 minutes is the 97th percentile, and represents the definition of a prolonged third stage.

Pathophysiology

The third stage involves separation of the placenta with capillary hemorrhage and shearing of the placental surface when the uterus contracts after delivery of the infant. Signs of separation include a gush of blood, cord lengthening, and the uterine fundus becoming more globular and firm.

Complications (Chapter 26)

- **PPH**—classically defined as an estimated blood loss (EBL) >500 cc after vaginal delivery or >1000 cc after cesarean delivery

- Retained placenta—placenta not expelled ≥30 minutes after infant delivery

- Uterine inversion—collapse of uterine fundus into the uterine cavity

Management

An algorithm including uterotonics use and active management of placental delivery is shown in Figure 9.1.

Uterotonics

**Oxytocin (Syntocinon®)**. Oxytocin binds to specific uterine receptors with immediate action causing increasing strength and frequency of contractions (Table 9.1). The mean half-life is 3 minutes with the plateau reached after 30 minutes. It can either be given as IV or IM, though IM injection has time of onset of 3–7 minutes and clinical effect is longer, lasting 30–60 minutes. It is metabolized by the liver and kidneys with a known 5% antidiuretic effect of vasopressin. If given in large volumes, greater than 40–50 cc/minute, and high concentration, or when given without dilution, it may result in hyponatremia and syndrome of inappropriate antidiuretic hormone hypersecretion (SIADH) with symptoms of headache, vomiting, drowsiness, and convulsions.

Many different ways of administering oxytocin for prophylaxis in the third stage of labor following vaginal delivery have been used, including IM, IV (either undiluted or as a diluted infusion), and intraumbilical. The IM dose is 5–10 IU, administered at the delivery of the anterior shoulder, or soon after, before delivery of the placenta. Oxytocin can also be given as an undiluted IV bolus, or a diluted IV infusion consisting of 10–40 IU diluted in 500–1000 cc of NS or LR. A recent randomized controlled trial (RCT) compared various oxytocin doses (10 IU, 40 IU, and 80 IU IV in 500 mL) given over 1 hour after placental delivery for primary PPH prophylaxis at vaginal delivery, and showed no differences overall in incidence of PPH. However, the 80 IU dose was associated with
less need for additional uterotonics, and less risk of decline in hematocrit $\leq 6\%$ [1]. The comparative effectiveness and safety of these various approaches have not been well quantified in high-quality RCTs so there is insufficient evidence to recommend a specific route or dosage [2]. The IM route is favored in some institutions, particularly in resource-poor settings, as it can be administered rapidly by providers with minimal training. Undiluted IV boluses are generally not recommended as they are associated with transient hypotension and tachycardia. Intraumbilical injection is not superior to IV administration and has not been shown to be effective in reducing blood loss or manual placental removal [3].

**Oxytocin is the prophylactic uter tonic of choice in the third stage of labor.** Compared with no uterotonics, prophylactic oxytocin reduces blood loss of >500 cc after vaginal delivery and the need for therapeutic uterotonics [4]. There are similar incidences of manual removal of the placenta and rates of blood transfusions with the use of oxytocin compared with no uter tonic.

**Compared with ergot alkaloids** (methergine or ergometrine [ergonovine]), oxytocin is associated with fewer side effects including nausea and vomiting, with no difference in length of the third stage or need for manual placental removal [4]. There is no clear evidence that oxytocin is superior to ergot alkaloids in blood loss prevention, however the side effect profile makes oxytocin a better first-line option [4].

**Compared with oxytocin alone,** ergometrine in addition to oxytocin is associated with increased side effects with only modest benefits. Benefits include a statistically significant reduction in the risk of PPH when compared with oxytocin alone for blood loss of 500 ml or more. There is no difference in PPH for blood loss >1000 cc. In addition, there are no differences in blood transfusion, retained placenta, or other neonatal outcomes. Unfortunately, adverse side effects are significant including nausea, vomiting, and hypertension, from the ergometrine added to oxytocin [5]. When compared with ergot alkaloids alone, oxytocin in addition to ergot alkaloids was not beneficial in PPH prevention [4].

As part of AMTSL, oxytocin is routinely administered after delivery of the anterior shoulder or immediately after delivery of the neonate and before delivery of the placenta.
of oxytocin administration before or after placental delivery, did not show differences in rates of blood loss, length of third stage of labor and rates of retained placenta [6,7]. Nonsignificant trends for less PPH but more retained placenta are associated with oxytocin given before compared with after placental separation [6].

**Oxytocin prophylaxis** with 20–80 IU in 500 cc over 30 minutes after cord clamping is recommended following delivery by cesarean section [8]. However, the optimal route and dosing in this situation has not been established (see Chapter 13).

**Oxytocin Agonists**

**Carbetocin** (not currently available in the United States) is a synthetic analogue of oxytocin; 100 µg IV has been shown to reduce the need for therapeutic uterotonics compared with oxytocin, for those women who underwent cesarean section, but not for vaginal delivery; and reduced need for uterine massage following any mode of delivery [9]. There is limited comparative evidence on adverse events [9,10]. Compared with syntometrine (oxytocin plus ergometrine), carbetocin is associated with less blood loss and fewer side effects when used prophylactically following vaginal delivery; however the risk of PPH is not decreased [9]. The World Health Organization (WHO) together with pharmaceutical companies is conducting a multicountry trial, in low, middle and high income countries, expecting to enroll 29,000 women to test effectiveness of a proprietary formulation of carbetocin compared with oxytocin to reduce PPH.

**Ergot Alkaloids (Methergine® or Ergometrine)**. Methyl-ergonovine and its parent compound ergometrine result in sustained tonic contraction of uterine smooth muscle by stimulation of α-adrenergic myometrial receptors. The dose is 0.2 mg IM injection or per os (PO) with a mean elimination half-life of 3.39 hours (range 1.5–12.7 hours). Intravenous administration is not recommended as it is associated with more severe side effects. Nausea and vomiting are common side effects, although the most concerning side effect is vasoconstriction of the vascular smooth muscle. This results in elevation of central venous pressure and systemic blood pressure increasing the risk of pulmonary edema, stroke, or myocardial infarction. Contraindications include cardiac disease, autoimmune diseases associated with Raynaud’s disease, and pulmonary disease. There is no high-quality evidence to support the use of carboprost for PPH prophylaxis, but it can be used as a second-line agent for treatment of PPH (see Chapter 26).

**Prostaglandin F2α (hemabate/carboprost)** causes contraction of uterine smooth muscle cells. Administration is either 0.25 mg IM or as a direct injection into the myometrium, which may be repeated every 15–20 minutes for a maximum of 8 doses or 2 mg. Side effects are secondary to smooth muscle constriction and include bronchoconstriction, venoconstriction, and constriction of GI smooth muscle. Common side effects are nausea, vomiting, diarrhea, pyrexia, bronchospassms, and case reports of hypotension and intrapulmonary shunting with arterial oxygen desaturation. It is contraindicated in patients with cardiac and pulmonary disease. There is no high-quality evidence to support the use of carboprost for PPH prophylaxis, but it can be used as a second-line agent for treatment of PPH (see Chapter 26).

**Tranexamic Acid**. TA is a synthetic derivative of the amino acid lysine that is metabolized in the kidney and functions as an antifibrinolytic. For prophylaxis, it is administered intravenously at a dose of 10 mg/kg (usually maximum of 1 g), before skin incision (e.g., 30 minutes prior), with rapid onset of action and a mean elimination half-life of 2–10 hours. Side effects include nausea, vomiting and dizziness [18].

TA given in combination with routine prophylactic uterotonics appears to decrease the risk of postpartum blood loss >400 cc, >500 cc, and >1000 cc, need for blood transfusion in women at low risk for PPH following a cesarean delivery with data from clinical trials of mixed quality and meta-analysis from such data [18,19]. For vaginal delivery, there is insufficient evidence to recommend the routine use of TA alone for PPH prophylaxis, as well as its use in a population at high risk for PPH. It has been reported it reduces the risk of blood loss >400 or >500 cc but not >1000 cc after vaginal delivery. There is variability amongst available studies, which are too small to assess rare outcomes of venous thromboembolism or maternal mortality. Therefore, **we recommend it be considered for routine prophylactic use at cesarean section in addition to prophylactic oxytocin**. Currently a large multi-country RCT “The Woman Trial” is testing the use of 1 g of TA initially, followed by 1 g if bleeding continues, in women diagnosed with PPH, compared with placebo. It is expected 20,000 women will be recruited and that this RCT will provide definitive data on use of TA for treatment of PPH (Chapter 13).
Summary

Oxytocin (5–10 IU IM or 10–40 IU IV in 500–1000 cc NS or LR bolus) should be routinely administered with delivery of the anterior shoulder or before delivery of the neonate, in all pregnancies as prophylaxis to prevent PPH. At cesarean, TA before skin incision should be given, in addition to oxytocin after delivery of the fetus. Methergine, syntometrine, and prostaglandin F2α should generally not be used for prophylaxis in normal third stage of labor, unless oxytocin is unavailable. Methergine, prostaglandin F2α, and misoprostol are often used for treatment of PPH (see Chapter 26).

Umbilical Cord

Cord Blood Collection

Cord gases are stable in a clamped segment of cord for up to 60 minutes and in a heparinized syringe for up to 60 minutes. Umbilical artery pH, pCO2, and base deficit may be helpful in indicating timing of insult and can be collected in cases of non-reassuring fetal heart rate tracings (NRHFTs), meconium, low Apgar scores (defined as below 7 at 5 minutes), growth restriction, perinatal birth, or any sentinel event including cord prolapse, uterine rupture, or placental abruption. Umbilical vein pH may be helpful in cases of utero-placental problems like growth restriction, placental abruption, asthma, and hypertension. While there are no RCTs on this issue, routine sending of umbilical cord gases is usually not necessary or recommended for normal labor, delivery, and Apgar scores, without risk factors.

Cord Blood

Cord blood is routinely sent for Rh status of the infant, especially in Rh-negative women. Cord blood collection for stem cells has increased in popularity in recent years. Obstetricians should support public banking of cord blood [20]. Public banking is recommended over private banks secondary to more stringent U.S. Food and Drug Administration (FDA) guidelines, given its increased legal responsibilities, cost-effectiveness, and greater access to cord blood by the general population. The chance of a child requiring an autologous transplant from privately banked cord blood is about 1/2700. Directed donation of umbilical cord blood when there is a disease in the family amenable to privately banked cord blood is associated with fewer transfusions for anemia, less IVH and lower risk for necrotizing enterocolitis than ECC at <30 seconds [21]. This intervention also benefits very preterm neonates <32 weeks, with decreased mortality, risk of blood transfusion and IVH [22]. In addition, DCC has been shown to be safe and does not compromise the preterm infant [23]. There are no clear differences in other outcomes including respiratory distress, death, initial adaptation phase, or long-term outcomes. There is no current recommendation on positioning of the preterm infant during DCC secondary to insufficient data. One RCT showed no differences in neonatal outcomes during a 120 second DCC after vaginal delivery with the neonate held either at the level of the vagina, or the level of the abdomen or chest, concluding that mothers can safely be allowed to hold their baby on their abdomen or chest for skin-to-skin and breastfeeding [24].

Milking of the cord following preterm delivery has been examined in a few RCTs with benefits similar to DCC. Compared with no milking, milking of the cord has been associated with higher hematocrit, lower transfusion rates and less need for circulatory and respiratory support [25–27]. One study examined the effect of DCC in addition to cord milking versus cord milking only, and found no differences [28].

In term neonates, DCC has also been shown to have some benefits, but also some risks. Compared with early clamping, DCC is associated with higher hematocrit and a lower risk of iron deficiency at less than a year of age. There was no increased risk of PPH, however, there was an increased risk of the neonate needing phototherapy for hyperbilirubinemia. The possible increased risk of neonatal jaundice requiring phototherapy must be weighed against the physiological benefit of greater hemoglobin and iron levels conferred by DCC in term infants, which may be of clinical value particularly in infants where access to good nutrition is poor [29].

Placenta

Cord Drainage with or without Traction

Immediate cord drainage and traction is associated with a shorter duration of the third stage of labor and a smaller decrease in hemoglobin compared with no drainage or traction [30]. Cord drainage with traction reduces the length of the third stage of labor and decreases blood loss compared with traction and no cord drainage [31].

Cord Traction

As mentioned above, the combination of cord drainage with traction appears to shorten the third stage of labor and leads to a smaller decrease in hemoglobin compared with no drainage or traction [30]. Cord traction alone has been associated with lower mean blood loss and decreased risk of PPH, shorter third stage of labor, and lower risk of needing manual placental removal compared with no cord traction [32].

Massage of the Uterus

There is limited evidence to evaluate the effect of massage of the uterus alone. Three trials have examined the use of massage in addition to oxytocin. In one small trial, use of uterine massage resulted in decreased mean blood loss and need for additional uterotonics. A larger trial did not demonstrate benefit when uterine massage was added to oxytocin treatment [33]. A more recent RCT from China also did not show a significant decrease in blood loss with addition of uterine massage to 10 IU oxytocin IM prophylaxis [34]. However, uterine massage has been evaluated as a component of AMTSL and found to be beneficial in prevention of a prolonged third stage and PPH. Therefore, uterine massage after delivery of the placenta is recommended.

Injection of Oxytocin in Umbilical Cord

Injection of oxytocin into the umbilical cord is not recommended. Compared with IV/IM oxytocin or to injection of saline alone, the umbilical vein injection of oxytocin does not have a significant effect on postpartum blood loss, need for transfusion, length of the third stage of labor or need for manual placental removal [3].

Active Management of the Third Stage of Labor

AMTSL usually consists of

- Prophylactic oxytocin at delivery of the anterior shoulder or immediately after delivery of the baby
- ECC and cutting
- Controlled cord traction
Expectant management usually consists of:

- No uterotonics
- No ECC, cutting, or traction
- Use of gravity and maternal expulsive efforts for placenta delivery

There is heterogeneity in the literature regarding different definitions for AMTSL. We have used the one utilized for evaluation of the existing RCTs of AMTSL by the Cochrane collaboration [35]. AMTSL has been used mostly in pregnancies at term after vaginal births, so recommendations are limited to this population. AMTSL reduces the risk of PPH and should be offered and recommended to all women, including women following preterm birth. Compared with expectant management, active management is associated with reduced risk of maternal primary hemorrhage of blood loss >1000 cc and >500 cc, and of maternal hemoglobin <9 g/dL following birth [35]. Adverse effects were identified including increases in vomiting after birth, increased maternal diastolic blood pressure, increased use of analgesia and pain, more women returning to the hospital with bleeding, and decrease in infant birth weight, possibly reflecting some degree of lower blood volume from placental transfusion if ECC is performed. There was no effect on risk of infant admission to the neonatal unit or incidence of jaundice requiring treatment. Each element of AMTSL should be assessed separately, as we have done in the preceding sections “Umbilical cord” and “Placenta.”

Repair of Laceration

Closure and Repair

Compared with nonclosure, closure of first- and second-degree perineal lacerations after vaginal delivery is associated with better healing seen at 10 days and 6 weeks and similar pain scores [36–38]. The continuous suturing techniques, compared with the interrupted, are associated with less short-term pain in second-degree and episiotomy repairs [39]. Furthermore, approximation (end-to-end) and overlap technique for third-and fourth-degree laceration repair are associated with similar outcomes in two RCTs [40,41], while in the most recent RCT end-to-end repair was associated with significantly lower rates of anal incontinence, with no difference in long-term outcomes [42]. One small randomized trial showed that use of antibiotics at the time of third- and fourth-degree repairs decreased perineal wound complications (wound disruption and purulent discharge) [43] (see also Chapter 30).

Suture

Absorbable synthetic materials should be used for all layers of the repair. Compared with catgut (plain or chromic), absorbable synthetic sutures (e.g., Vicryl) used for perineal repair decreased women’s experience of short-term (3-day) pain, less need for analgesia, reduced rate of suture dehiscence up to day 10, and less need for resuturing at <3 months. There is no significant difference in long-term pain or dyspareunia experienced by women [44]. More women with catgut sutures required resuturing compared with synthetic sutures, while more women in the standard synthetic suture group required suture removal compared with the rapidly absorbed group. Clinical experience has shown suture removal is necessary <5% of the time when using 3–0 or finer sutures and performing subcuticular skin closures [44]. Polyglyactin (Vicryl) and polydioxanone (PDS) have similar outcomes [45].

Anal Ultrasound

Anal endosonography with clinical examination immediately after delivery in nulliparous women with second-degree lacerations detected more sphincter tears than clinical examination. Anal endosonography with immediate repair of these tears is associated with less severe fecal incontinence at 1 year compared with clinical examination only [46].

Perineal Pain Control

Nonsteroidal Anti-Inflammatory Drugs

There are no RCTs to accurately assess the effectiveness of oral NSAIDs for perineal pain control. However, clinical experience shows a reduction in perineal pain. Rectally administrated NSAIDs appear to provide effective pain relief in postpartum women. Rectal indomethacin or diclofenac are associated with less pain up to 24 hours after birth and less requirement for additional analgesia in the first 24 hours and 48 hours postpartum [47]. No information is available on pain experienced >72 hours after birth or other outcomes of importance such as the impact on daily activities, resumption of sexual intercourse, and the impact on the mother-baby relationship. More studies are needed to assess the acceptability of this route of administration and comparison to oral NSAIDs.

Topical Anesthetics

Compared with placebo, topical anesthetics applied to the perineum are associated with similar pain relief up to 24 hours to 72 hours postpartum, but women are more satisfied after administration of an anesthetic [48]. Epifoam (1% hydrocortisone acetate and 1% pramoxine hydrochloride in the mucoshesive foam base) use is associated with less additional analgesia, while lignocaine/lidocaine showed no difference with regard to additional analgesia use compared with placebo [48]. A lidocaine/prilocaine cream appears to be as effective as local mepivacaine injection for pain relief during repair [49]. Compared with indomethacin vaginal suppositories, topical anesthetics have similar mean pain scores.

Therapeutic Ultrasound for Perineal Pain

There is insufficient evidence to evaluate the use of ultrasound in treating perineal pain and/or dyspareunia following vaginal delivery. Women treated with active ultrasound for acute perineal pain are more likely to report improvement in pain, less pain at 10 days and 3 months, but more likely to have bruising at 10 days compared with placebo [50]. Additionally, women with persistent perineal pain or dyspareunia treated with ultrasound are less likely to report pain with sexual intercourse.

Anesthesia

Epidural is commonly used during labor for pain control. Spinal, epidural, paracervical block, or general anesthesia may be used if complications arise in the management of retained placenta, intractable PPH, uterine inversion, or assisted vaginal delivery (Chapter 11).

Breast-Feeding

Hypertension and headache are associated with misoprostol and ergotamine use. There are no other known complications with breastfeeding after use of other uterotonics. There are no differences in breastfeeding or onset of jaundice with AMTSL.

Delivery Note

If a complicated delivery occurs with possible fetal compromise, a detailed note should address the pertinent and immediate neonatal status, including Apgar scores, umbilical cord pH, and base deficit (if obtained), and assessment of fetal heart testing prior to delivery.
PART 2—CARE OF THE JEHOVAH’S WITNESS PREGNANT WOMAN

Key Points
- Members of the Jehovah’s Witness faith refuse blood transfusions due to their beliefs that accepting them violates God’s law, and would lead to excommunication and eternal damnation.
- Refusal of blood can lead to an increased risk of maternal (and at times therefore fetal) death, especially in cases of obstetric hemorrhage.
- All obstetrical providers are responsible to ask each woman at her first prenatal visit (or preconception) if she has any objection to receiving any blood product in case of necessity.
- The woman’s wishes should be respected, following the principle of autonomy.
- After counseling, the patient should be asked to sign the consent for blood products, as well as the Health Care Power of Attorney.
- The third stage should be managed actively, and PPH prevented as much as possible. The use of cell saver is safe and effective in pregnancy.

Historic
“That is why I have said to the sons of Israel: ‘No soul of you must eat blood and no alien resident who is residing as an alien in your midst should eat blood.’” (Holy Bible: Leviticus 17:12). Charles Russell started the Jehovah’s Witness Christian sect in Pennsylvania in 1872.

Members of the Jehovah’s Witness faith refuse blood transfusions due to their beliefs that prohibit use of blood products, because they believe that accepting them violates God’s law, and would lead to excommunication and eternal damnation, as they take literally the statement reported in the Bible.

Complications
The risk of obstetric hemorrhage is approximately 6% in Jehovah’s Witness women [51]. The risk of maternal death is about 0.5%, a 44-fold increase compared with non-Jehovah’s Witness controls [51].

Management
There are no trials to assess the efficacy of interventions specifically in women who are Jehovah’s witnesses. Refer to the guidelines of third stage and abnormal third stage for general recommendations.

Preconception Counseling
Counsel regarding complications, and management in pregnancy.

Prenatal Care
All obstetrical providers are responsible to ask each woman at her first prenatal visit (or preconception visit) if she has any objection to receiving any blood product in case of necessity. The woman who states she would decline blood transfusion even if medically necessary and/or is a Jehovah’s Witness should be managed as follows:
- Counsel the woman and any family member present regarding the reasons and the risks of blood product refusal, including the possibility of maternal and fetal death. The counseling should be documented on the medical record. The patient might want to consult with the “elders” before signing informed refusal. Her wishes should be respected, following the principle of autonomy. A physician has the right of refusing to provide care for a Jehovah’s Witness only if an alternative caregiver agrees to accept and care for the patient [52].
- After counseling, the patient should be asked to sign the consent for blood products (Figure 9.2), as well as the Health Care Power of Attorney [52]. These two consents should be kept in the medical record chart and available at labor and delivery. Approximately 39% of Jehovah’s Witness pregnant women accept a variety of donated blood products, and 55% accept either intraoperative normovolemic hemodilution or transfusion of their own blood obtained by a cell saver [53].
- Consider including a copy of this guideline in the medical record, for reference.
- Selected high-risk patients, for example, those with hemoglobin <9 mg/dL, may be considered for appropriate replacement of iron, folic acid, and erythropoietin, which can be coordinated by a maternal-fetal medicine and/or a hematologist specialist.
- A routine consult with the Maternal-Fetal Medicine service is not required, but can be considered.
- A routine ethics consult is not indicated, but can be considered in specific cases.

Antepartum Testing
No specific antepartum testing is indicated.

I hereby consent to the blood products marked below:

- ___Whole blood
- ___Fresh frozen plasma
- ___Cryoprecipitate
- ___Albumin
- ___Erythropoietin
- ___Immune globulins (blood fraction, Rh immunoglobulin)
- ___Clotting factors
- ___PolyHeme (human hemoglobin), hemopure products (bovine hemoglobin)
- ___Recombinant factors
- ___Platelet cell fractions (platelet gel)
- ___Other surgical procedures, medical tests, or current therapy
- ___Using my own blood, i.e., tagged red cells, white cells, blood patching
- ___Hemodialysis equipment (nonblood primed)
- ___Intraoperative blood salvage (cell saver)
- ___Intraoperative hemodilution

Patient’s Name ________________________________
Patient’s Signature ________________________________
Date ________________________________

Figure 9.2 Blood product consent for Jehovah’s Witness patients.
Delivery

- Consents: Upon admission and prior to the surgery/delivery, all Jehovah’s Witness patients should have signed a consent form (Figure 9.2) and the previously mentioned healthcare power of attorney. If consents have not been previously signed or are not available, the patient should be counseled and consents signed.
- The third stage should be managed actively with the goal of PPH prevention. Oxytocin, methylergonovine, 15-methylprostaglandin F2α, misoprostol, and other medical and, if necessary, surgical therapies should be employed [54].
- If an operative delivery or bleeding disorder is anticipated, the patient has a history of a low-lying placenta or a placenta previa, or if an operative delivery is to occur, a cell saver can be on standby in labor and delivery throughout the patient’s labor and delivery. Use of the cell saver is safe in obstetrics [55].

Anesthesia

An anesthesiology consult should be obtained. The anesthesiologist will review the patient’s medical record and consents, including consent or refusal for blood products.

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Intrapartum fetal monitoring
Serena Xodo and Suneet P. Chauhan

KEY POINTS
- Compared with intermittent auscultation (IA), the use of continuous electronic fetal heart rate (FHR) monitoring significantly increases the rate of operative interventions (vacuum, forceps, and cesarean delivery [CD]) for nonreassuring patterns, but it does decrease the likelihood of neonatal seizures.
- With some persistent category II or III patterns, limited evidence (one randomized controlled trial [RCT]) showed that intrapartum resuscitation with maternal oxygen, change in maternal position, discontinuation of labor stimulation and/or tocolytics, or amnioinfusion (if variable decelerations) reduce the need to proceed with emergent CD but do not reduce the likelihood of injury from intrapartum hypoxia. The use of maternal oxygen has been also associated with possible harm, but the evidence is still insufficient for a recommendation. The labor of women at risk for poor peripartum outcomes should be monitored with continuous electronic FHR tracing.
- Reinterpretation of the FHR tracing, especially knowing the neonatal outcome, should be avoided.
- There is insufficient evidence to assess computerized FHR monitoring, but the limited evidence so far is promising.
- For singletons at 36 weeks or more, fetal electrocardiogram (ECG) ST-segment analysis (STAN) used as an adjunct to conventional intrapartum electronic fetal heart-rate monitoring does not improve perinatal outcomes.
- Fetal pulse oximetry (FPO) is not associated with significant maternal or neonatal benefits compared with continuous FHR monitoring alone, except for a significant decrease in CD for nonreassuring FHR (NRFHR).

BACKGROUND
Electronic FHR monitoring during labor is the most common obstetric procedure in the United States [1]. Between 1997 and 2003 in the United States, monitoring was utilized in 84% (23,273,458) of the over 27 million births [1]. For admission tests at the beginning of labor, see Chapter 7. For meconium, see Chapter 31. For hyperstimulation and hypercontractility should be abandoned. Terms such as “asphyxia,” “hypoxia,” and “fetal distress” should not be used in the interpretation of FHR tracing.

FHR DEFINITIONS
Intrapartum fetal monitoring usually refers to monitoring of the FHR. Outside the United States, this monitoring is often referred to as cardiotocography (CTG), while the term intrapartum FHR monitoring is more commonly used in the United States. Adapted from the 2008 National Institute of Child Health and Human Development Workshop [2], the definitions of FHR pattern are described in Table 10.1 and depicted in Figures 10.1 through 10.3. The definitions were developed for intrapartum monitoring but may be applicable to antepartum monitoring. No distinction is made between short- and long-term variability [2]. Accelerations, decelerations, bradycardia, and tachycardia can be quantified by describing the nadir/zenith and the duration in minutes and seconds of the FHR change. A recurrent deceleration occurs with >50% of uterine contractions in any 20-minute period.

A three-tier system for categorization of FHR patterns has been recommended [2]. According to this classification, category I tracings are predictive of normal fetal acid–base status at the time of observation and may be followed in a routine manner. Category II tracings are indeterminate and not predictive of abnormal fetal acid–base status but require evaluation and continued surveillance and reevaluation. Category III tracings are abnormal and predictive of abnormal fetal acid–base status (Table 10.2; Figures 10.4 through 10.6). These tracings require prompt evaluation and should include efforts to expeditiously resolve the abnormal pattern. It is noteworthy that a full description of an electronic fetal monitoring (EFM) tracing requires a qualitative and quantitative description of the following: (1) uterine contractions, (2) baseline FHR, (3) baseline FHR variability, (4) presence of accelerations, (5) periodic or episodic decelerations, and (6) changes or trends of FHR patterns over time [2].

In relation to uterine activity, tachysystole is defined as >5 contractions in 10 minutes averaged over 30 minutes, whether the contractions are spontaneous or stimulated. The terms hyperstimulation and hypercontractility should be abandoned. Terms such as “asphyxia,” “hypoxia,” and “fetal distress” should not be used in the interpretation of FHR tracing.

PH DEFINITIONS
- Acidemia: increased concentration of hydrogen ions in blood.
- Acidosis: a pathologic condition marked by increased concentration of hydrogen ions in tissue.
- Hypoxemia: decreased oxygen content in blood.
- Hypoxia: a pathologic condition marked by decreased level of oxygen in tissue.
- Asphyxia: acidemia, hypoxia, and metabolic acidosis; must have all of the following: (1) umbilical arterial pH <7.00; (2) Apgar ≤3 at >5 minutes; and (3) neonatal neurologic sequelae (e.g., seizures, coma, and hypotonia) [3]. This term should be used with caution and never before birth (see also Chapter 31).

INCIDENCE
The prevalence of CD for nonreassuring FHR (NRFHR) tracing is about 3% or more, and it is increasing [4]. NRFHR is usually the second most common reason for CD, after arrest of labor [5].
The risk of CD for NRFHR is >20% in patients with moderate/severe asthma, severe hypothyroidism, severe preeclampsia, postterm, or fetal growth restriction (FGR) with abnormal Doppler studies [4].

Use of likelihood ratio suggests that fetal movement count, abnormal FHR on admission, vibroacoustic stimulation, amniotic fluid index, contraction stress test, and modified and completed biophysical profile are poor diagnostic tests to identify which patients will require emergent CD for NRFHR [4]. Umbilical artery systolic/diastolic
ratio, however, is a reliable test to predict the need for CD for NRFHR.

FETAL HEART MONITORING: CONTINUOUS ELECTRONIC FHR MONITORING VERSUS INTERMITTENT AUSCULTATION

Despite the ubiquitous use, there are concerns about the efficacy of EFM. As noted by the American College of Obstetricians and Gynecologists (ACOG) practice bulletin [6], the efficacy of monitoring is adjudicated by comparing the neonatal and infant morbidity, including seizure and cerebral palsy, or mortality averted versus the unnecessary interventions (operative vaginal delivery or CD) undertaken. Since all the randomized clinical trials (RCTs) with EFM compare it to IA, the efficacy is determined by calculating the relative risks (RR) of interventions, neonatal seizures, cerebral palsy, or death.

Compared with IA, continuous EFM (CTG) shows no significant improvement in overall perinatal death rate (RR 0.86, 95% CI 0.59–1.23), but was associated with a halving of neonatal seizures (RR 0.50, 95% CI 0.31–0.80). There was no significant difference in cerebral palsy rates (RR 1.75,
Figure 10.4 Category I. (A) Normal baseline, moderate variability, absence of late and variable decelerations, accelerations present. (B) Normal baseline, moderate variability, absence of late and variable decelerations, early decelerations present. (C) Normal baseline, moderate variability, absence of late and variable decelerations.

Figure 10.5 Category II. (A) Bradycardia with moderate variability. (B) Tachycardia. (C) Minimal variability. (D) Absent variability. (E) Marked variability. (F) Recurrent variable decelerations, minimal variability. (G) Prolonged deceleration. (H) Recurrent late decelerations, moderate variability. (I) Variable decelerations.
Figure 10.5  Continued
**Figure 10.5**  Continued

**Figure 10.6** Category III. (A) Recurrent late decelerations, absent fetal heart rate (FHR) variability. (B) Recurrent variable decelerations, absent FHR variability. (C) Bradycardia, absent FHR variability.
There was a **significant increase in cesarean sections** associated with continuous CTG (RR 1.63, 95% CI 1.29–2.07). Women were also **more likely to have an instrumental vaginal birth** (RR 1.15, 95% CI 1.01–1.33). Data for subgroups of low-risk, high-risk, preterm pregnancies and high-quality trials were consistent with overall results [6,7]. EFM is associated with 1 additional CD for every 58 women monitored continuously; 661 women need to have EFM during labor to prevent one neonatal seizure [7].

While the efficacy of EFM is debated, it is noteworthy that there are concerns regarding the 13 RCTs, which are sampled over 37,000 women, on continuous HFR monitoring versus IA [7]. Only two of these trials were of high quality [8,9], and only three trials reported data in low-risk women [8–10]. The authors of the meta-analysis on this subject [7] noted that to test the hypothesis that continuous monitoring can prevent 1 death in 1,000 births, more than 50,000 women would have to be randomized. Additionally, there are concerns that the meta-analysis combines results from RCTs published prior to the introduction of the Consolidated Standards of Reporting Trials (CONSORT) guidelines [11], and that inadequate study size may not reflect the outcomes in contemporary obstetrical practice [12].

Thus, if IA is available on the labor and delivery, it is reasonable to discuss the options with the patients [13]. Admittedly both patients and clinicians prefer continuous HFR monitoring as the method to evaluate the fetus during labor [14]. The explanations for the preference include the ease of use, the reassurance women derive from hearing the heart beat during labor, and the different value patients and clinicians place on the route of delivery and neonatal outcomes. Compared with pregnant patients and mothers, obstetricians overestimate the burden posed by cesarean, and, contrary to obstetricians, women value a newborn with permanent neurologic handicap over neonatal death [14].

Clinicians choosing to utilize IA should be aware of some of the problems associated with its use. **Continuous electronic HFR monitoring has a significantly better ability to detect acidemia** (umbilical arterial pH < 7.15 in this study) than IA: a sensitivity of 97% versus 34% and a higher positive predictive value of 37% versus 22%, respectively [15]. Continuous tracing of the FHR was not only superior with detection of respiratory and mixed acidosis but metabolic as well. It is possible that the ominous FHR patterns are poorly assessed by IA. Logistically it may not be feasible to adhere to guidelines of how frequently the heart rate should be auscultated with IA. One prospective study noted that the protocol for IA was successfully completed with only 3% of the cases [16].

Even though the use of continuous electronic monitoring of the FHR does not decrease the prevalence of cerebral palsy, it assists in determining if the injury occurred during the ante- or intrapartum periods. Review of the FHR tracing of neurologically injured newborns indicates that the majority of them had an abnormal pattern consistent with asphyxial injury prior to the onset of labor [17]. Moreover, a pregnancy with chronic fetal distress may develop superimposed acute asphyxia, in which case the impairment may be more severe than if the sentinel event and injury occurred during labor.

Not all pregnancies should be monitored with IA because **those at risk for adverse outcomes like cerebral palsy, neonatal encephalopathy, and perinatal death should be monitored with continuous FHR tracing during labor** [18]. Thus, high-risk pregnancies that underwent antepartum surveillance should not be evaluated with IA, nor should those who are likely to have CD for an NRFHR pattern [4]. These include FGR, oligohydramnios, polyhydramnios, placenta previa, postterm (≥42 weeks), multiple gestation, isoimmunized pregnancy, prior intrauterine fetal demise (IUFD), or maternal renal disease, diabetes, preeclampsia, collagen disorders, hemoglobinopathies, and cardiovascular disease. Additionally, parturients should be monitored with continuous tracing of the FHR if they have been induced or augmented, or have dysfunctional labor, tocolytics administered more than once an hour, suspected FHR abnormalities with auscultation, abnormal fetal presentation, regional anesthesia, abruption, infection, preterm labor, prior CD (vaginal birth after cesarean [VBAC] attempt), hypertonic uterus, and meconium staining of the amniotic fluid [19].

FHR monitoring should be continued until delivery. If CD is performed, internal scalp monitoring can be continued until delivery, while external monitoring can be discontinued when the abdominal preparation begins.

## REVIEWING ELECTRONIC FHR MONITORING

When EFM is utilized during labor, the nurses or physicians should review it frequently. If the patient is at low-risk pregnancy, the FHR tracing should be reviewed at least every 30 minutes in the first stage of labor and at least every 15 minutes during the second stage. The corresponding frequency for high-risk parturients is 15 and 5 minutes, respectively. The maternal pulse should be taken to make sure the FHR is indeed fetal, and not maternal. **Healthcare providers should document** that they have reviewed the tracing by a narrative note or use of comprehensive flow sheets or by placing one’s initials on the monitor strip, if it is reassuring [19]. Among low-risk patients, it is more feasible to confirm that the strips are reviewed according to the guidelines, whereas among high-risk patients, compliance during active phase, and especially during second stage of labor, is more demanding and difficult. The FHR tracing, as a part of the medical chart, should be labeled and available for review if the need arises. Alternatives like computer storage of the FHR tracing that do not permit overwriting or revisions are reasonable, as are microfilm recordings [19].

Due to the **inter- and intraobserver variability**, FHR tracing should be interpreted cautiously and preferably without knowing the neonatal outcome [20–22]. When four obstetricians, for example, examined 50 cardiotograms, they agreed in only 22% of the cases. Two months later, during the second review of the same 50 tracings, the clinicians interpreted 21% of the tracings differently than they did during the first evaluation [21]. Factors that influence the interpretation of cardiotograms include the clinician’s experience, whether the tracing is normal versus equivocal or ominous with greater agreement if the tracing is reassuring, and the time of the day, with possibly greater error at night. With retrospective reviews, the foreknowledge of neonatal outcome alters the impressions of the tracing. Given the same intrapartum tracing but opposite neonatal outcomes, the reviewer is more likely to find evidence of fetal hypoxia and criticize the obstetrician’s management if the outcome was supposedly poor versus good [22].

The positive predictive value of NRFHR for cerebral palsy is about 0.1% [8]. The **false-positive rate is extremely high** (99%) for FHR tracing and abnormal neonatal outcome, especially cerebral palsy [23].
EXTERNAL VERSUS INTERNAL FHR MONITORING

External FHR monitoring is accomplished via a Doppler ultrasound device applied to the maternal abdomen. Internal scalp electrode (scalp lead) for FHR monitoring measures the R-R interval between consecutive beats. This provides an accurate representation of FHR variability. There are no RCTs comparing these two monitoring techniques. Internal monitoring is used, in general, for an FHR that cannot be consistently assessed by external monitoring. Contraindications to internal monitoring include maternal infections such as human immunodeficiency virus (HIV), active hepatitis B or C, and fetal thrombocytopenia. Otherwise internal fetal monitoring is safe.

COMPUTERIZED FHR MONITORING (EXPERT SYSTEMS)

Measured by interobserver agreement, the reliability of electronic FHR monitoring is not very good. Computerized evaluation of the intrapartum FHT, also called Expert Systems (ES), has been evaluated to try to improve the sensitivity for perinatal morbidity and mortality of FHR monitoring (CTG), and also try to decrease false positives associated with FH monitoring.

Two RCTs comparing CTG monitoring during labor with an ES versus CTG without an ES have been published, but only one trial (n = 220) provides data for quantitative analysis. There is no strong evidence that CTG with an ES has an effect on the incidence of CD (RR 0.61; 95% CI 0.35–1.04) when compared with CTG with fetal scalp blood sampling (FSBS). There is no strong evidence supporting a reduction in the incidence of neonatal seizures (RR 0.33; 95% CI 0.01–8.09) or fetal acidaemia (RR 0.50; 95% CI 0.09–2.67) in women monitored using a CTG with an ES versus a CTG without an ES. Perinatal mortality was not reported. No fetal deaths occurred. There was no strong evidence supporting a reduction in the incidence of forceps-assisted vaginal birth (RR 0.50; 95% CI 0.05–5.43), hypoxic ischemic encephalopathy (RR 0.33; 95% CI 0.01–8.09), admission to the newborn intensive care unit (RR 0.40; 95% CI 0.08–2.02) or an Apgar less than seven at 5 minutes (RR 0.50; 95% CI 0.13–1.95) [24].

Another RCT evaluated computer analysis by the Omniview-SisPorto 3.5 system of intrapartum CTG tracings to standard clinician analysis of prerecorded cases. Prediction of abnormal umbilical artery pH was more reproducible and accurate when clinicians had access to the computerized analysis of the CTGs [25].

Therefore, while limited results look promising, there is insufficient evidence to assess if computerized evaluation of electronic FHR monitoring improves perinatal outcomes.

OTHER FETAL MONITORING TESTS

Digital Scalp or Vibroacoustic Stimulation

The presence of acceleration usually assures that the fetus is not acidic (pH < 7.20). If spontaneous acceleration is not present, and/or NRFFHR is present, digital scalp or vibroacoustic stimulation should be done to elicit an acceleration (see definition in Table 10.1). Allis clamp and scalp puncture have been used to elicit an acceleration but are less safe. Digital scalp stimulation (gentle stroking of the fetal scalp for 15 seconds) is the test with the best predictive accuracy among these four [26]. There are currently no RCTs that address the safety and efficacy of digital scalp or vibroacoustic stimulation used to assess fetal well-being in labor in the presence of NRFFHR. If the FHR increases, then labor should continue since an acceleration following fetal stimulation indicates that the likelihood of low scalp pH is 2% [26]. In the absence of an acceleration, the likelihood is 38%.

Fetal Scalp Blood Sampling

FSBS to measure scalp pH and lactate was introduced in 1962 by Saling in Berlin, Germany, as a stand-alone fetal status test, before the commercialization of EFM machines in 1968 [27]. It is important to understand fetal intrapartum physiology to assess FSBS. The fetus reacts to intrapartum decreased supply in oxygen by using anaerobic glycolysis, which leads to metabolic acidosis through conversion of pyruvate to lactate. Low pH is a combined measure of both metabolic acidosis (including base deficit) and the more labile component, respiratory acidosis [28].

At FSBS, a pH value >7.25 is considered to be normal, demonstrating a well oxygenated fetus. Values between 7.25 and 7.20 are considered subnormal, thus indicating the need to repeat the sampling within 20–30 minutes. Values of pH < 7.20 (or <7.15 in the second stage of labor) require intervention, such as intrauterine resuscitation or operative delivery. As for lactate concentration, normal values have been described as being <4.2 mmol/L. Values between 4.2 and 4.8 are defined intermediate, and values >4.8 mmol require intervention [29]. If the results are scalp pH < 7.20 or lactate >4.8 mm/L, then delivery should be accomplished expeditiously, usually by CD [30].

The technical aspects of FSBS require ruptured membranes and a cervical dilatation greater than or equal to 3 cm. An amnioscope is then placed vaginally to allow adequate visualization of the fetal head. A small blood sample is then taken from the fetal scalp. Usually 30–50 µL of blood are sufficient to perform the test. For testing lactate a much smaller amount of blood is required [28]. To be beneficial, the scalp pH machine needs to be reliable and readily available with prompt results.

There is no RCT that compares specifically FSBS versus no FSBS as primary, or even secondary, intervention for fetal monitoring intrapartum. Examining then indirect data, the latest Cochrane Review reports, in the meta-analysis on continuous CTG monitoring (with or without FSBS), that FSBS is associated with an increase in instrumental deliveries (P = 0.04 and a decrease in neonatal acidosis (P = 0.04) [31]. Retrospective studies have also failed to show a positive effect of FSBS on neonatal outcome [32,33].

The only level 1 evidence available on FSBS is that comparing pH or lactate measurement for FSBS [28]. The two RCTs [34,35], enrolled a total of 3348 women in labor having a NRFFHR pattern, and investigated maternal and fetal/neonatal infant outcomes following FSBS for pH or lactate measurement. The Cochrane Review noted that lactate sampling is more likely to be successful than pH sampling; but there were no differences between the two techniques (lactate versus pH analysis of FSBS) in terms of mode of birth, or neonatal outcomes evaluated by umbilical cord blood gases, Apgar score, encephalopathy, or admission to the neonatal intensive care unit (NICU) [29].

Given these data, as well as the several risks of FSBS [27], the ACOG has recommended against the use of FSBS [33,36]. Others have also cautioned against its use [27,30]. However, this procedure is still frequently performed in some Northern European countries (e.g., Sweden).
Fetal Electrocardiogram for Fetal Monitoring in Labor

The use of the fetal ECG has been evaluated as an adjunct to continuous electronic FHR monitoring during labor [36–43]. The use of internal monitoring with a scalp lead is mandatory to obtain ECG. In most labors, technically satisfactory cardiographic traces can be obtained by external ultrasound monitors, which are less invasive than internal scalp electrodes. One study assessed PR intervals and is insufficient to make recommendations [41]. Six RCTs assessed the ST segment [34–40]. A meta-analysis of six RCTs including 26,529 laboring singletons with cephalic presentation at 34 weeks or more reported that, compared with standard CTG only, ST analysis (STAN) plus CTG was associated with similar perinatal composite outcome (1.5% versus 1.6%; RR 0.90, 95% CI 0.74–1.10), neonatal metabolic acidosis (0.5% versus 0.7%; RR 0.74, 95% CI 0.54–1.02), admission to the neonatal intensive care unit (5.4% versus 5.5%; RR 0.99, 95% CI 0.90–1.10), perinatal death (0.1% versus 0.1%; RR 1.71, 95% CI 0.67–4.33), neonatal encephalopathy (0.1% versus 0.2%; RR 0.62, 95% CI 0.25–1.52), CD (13.8% versus 14.0%; RR 0.96, 95% CI 0.85–1.08), and operative delivery (either cesarean or operative vaginal delivery) (25.9% versus 25.1%; RR 0.93, 95% CI 0.86–1.01) [42]. The Cochrane review confirms these findings, and found an 8% decrease in operative vaginal deliveries associated with STAN [36].

As there is no difference in perinatal outcomes between STAN with CTG compared with CTG alone, this modality cannot be yet recommended for routine clinical care. Nonetheless, given heterogeneity in inclusion criteria among RCTs, issues regarding learning curve for STAN, Hawthorne effect in RCTs, different (3 vs. 4) tier classification used in RCTs, and sample size and power issues given abnormal perinatal outcomes (e.g., metabolic acidosis) are uncommon, more research is needed before the STAN technology can be deemed of no value for fetal monitoring in labor [43].

Fetal Pulse Oximetry

A normal fetal oxygen saturation (FSPO2) in labor is 35–65%. Fetal pulse oximetry (FPO) showing FSPO2 <30% for at least ≥2 minutes is associated with a higher risk for declining fetal arterial pH and metabolic acidosis. The fetal oxygen sensor lies against the fetal cheek. The use of the FPO has been evaluated as an adjunct to continuous electronic FHR monitoring during labor [44–49]. RCTs on FPO are at high risk of bias because the impractical nature of blinding participants and clinicians. In RCTs not requiring FSBS prior to study entry, there was no evidence of differences in the overall cesarean section rate between those monitored with FPO and those not monitored with FPO or for whom the FPO results were masked (RR 0.99, 95% CI 0.86–1.13). There was evidence of a higher risk of cesarean section in the group with FPO plus CTG than in the group with fetal ECG plus CTG (one study only, n = 180, RR 1.56, 95% CI 1.06–2.29). No studies reported details of long-term disability [44].

There was evidence of a decrease in cesarean section for nonreassuring fetal heart rate tracing (NRFHT) in the FPO plus CTG group compared with the CTG group (RR 0.65, 95% CI 0.46–0.90) in women ≥34 weeks without use of FSBS. There was no evidence of differences between groups in cesarean section for dystocia, although the overall incidence rates varied between the RCTs [44]. No differences are seen for endometritis, intrapartum or postpartum hemorrhage, uterine rupture, low Apgar scores, umbilical arterial pH or base excess, admission to the NICU, or fetal/neonatal death [44,45].

Given the above evidence, routine use of FPO in labor is not recommended, but can be considered in venues that do not use FSBS, to reduce rates of cesarean for NRFHT.

MANAGEMENT OF ABNORMAL FHR

The best time to evaluate the FHR tracing is after a contraction. The frequency with which the FHR should be evaluated depends on the risk status of pregnancy (see above). To correctly interpret the FHR pattern, the clinician must be cognizant of the gestational age, medications (Table 10.3) [50], prior fetal assessment, and obstetric and medical conditions. There is no evidence to support the prophylactic use of betamimetics during the second stage of labor. Compared with placebo, prophylactic betamimetic therapy is associated with an increase in forceps deliveries in one trial (Figure 10.7) [51].

Amnioinfusion

Amnioinfusion means, as the word says, infusing fluid (usually saline solution) in the amniotic cavity. Amnioinfusion has been proposed in order to prevent or relieve possible umbilical cord compression during labor either after rupture of membranes (ROM), or in cases of oligohydramnios with intact membranes. This technique consists of introducing into the uterine cavity saline or ringers lactate transcervically through a catheter in women with ROM, or transabdominally through a spinal needle when membranes are intact. Amnioinfusion can be used prophylactically (e.g., in cases of oligohydramnios), or therapeutically (e.g., when repetitive variable decelerations occur during labor). This type of decelerations is typically caused by umbilical cord compression, which occurs more frequently in case of oligohydramnios [52]. For prophylactic amnioinfusion for oligohydramnios before labor, see Chapter 19.

In the Cochrane review, therapeutic transcervical amnioinfusion for potential and suspected umbilical cord compression (mostly associated with variable decelerations) was found to significantly reduce the following outcomes: cesarean section (13 trials, 1493 participants, RR 0.62, 95% CI 0.46–0.83); FHR decelerations (seven trials, 1006 participants, RR 0.53, 95% CI 0.38–0.74); Apgar score less than seven at 5 minutes (12 trials, 1804 participants; RR 0.47, 95% CI 0.30–0.72); meconium below the vocal cords (three trials, 767 participants; RR 0.45, 95% CI 0.25–0.81); and maternal stay longer than 3 days (four trials, 1015 participants, RR 0.45, 95% CI 0.25–0.78) [53].

Transcervical amnioinfusion can be done by bolus or continuous infusion technique, with similar ability to relieve recurrent variable decelerations. Neither pumps nor warmers are necessary with amnioinfusion. In fact, the use of infusion pump during amnioinfusion significantly increases the risk of NRFFH. Either lactated Ringer’s solution or normal saline can be used to place a crystalloid solution into the uterus without altering the neonatal electrolyte balance [53].

Transabdominal amnioinfusion showed similar results, though a fewer sample of patients was involved in the trials. Mean cord umbilical artery pH was higher in the amnioinfusion group (seven trials, 855 participants; average mean difference 0.03, 95% CI 0.00–0.06) and there was a trend toward fewer neonates with a low cord arterial pH (less than 7.2 or as defined by trial authors) in the amnioinfusion group (8 trials, 972 participants, RR 0.58, 95% CI 0.29–1.14) [53]. ACOG recommends amnioinfusion for management of recurrent variable decelerations [54].
Tocolysis is a possible tool in the management of intrapartum NRFHR tracing. The rationale behind using intrapartum tocolysis is that uterine relaxation improves uteroplacental blood flow and therefore fetal oxygenation. The most commonly studied tocolytics for intrapartum fetal resuscitation are beta-mimetic drugs (terbutaline 0.25 mg intravenous [iv], hexoprenaline 10 mg iv, or ritodrine 10 mg in 9 mL saline administered iv over a 1-minute period).

There are some important safety concerns regarding using these drugs. The most serious reported side effects associated with administration of beta-agonistics are pulmonary edema and myocardial ischemia [55]. Six RCTs have been performed with the aim to evaluate the use of betamimetics as tocolytics for acute intrapartum fetal resuscitation [56–61]. When betamimetic administration was compared with no treatment, neonatal outcome seemed to improve [57,58]. Four RCTs assessed the efficacy of betamimetics by comparing these drugs with either magnesium sulfate (4.0 g iv magnesium sulfate) [59], or atosiban (6.75 mg in 4.9 mL saline administered iv over a 1-minute period) [56,60], or nitroglycerin (0.4 mg iv) [61]. Betamimetics provided a more effective tocolysis, with similar successful resuscitation rates. The tocolytic potency of a single bolus of atosiban for tocolysis in term labor, especially for spontaneous contractions, needs further research [56].

ACOG recommends the administration of tocolytic medications (e.g., terbutaline) when tachysystole is associated with FHR abnormalities [54]. The time gained with this intervention

### Table 10.3 Effect of Medications and FHR Patterns

<table>
<thead>
<tr>
<th>Medications</th>
<th>Study Design</th>
<th>Effect on FHR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Butorphanol</td>
<td>Case : control</td>
<td>Transient sinusoidal FHR pattern</td>
</tr>
<tr>
<td>Cocaine</td>
<td>Case : control</td>
<td>No characteristic changes in FHR pattern</td>
</tr>
<tr>
<td>Corticosteroid</td>
<td>RCT</td>
<td>Decrease in FHR variability with betamethasone but not dexamethasone</td>
</tr>
<tr>
<td>Magnesium sulfate</td>
<td>RCT and retrospective</td>
<td>A significant decrease in the FHR baseline and variability; inhibits the increase in accelerations with advancing gestational age</td>
</tr>
<tr>
<td>Meperidine</td>
<td>RCT</td>
<td>No characteristic changes in FHR pattern</td>
</tr>
<tr>
<td>Morphine</td>
<td>Case : control</td>
<td>Decreased number of accelerations</td>
</tr>
<tr>
<td>Nalbuphine</td>
<td>RCT</td>
<td>Decreased the number of accelerations, long- and short-term variations</td>
</tr>
<tr>
<td>Terbutaline</td>
<td>Retrospective</td>
<td>Abolishment or decrease in frequency of late and variable decelerations</td>
</tr>
<tr>
<td>Zidovudine</td>
<td>Case : control</td>
<td>No difference in the FHR baseline, variability, number of accelerations or decelerations</td>
</tr>
</tbody>
</table>

Source: Adapted from Chauhan SP and Macones GA, Obstet Gynecol, 105(5 Pt 1), 1161–1169, 2005.

Abbreviation: RCT, randomized clinical trial.

![Figure 10.7](image)
may be useful for preparing for cesarean section or operative delivery, setting up regional analgesia, transferring a woman at home or in a unit without the necessary surgical or neonatal facilities to an appropriate hospital, or reviewing the need for urgent delivery.

Intrauterine Resuscitation

In the presence of NRFHR, concomitantly with performing scalp stimulation and/or scalp pH, intrauterine resuscitation can be attempted with maternal position change, hydration, oxygenation, and/or stopping labor stimulants (i.e., oxytocin) (Figure 10.7). Intravenous fluid (IVF) bolus of 1000 mL, lateral positioning, and oxygen administration at 10 L/minute via nonbreather face mask have been studies together in one RCT, and are effective resuscitative measures to improve fetal oxygen saturation (FSpO₂) during labor [62]. The evidence is from patients with normal FHR, and at present it is a reasonable assumption that the findings are applicable to NRFHR pattern.

- Maternal position change
  Maternal position change to improve fetal status in cases of intrapartum NRFHT has not been studied separately in an RCT, but only together with IVF and oxygenations (see above) [62]. Indirect evidence for position change comes from studies on maternal positioning during normal labor. There have been two reports comparing the effects of right lateral, left lateral, and supine maternal positions on fetal oxygen status, each suggesting that lateral positioning is more favorable than a supine position [63,64]. In 2013, a Cochrane review was published on the effect of maternal positions during the first stage of labor on different important outcomes for the mother and the baby. A total of 25 trials were included, with 5218 women considered. It was found that women should be encouraged and supported to use upright and mobile positions, as it seems that they benefit in terms of: a shorter duration of first stage of labor, a reduction in the risk of cesarean birth, a less use of epidural as a method of pain relief, and a lower chance of babies being admitted to the neonatal unit [65]. However, this review does not answer directly the question whether changing maternal position could be considered as a therapeutic measure in case of NRFHT in labor.

- Hydration
  There is insufficient evidence (no trial) to assess by itself the effect of intrapartum maternal hydration on fetal status. Maternal hydration has been studied in RCTs only together with IVF and oxygenations (see above) [62]. Except for the situation where maternal hydration is necessary to correct either maternal hypotension or hypovolemia, the impact of IVFs intrapartum on NRFHT has not been established. Little is known about the effects of additional IVF volume on maternal transfer of oxygen to the fetus [62].

- Oxygenation
  Oxygen is frequently given to improve fetal status, though evidence of fetal benefits is lacking. Oxygen can be used either prophylactically, i.e., to prevent NRFHT, or therapeutically, i.e., to improve NRFHR which is already present. Only two RCTs have been done on the effects of maternal oxygen therapy for prophylaxis [66]. The evidence available does not support the use of prophylactic maternal oxygen therapy during labor; in fact, abnormal cord blood pH values (less than 7.2) were recorded significantly more frequently in the oxygenation group than the control group (RR 3.51, 95% CI 1.34–9.19) [66]. Evidence from animal studies suggests that giving oxygen to the mother raises the markers of free radical activity, which leads to edema and hemorrhage in vital organs for the fetus, such as brain and lungs [67].

No RCTs on the therapeutic use of maternal oxygen for NRFHT exist. Unfortunately, the limited data available on women in labor does not clarify whether oxygen in labor is beneficial or harmful for the fetus. Oxygen has been associated with decreased umbilical cord pH [68], increased need for neonatal resuscitation [69], and raised level of markers of free radical activity [70]. Moreover neonatal resuscitation with 100% oxygen is no longer recommended [71]. In two FPO studies, maternal administration of oxygen by simple face mask was associated with a 5% increase in fetal oxygen saturation in normally oxygenated fetuses, and 20% in hypoxemic fetuses, returning abnormal to normal level of oxygenation [72,73].

- Labor stimulant
  There is insufficient evidence (no trial) to assess by itself the effect of labor stimulant discontinuation on fetal status. Nonetheless, labor stimulants such as oxytocin should be discontinued in cases of NRFHT, as it is well known that contraction can be associated with decreased fetal oxygenation, especially too frequent.

Other Interventions

Additional steps with the management of NRFHR tracing might include (no trials available) the following:

- Tocomonitoring
  Uterine contraction assessment can be considered especially for suspected for tachysystole.

- Cervical examination
  Assessment of cervical status can be helpful to assess rapid dilation or descent, and to ensure that the umbilical cord has not prolapsed.

- Maternal blood pressure
  Maternal blood pressure monitoring may be helpful, especially among those who have received regional anesthesia. If hypotension is present in conjunction with NRFHR pattern, ephedrine or phentolamine may be utilized [6].

- Piracetam for NRFHR in labor
  Piracetam, a derivative of γ-aminobenzoic acid, is thought to promote the metabolism of the brain cells when they are hypoxic. There is not enough evidence to evaluate the use of piracetam for NRFHR in labor. Compared with placebo, piracetam is associated with a nonsignificant trend to reduced need for cesarean section, and similar incidences of low Apgar scores, or neonatal respiratory problems and signs of hypoxia [74].

- Operative delivery for NRFHR in labor
  There are no contemporary trials of operative delivery versus conservative management of suspected NRFHR. In the only old (1959) trial, there is no difference in perinatal mortality [75].

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KEY POINTS

- In every hospital providing labor and delivery services, anesthesia personnel must be available on a 24-hour basis, with the ability to perform a cesarean delivery (CD) within 30 minutes from decision.
- Intravenous (IV) pain relief is inferior to neuraxial analgesia, minimally effective, and associated with several maternal and fetal/neonatal side effects.
- Neuraxial analgesia provides the best pain relief in labor, and should be available to all laboring women upon request.
- Airway complications can occur during induction and emergence of general anesthesia as well as the postoperative period and constitute the most common cause of anesthesia related death.
- In the absence of a medical condition associated with coagulopathy, it is not necessary to obtain a platelet count before neuraxial techniques. In general, women with platelet counts of ≥75,000/µL can safely receive neuraxial analgesia.
- As there is no benefit to obstetric outcome from delaying neuraxial analgesia until an arbitrary cervical dilation has been reached, the decision of when to initiate neuraxial analgesia should be made individually with each woman.
- Patients should be counseled regarding the benefits and risks of neuraxial analgesia in labor. These include the following:
  - Neuraxial anesthesia is associated with much better pain relief compared with any other intervention for pain relief, and with lower risk of umbilical cord pH < 7.20 compared with systemic opioids.
  - Neuraxial analgesia is associated with an increased incidence of hypotension, pruritus, urinary retention, and maternal fever; increased need for oxytocin administration; increased duration of the second stage (by about 15 minutes); and increased risk of instrumental vaginal birth.
  - Rare complications of neuraxial analgesia include postdural puncture headache (PDPH), respiratory depression from opioid use, epidural or spinal hematoma or abscess.
  - Discontinuation of neuraxial analgesia in the second stage does not impact obstetric outcome.
  - Use of low doses of anesthetic medications, prophylactic prehydration, and phenylephrine or ephedrine can decrease the incidence of maternal hypotension and associated nonreassuring fetal heart rate (NRFHR).
  - Compared with the standard epidural approach, combined spinal epidural (CSE) has been shown to produce a quicker (by about 6 minutes) onset of analgesia, resulting in a lower total dose of local anesthetic over the course of the labor, achieve a lower median visual analog pain score earlier in labor, increase the incidence of maternal satisfaction, have a lower incidence of incomplete block, but is associated with an increased likelihood of maternal pruritus.
  - For CD, neuraxial anesthesia is the anesthetic technique of choice. Spinal (intrathecal) anesthesia has advantages over epidural anesthesia including quicker onset of surgical anesthesia, simplicity, lower total drug dose, and superior abdominal muscle relaxation. Compared with epidural anesthesia, the spinal technique is associated with a similar failure rate, need for supplemental intraoperative analgesia, need for conversion to general anesthesia intraoperatively, maternal satisfaction, need for postoperative pain relief and neonatal intervention.
  - Hypotension following spinal analgesia for CD can be decreased by crystalloid or colloid administration, phenylephrine, ephedrine, and lower limb compression.
  - General anesthesia for CD should be avoided if at all possible as it is associated with a threefold risk of maternal death compared with neuraxial analgesia. Risks include inability to intubate or ventilate the patient at the induction of general anesthesia as well as airway complications at emergence from anesthesia. There are no evident advantages to general anesthesia in the absence of a contraindication to a neuraxial approach.

HISTORIC NOTES

Three months after the first successful public demonstration of the anesthetic properties of ether, on January 19, 1847, Dr. James Simpson Young of the University of Edinburgh used diethyl ether as anesthesia for childbirth to a woman with a deformed pelvis [1]. Before this time, pain during labor and childbirth were seen by both medical professionals and the lay community as a necessary and indeed inseparable part of pregnancy. By 1860 anesthesia for parturition had become common practice thanks to a majority of women demanding it be a part of their medical care. The practice of obstetric anesthesia has changed markedly since. Women in labor now receive analgesia, rather than anesthesia, with the goal of enabling maternal mobility during labor. Refined anesthetic techniques for women requiring CD have substantially decreased the number of maternal deaths directly related to anesthesia.

DEFINITIONS

Pain: an unpleasant sensory and emotional experience associated with actual or potential tissue damage

Analgesia (from the Greek: “no pain”): loss of the ability to feel pain without the loss of consciousness

Anesthesia (from the Greek: “no sensation”): an induced, temporary state of analgesia, paralysis, amnesia, and unconsciousness
GENERAL COMMENTS
In 2008, 60% of women in the United States chose to receive epidural or spinal anesthesia during labor [2]. While not all laboring women desire the services of an anesthesiologist, maternal request is a sufficient medical indication for pain relief in labor [3]. Hospitals providing labor and delivery services must have anesthesia personnel available on a 24-hour basis, with the ability to perform a CD within 30 minutes from decision [4]. Availability of licensed practitioners to administer anesthetics and support vital functions in emergencies is recommended [3,4].

PHYSIOLOGY OF LABOR PAIN
Labor is associated with two sources of pain: visceral pain at T10 through L1 from uterine contractions and cervical dilatation during the first stage of labor and somatic pain transmitted by the pudendal nerve at S2 through S4 from descent and consequent pressure of the fetal head on the pelvic floor, vagina, and perineum during the second stage of labor [5]. This pain is amenable to safe intervention as the anesthesiologist can interrupt the transmission of peripheral afferents from the cervix, lower uterine segment and perineum by use of neuraxial techniques that block spinal cord transmission of labor pain.

MATERNAL PHYSIOLOGIC CHANGES
Pregnancy is associated with unique physiologic changes. As a result of these changes, after the first trimester, parturients are at increased risk for aspiration of gastric contents, have decreased oxygen reserve, and are much more likely to be difficult to intubate than a nonpregnant woman (see also Chapter 3). Maternal mortality associated with general anesthesia is estimated at approximately 32/1,000,000 live births versus 1.9/1,000,000 live births for neuraxial analgesia [5].

Pregnant patients should be positioned with left uterine displacement when supine after 20 weeks of gestational age. Left uterine displacement prevents aortocaval compression, which can result in a marked decrease in venous return and a subsequent drop in cardiac output. The ability to compensate for aortocaval compression is compromised in the presence of neuraxial analgesia or anesthesia.

When considering anesthesia or analgesia, one must take into account that pregnant women are more sensitive to sedative hypnotics, local anesthetics, and the inhaled anesthetic agents than nonpregnant women. In addition the pregnant patient should be considered two patients and occasionally the needs of one must be prioritized over the other. Informed consent should be attempted in case of an emergency, and the patient’s wishes for an unmedicated birth always respected. Successful initiation of breast-feeding is not affected by neuraxial analgesia.

NONPHARMACOLOGIC ANALGESIC TECHNIQUES
Increasingly, many pregnant women are seeking alternative approaches to labor pain relief such as water immersion, acupuncture and aromatherapy. While these techniques may not provide complete pain relief, they may diminish labor pain helping parturients delay or even avoid the use of pain relief medications if that is their goal.

Acupuncture and Acupressure
Acupuncture and acupressure are therapeutic techniques based on the concept that energy flows throughout the body and strategic placement of needles or pressure can restore its balance and provide pain relief. Theories on its mechanism of action suggest it stimulates the release of endorphins in the muscles, brain, and spinal cord.

Compared with no intervention or placebo, acupuncture for labor pain is associated with less intense pain, increased satisfaction with pain relief, and reduced use of pharmacological analgesia, in small trials. Compared with no intervention or placebo, acupressure has been associated with less pain intensity, in small trials [6]. Therefore, acupuncture and acupressure may have a role with reducing pain, increasing satisfaction with pain management and reduced use of pharmacological management.

Water Immersion
Water immersion is a form of hydrotherapy in which water is used to relieve pain during labor. A Cochrane review suggests that immersion during the first stage of labor was associated with a reduction in the use of neuraxial anesthesia compared with controls [7]. There were no difference in perineal trauma, need for assisted vaginal deliveries or cesarean deliveries. There is no evidence that immersion improves perinatal outcome, however, there are case reports of rare but serious adverse effects in the newborn including respiratory distress when practiced during the second stage of labor. Given these rare but serious risks to the newborn, it is the opinion of The American College of Obstetricians and Gynecologists and the American Academy of Pediatrics that the practice of immersion in the second stage of labor (underwater delivery) should be avoided in routine obstetrical care [8]. See also Chapters 7 and 8.

Biofeedback
Biofeedback is a therapy that allows women to gain control over their body’s physiologic response to pain by controlling heart rate, blood pressure and muscle tension. Despite some positive results shown in some very small trials, there is insufficient evidence that biofeedback is effective for the management of pain during labor [9].

Aromatherapy
Aromatherapy is the use of plant derived aromatic essential oils to promote the sense of physical and psychological well-being. The limited evidence available does not show benefit, or harm, from aromatherapy for labor pain relief compared with no aromatherapy [10].

Hypnosis
Hypnotherapy is the use of hypnosis to deeply relax the body through focused concentration. For childbirth hypnosis is often used to focus attention on feelings of comfort or numbness as well as to enhance women’s feelings of relaxation and sense of safety. In a review of six studies (1032 women) there was no significant difference between women receiving hypnosis versus controls in the use of pharmacologic pain relief [11].

Massage
There is insufficient evidence to assess the effect of massage therapy on labor pain. However a meta-analysis involving 326 women found that women who used massage felt less pain when compared with women given usual care during first stage of labor [12].

Relaxation
A review of eleven randomized controlled trials, with data reported on 1374 women, found that relaxation techniques and yoga may help manage labor pain [13].
Transcutaneous Nerve Stimulation
Transcutaneous electrical nerve stimulation (TENS) units emit low voltage electrical impulses and may be used to stimulate acupressure points. A systemic review of nine trials including more than 1000 women concluded that TENS did not reduce labor pain and did not reduce the use of additional analgesic agents [14].

SYSTEMIC ANALGESIA
Parenteral Opioids
Opioids are the most widely used of the systemic medications for labor analgesia. Opioids are low cost, easy to use, and do not require specialized personnel or equipment to be delivered safely. There are a variety of opioid agonist–antagonists in clinical use such as buprenorphine (Buprenex), butorphanol (Stadol), or nalbuphine (Nubain), or pure opioid agonists such as meperidine, fentanyl, morphine, hydromorphone, or their derivatives. Characteristics of some of these drugs are shown in Table 11.1. All provide moderate pain relief with respect to another. Opioids may be delivered intramuscularly (IM) or IV. IM administration is easy for the healthcare provider, but painful for the patient. IV administration does require an IV line be placed, but allows faster onset of analgesia, predictable magnitude of peak plasma concentration, and titrating of dose to effect. IV opioids may be delivered as intermittent boluses or via patient-controlled analgesia (PCA) technology.

Efficacy
These drugs are similarly efficacious in relief of maternal labor pain. In one small study, compared with placebo, meperidine (Demerol, pethidine) 100 mg IM in early labor is associated with a very modest (17 mm) reduction in the visual analog scale (VAS) pain score at 30 minutes [15,16]. Request for epidural analgesia is delayed to 232 minutes compared with 75 minutes for parturients receiving placebo, with no further requests for analgesia in 32% versus 4% of women receiving meperidine or placebo, respectively [16].

There is not enough evidence to definitively evaluate the comparative efficacy of the various systemic opioids used to provide labor analgesia. There are problems with the methodological quality of some trials, and lack of consistency in the way various outcomes are reported. Nonetheless, using a variety of measures, including pain relief, maternal satisfaction with analgesia, interval to delivery, and obstetric outcome, there is no evidence of significant differences between meperidine, tramadol (Ultram, Meptazinol), or pentazocine (Talwin) [15].

Maternal Safety
There is not enough evidence to definitively evaluate the comparative safety of the various systemic opioids used to provide labor analgesia. Maternal side effects are largely dose dependent rather than drug dependent. Across all opioids there exists the potential for maternal nausea, histamine release and pruritus, delayed gastric emptying, constipation, urinary retention, dysphoria, drowsiness, hyperventilation and hypotension. There is weak evidence of more frequent adverse effects such as maternal nausea, vomiting, and drowsiness with meptazinol compared with meperidine. However, both tramadol and pentazocine have fewer maternal side effects when compared with meperidine [15], making these two agents the ones with the best evidence of both moderate efficacy and lowest incidence of side effects among the opioid analgesic agents. Respiratory and neurobehavioral depression can be reversed with the use of the pure opioid antagonist naloxone. Meperidine, the first synthetic opioid to be used for labor analgesia, is metabolized to normeperidine in the liver, which can cause respiratory depression and convulsions, neither of which can be reversed by naloxone.

Fetal Safety
IV opioids cross the placenta owing to their low molecular weight and lipid solubility and as a result can affect the neonate. When compared with neuraxial analgesia, IV opioids are associated with increased incidence of NRFHR, lower fetal base excess, and decreased fetal respirations and tone at birth. Drugs with active metabolites, such as meperidine (active metabolite normeperidine), are associated with more prolonged neonatal sedation. Agonist/antagonists such as nalbuphine can result in both neonatal cardiac and respiratory depression although there is no data that demonstrates adverse outcomes. Naloxone is a pure opioid antagonist and is the drug of choice in treatment of neonatal respiratory and neurobehavioral depression secondary to opioid agonist agents. Repeated doses might be necessary, but excess use can be associated with neonatal withdrawal seizures. Compared with neuraxial analgesic techniques, systemic medications are much less effective at decreasing visual analog pain scores (see Efficacy and Advantages in the section “Epidural Analgesia”).

### Table 11.1 IV or IM Opioid Agents for Maternal Pain Relief in Labor

<table>
<thead>
<tr>
<th>Agent</th>
<th>Usual Dose (mg)</th>
<th>Frequency (Every Hour)</th>
<th>Onset (Minutes)</th>
<th>Neonatal Half-Life (Hours)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tramadol</td>
<td>50–100 mg</td>
<td>4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pentazocine</td>
<td>30 mg IV or IM</td>
<td>3–4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Meperidine</td>
<td>25–50 IV</td>
<td>1–2</td>
<td>5</td>
<td>13–22</td>
</tr>
<tr>
<td></td>
<td>50–100 IM</td>
<td>2–4</td>
<td>30–45</td>
<td>&gt;60</td>
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<tr>
<td>Fentanyl</td>
<td>50–100 µg IV</td>
<td>1</td>
<td>1</td>
<td>5</td>
</tr>
<tr>
<td>Nalbuphine</td>
<td>10 IV or IM</td>
<td>3</td>
<td>2–3 (IV)</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>15 (IM)</td>
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<td>15 (IM)</td>
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<td></td>
<td>1–2 (IV)</td>
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<tr>
<td></td>
<td>10–30 (IM)</td>
<td></td>
<td></td>
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<tr>
<td>Butorphanol</td>
<td>1–2 IV or IM</td>
<td>4</td>
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<td>Morphine</td>
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<tr>
<td></td>
<td>10 IM</td>
<td></td>
<td>30–40 (IM)</td>
<td></td>
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</tbody>
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Abbreviations: IV, intravenous; IM, intramuscular.
PERIPHERAL NERVE BLOCKS

Paracervical Block
During the first stage of labor, pain from cervical dilation (but not uterine contractions) can be blocked by a paracervical block. There are no trials to assess the effectiveness of paracervical block in labor. Risks include maternal local anesthetic toxicity from IV injection, fetal local anesthetic toxicity from inadvertent fetal injection, and a strong association with fetal bradycardia.

Pudendal Block
During the second stage of labor, perineal pain can be blocked with a pudendal block. There are no trials to assess the effectiveness of pudendal block in labor. There is a small risk of maternal local anesthetic toxicity in the event of accidental intravascular injection.

Lumbo-Sacral Sterile Water Injection
For patients that desire nonpharmacologic analgesic techniques, back pain during the first stage of labor can be reduced with lumbo-sacral sterile water injection. In a randomized controlled trial studying 128 term parturients, women receiving sterile water injection versus acupuncture reported greater pain relief and higher degrees of relaxation during labor [17]. In another study, women with severe lower back pain, four injections of sterile water, either 0.1 mL intracutaneously or 0.5 mL subcutaneously in the lumbar–sacral region, has been shown to be effective in reducing severe back pain [18].

NEURAXIAL (REGIONAL) LABOR ANALGESIA (EPIDURAL, SPINAL, OR CSE)

Background
Epidural, spinal, and combined spinal–epidural analgesia techniques are the most effective methods of providing pain relief to the laboring parturient. A large meta-analysis looking at over 300 studies comparing neuraxial analgesic techniques to a variety of other techniques including systemic opioids, inhaled anesthetics, and nonpharmacologic concluded that neuraxial techniques are superior to all other methods in terms of parturient pain relief [19]. In a jointly published opinion, both the American College of Obstetricians and Gynecologists and the American Society of Anesthesiologists (ASA) state: “in the absence of a medical contraindication, maternal request is a sufficient medical indication for pain relief during labor” [20]. As labor results in severe pain for many women, neuraxial anesthesia, when available, should be offered to all women in labor, assuming no medical contraindication.

Overview of Techniques
Neuraxial labor analgesia is also commonly called regional analgesia. It can be provided via the epidural space, the intrathecal (spinal) space, or both.

Medications injected into the epidural space have a relatively slow analgesic onset of 8–15 minutes. Block height is determined by the volume of medication injected into the epidural space. Injecting low concentration local anesthetic produces analgesia and injecting high-concentration local anesthetic produces anesthesia.

By contrast, medication injected into the intrathecal space has a quick onset of 2–5 minutes. Block height is determined primarily by the baricity (density relative to cerebrospinal fluid [CSF]) of the injectate and, to some extent the total volume of drug.

Epidural Analgesia

Technique
The epidural space (Figure 11.1) is located using a loss of resistance (to air or saline) technique [21]. A small catheter is placed into the epidural space through which local anesthetics and opioids can be intermittently injected (bolus) or given as a continuous infusion.

The most common local anesthetic choices include low-concentration, long-acting amides such as bupivacaine and ropivacaine in combination with a lipid soluble opioid such as fentanyl and sufentanil. The addition of opioids to local anesthetics produces an additive effect that results in a lower concentration of local anesthetic being required to produce adequate labor analgesia. A lower concentration of local anesthetic increases maternal mobility and decreases the potential for maternal systemic toxicity. There is insufficient evidence comparing concentrations, type of anesthetic used, and several other technical aspects of epidural anesthesia in labor to recommend one specific drug combination [22,23].

The epidural infusion rate can be controlled either by the anesthesiologist or by the patient. Patient-controlled epidural analgesia (PCEA) typically employs a background infusion of local anesthetic and opioid together with enabling the parturient to augment the infusion with a bolus dose every 10–15 minutes. This technique can also be used with a spinal catheter with appropriately reduced doses.

**Indications**

The primary indication for neuraxial analgesia is maternal request during labor [5]. However there are numerous maternal conditions that make neuraxial anesthesia the prudent choice. Examples of such conditions include anticipated difficult intubation, history of malignant hyperthermia, high risk for CD, the presence of a comorbidity that would benefit from the reduced catecholamine levels that result from adequate labor analgesia (e.g., selected respiratory or cardiac disease) and prophylaxis against autonomic hyperreflexia in the woman with a high spinal cord lesion. Functioning epidural catheters can also be used to provide anesthesia for instrumented delivery and extraction of a retained placenta.

In some cases, an epidural catheter can be intentionally inserted into the intrathecal space to provide continuous spinal analgesia. However, the resulting PDPH incidence may be high. Nonetheless, this risk may be acceptable in certain situations such as the presence of certain maternal comorbidities (typically cardiac), the morbidly obese parturient with an airway that appears extremely difficult or impossible to intubate, or following an inadvertent dural puncture during a difficult epidural placement.

**Efficacy and Advantages**

Compared with nonpharmacologic techniques or to IV opioids, epidural analgesia provides significantly better pain relief [24]. In >85% of cases, the pain relief is optimal. As labor results in severe pain for many women, and neuraxial analgesia is the most effective intervention to decrease or eliminate this pain, neuraxial analgesia should be offered, when available, to all women in labor (assuming no medical contraindication).

Compared with systemic opioids, epidural anesthesia is associated with superior pain relief and lower risk of umbilical cord pH <7.20. There is no evidence of a significant difference in the risk of CD (relative risk [RR] 1.10, 95% confidence interval [CI] 0.97–1.25 27 trials, 8417 women) [24]. In addition there is no evidence of a significant difference in the risk of long-term backache, low neonatal Apgar scores at 5 minutes, and other maternal and neonatal outcomes. No studies reported on rare but potentially serious adverse effects of epidural analgesia [15]. Neuraxial analgesia is the least depressant method of analgesia for the fetus/neonate.

There is no reason to delay epidural placement until an arbitrary cervical dilation has been reached. Data suggest that neuraxial analgesia can be offered as early as 2 cm or less without adversely affecting labor outcome or incidence of CD. Epidural placement at ≤2–5 cm is associated with similar maternal (instrumental delivery, CD, etc.) and neonatal outcomes compared with epidural placement at ≥3–5 cm in women in spontaneous labor or receiving oxytocin [25–29]. Therefore, the decision of when to place epidural analgesia should be made individually with each woman [5].

PCEA gives the patient control over her analgesia. Patients receive less local anesthetic overall (and therefore have less motor blockade) and receive fewer manual boluses from the anesthesiologist (“top offs”) [22,30]. As patient satisfaction is high with epidural analgesia, PCEA is not associated with additional improvement in maternal satisfaction.

There is insufficient evidence to support the hypothesis that discontinuing epidural analgesia late in labor (e.g., at the beginning of the second stage) reduces the rate of instrumental delivery. There is evidence that it increases the rate of inadequate pain relief in the second stage of labor [31].

**Safety, Disadvantages, and Complications**

Compared with systemic analgesia, epidural analgesia is associated with an increased need for oxytocin administration, increased duration of the second stage by an average of 15 minutes, increased risk of operative vaginal birth (forceps or vacuum-assist), and increased risk of mild maternal fever, hypotension, and urinary retention. (RR 1.42, 95% CI 1.28–1.57) [24]. A recent noninferiority trial assessed the effects of routine epidural analgesia during labor versus epidural upon maternal request suggests that routine epidural analgesia was associated with a statistically significant increase in the rate of operative deliveries (difference 8.1%, 95% CI −0.01 to 16.3) [32].

The etiology of the occasional increase in maternal temperature associated with epidural use is uncertain. This effect can lead to increased maternal and neonatal antibiotic treatments, as well as neonatal sepsis evaluations, but with no difference in the corresponding rate of neonatal infection or sepsis [33]. In centers with structured protocols for neonatal sepsis workups, there are no increases in the incidence of neonatal sepsis workups in babies born to mothers with epidural analgesia.

The incidence of maternal hypotension following neuraxial analgesia during labor is approximately 14%. Hypotension is associated with an increased incidence of NRFHT that rarely (1%–2%) necessitates CD. Left uterine displacement should be maintained whenever possible as it will increase maternal preload and increase uterine perfusion. Measures such as fluid preloading and concurrent administration of phenylephrine or ephedrine can mitigate or prevent maternal hypotension.

Although the risk of long-term backache in women who utilize epidural analgesia is similar to controls (IM meperidine), it is quite common for a woman to experience soreness or tenderness at the site of epidural insertion for 2–3 days [34].

Disadvantages of epidural analgesia include a slower onset compared with intrathecal (spinal) injection, incomplete blockade of pain (in about 10%–15% of patients), and inadvertent intrathecal or intravascular catheter placement. While the PCEA gives the patient more control and requires less intervention from anesthesiologists, this can lead to underdosing and therefore inadequate analgesia: for example, if the patient does not self-administer a bolus (if she is asleep), or if there is a pump malfunction.

Other complications include a 1%–2% chance of PDPH following accidental lumbar puncture, and rarely, epidural hematoma, respiratory arrest due to the unrecognized injection...
of an epidural dose of opioid into the intrathecal space, systemic local anesthetic toxicity due to unrecognized IV injection, or total spinal anesthesia from an unrecognized intrathecal injection of a dose of local anesthetic meant for the epidural space. See sections “Spinal Analgesia” and “Anesthetic Emergencies” for further details on these complications and strategies for their prevention and treatment. Women should be counseled about these risks before labor [33].

Preloading with IV fluids to prevent hypotension: Preloading with IV fluids (500–1000 mL, or weight-based formula 10–20 mL/kg) prior to traditional high-dose local anesthetic blocks may have some beneficial fetal and maternal effects in healthy women [35]. Using low concentration analgesic solutions for epidural techniques, preloading with IV fluids is associated with no significant difference in maternal hypotension, although only a very large effect was excluded. There is, however, a trend toward less frequent FHR abnormalities [35], therefore some fluid preloading may be beneficial.

Propylactic ephedrine to prevent NRFHR: Compared with no ephedrine, ephedrine 10 mg IV bolus, followed by a 20 mg continuous infusion over 60 minutes, started in the first minutes after the epidural test dose, significantly decreases the incidence of NRFHT from 15% to 3% [36]. This evidence though is insufficient to make a recommendation.

Spinal Analgesia

Technique

Using a small-bore spinal needle (typically a pencil point design that minimizes trauma to the dura thereby reducing the incidence of PDPH), a dural puncture is made and proper location confirmed by CSF aspiration (Figure 11.1). A single injection of an analgesic dose of an opioid such as fentanyl, or sufentanil, with or without a local anesthetic such as bupivacaine or ropivacaine, is administered.

Indications

Indications for single injection spinals include advanced labor where delivery is imminent, forceps deliveries in women without epidurals, and for patients with retained placentas.

Advantages

Advantages for single injection spinals include ease and speed of onset, completeness of block, and lower incidence of PDPH compared with epidural analgesia.

Disadvantages and complications

Disadvantages for single injection spinals are inability to redose.

Postdural puncture headache. PDPH occurs with a frequency of approximately 1%–2% after either spinal or CSE analgesia when administered using a small-gauge pencil-point needle. PDPH is thought to be caused by leak of CSF through the punctured dura. The leak of CSF and subsequent decreased spinal fluid pressure leads to downward traction or stretch on the meninges with resulting symptoms. PDPH is characterized by a positional frontal-occipital headache that is exacerbated by the upright position (gravity worsening stretch on the meninges), with improvement in symptoms when the patient is supine. Diploplia, tinnitus, nausea, and vomiting caused by stretch on the cranial nerves are also common symptoms. Although pencil-point design spinal needles have significantly reduced the incidence of PDPH following spinal analgesia, there are no proven interventions to prevent PDPH following inadvertent dural puncture with an epidural needle. Conservative early interventions include analgesics, supine positioning, caffeine, and hydration. In about 1/3 of cases, the headache persists and is severe enough to require an epidural blood patch procedure. An epidural blood patch is performed by injecting 10–25 mL of autologous blood into the patient’s epidural space at the level of the dural puncture using meticulous sterile technique. If the level of the dural puncture is unknown, a more caudal interspace should be chosen. After the procedure, the patient should rest in the supine position for 1–2 hours. Resolution of symptoms with blood patch occurs in 70%–85% of women. Expected side effects following epidural blood patch include backache and leg pain [37].

Respiratory depression or arrest. Respiratory depression or arrest due to intrathecal opioids occurs rarely (1 in 5,000–10,000 patients). Naloxone reverses this complication and should be readily available, along with airway management equipment when administering labor analgesia.

Hematoma. Hematoma after epidural or spinal analgesia is an extremely rare complication of neuraxial anesthesia (1/1,500,000–250,000). Symptoms include bilateral leg weakness, urinary incontinence and back pain. Prolonged motor paralysis without regression of block should raise suspicion. If suspected the patient should undergo prompt definitive imaging (MRI) of her neuraxis, followed by surgical decompression after diagnosis. Surgical decompression within 6 hours following the onset of symptoms often prevents permanent neurologic injury. The risk of persistent neurologic injury from epidural is about 1/240,000 [21].

Combined Spinal Epidural

Technique

The epidural space is identified with loss of resistance technique (Figure 11.1). A spinal needle is then introduced into the intrathecal space. An intrathecal dose of local anesthetic and opioid is injected through the spinal needle, which is then removed, leaving the epidural needle in place. An epidural catheter is inserted and an epidural local anesthetic and opioid infusion is started. The intrathecal dose generally lasts about 2 hours, after which the epidural catheter will provide continuous analgesia.

Indications

A CSE technique provides the benefit of immediate onset of spinal analgesia coupled with the indefinite duration of an epidural catheter technique. The CSE technique is particularly useful for women in advanced labor requesting pain relief. There are data that associate CSE with an increased rate of cervical dilatation and shorter length of labor compared with both IV opioid and epidural analgesia [38–40]. Indications for a CSE are the same as those for both epidural and spinal techniques.

Advantages

Both CSE and epidural techniques provide effective pain relief in labor. The type of opioid and concentration of local anesthetic used in the CSE or epidural technique impact maternal mobilization and other outcomes more than the technique itself [41]. There appears to be little basis for offering CSE over epidurals in labor, with no difference in overall maternal satisfaction despite a slightly faster onset (about 6 minutes) with CSE, and less pruritus with epidurals. There is some evidence that CSE compared with an epidural is associated with a faster
rate of cervical dilatation in nulliparous women less than 5 cm [39], and women close to imminent birth may benefit from the speed of onset a CSE offers. There is no difference in ability to mobilize, obstetric outcome, or neonatal outcome [41]. A recent review revealed however there is a statistically significant difference in risk of instrumental delivery between CSE and epidural analgesia with CSE having decreased rates of instrumentation (RR 0.81; 95% CI 0.67–0.97) [41].

Disadvantages
Disadvantages of CSE are similar to epidural and spinal techniques. Although data is limited, no differences are reported between IV preloading and no preloading to prevent hypotension [35]. Women who receive intrathecal opioids experience **more pruritus** than with a standard epidural [41]. However, pruritus is very common after both spinal and epidural opioids and results from &mu;-opioid receptor stimulation in the brainstem. Naloxone or nalbuphine are effective interventions. Diphenhydramine and other antihistamines are not effective in treating opioid-induced pruritus because opioids administered via the intrathecal or epidural route do not cause histamine release. The significantly higher incidence of urinary retention, instrumental deliveries and rescue analgesia interventions with traditional high concentration epidurals would favor the use of low-dose epidurals [41]. It is not possible to draw any meaningful conclusions as to possible differences between CSE and epidural in producing rare complications such as nerve injury and meningitis.

Contraindications to Regional Anesthesia

**Coagulopathy**
A routine platelet count is not necessary before administering neuraxial analgesia in the healthy parturient. Indications for platelet count may include hypertensive disorders of pregnancy, idiopathic thrombocytopenia purpura (ITP), abortion and disorders of coagulation. In the absence of DIC, a platelet count of ≥100,000/mL is considered safe. Several studies have also confirmed that platelet levels of 50,000–99,000/mL are not associated with higher risk of complications [3]. Women receiving low-dose aspirin are not at increased risk for complications from neuraxial blocks. Women receiving prophylactic unfractionated heparin, regardless of dosing schedule, should have the medication held for at least 4 hours, ideally 6 hours, before placing a neuraxial block or removing an epidural catheter. Women receiving therapeutic unfractionated heparin are candidates for regional analgesia if the activated partial thromboplastin time (aPTT) is normal. If the aPTT is elevated, protamine may be used to reverse the heparin effect. Women receiving low–molecular weight heparin are not candidates for regional analgesia for 12–24 hours from last dose, given a higher rate of epidural hematoma with placement during this period. Therefore, consideration should be given to converting women who require anticoagulation from low molecular to unfractionated heparin as they approach term.

The coagulation status of parturients with medical conditions associated with coagulopathy such as the HELLP (hemolysis, elevated liver enzyme values, and low platelet levels) syndrome should be thoroughly evaluated before initiating a neuraxial block. Patients with platelets less than 75,000/mL should be examined for stigmata of coagulopathy (easy bruising, bleeding from the IV site, and so on) before instrumentation. A prothrombin time, a partial thrombin time, and a platelet count should all be reviewed before proceeding. A fibrinogen level and a d-dimer level are useful to assess the presence of DIC if suspected. A bleeding time is not indicated. Patients with known platelet dysfunction including those on antiplatelet medication (e.g., clopidogrel) should not receive neuraxial analgesia. Aspirin therapy is considered an acceptable risk [42].

**Infection**
**Systemic.** Patients with suspected meningitis (bacterial or viral), sepsis, or viremia should not receive neuraxial blockade. Patients with suspected chorioamnionitis can receive neuraxial analgesia/anesthesia following the administration of appropriate IV antibiotics. Chronic herpes simplex virus (HSV) outbreak is not a contraindication to neuraxial techniques. However a primary outbreak places the parturient at risk for herpetic viremia and the theoretical risk for central nervous system (CNS) infection and should be weighed against the risk of alternative methods of analgesia. Human immunodeficiency virus (HIV)/acquired immunodeficiency syndrome (AIDS) is not a contraindication to spinal or epidural anesthesia, or epidural blood patch.

**Localized.** Patients with localized skin or soft tissue infections should not be instrumented at those sites.

ANESTHESIA AND MATERNAL COMORBIDITIES

**Hypertensive Disorders**

**Advantages of Neuraxial Analgesia**
Pregnant women with hypertension may benefit from neuraxial analgesia, as it may improve uterine perfusion through several pathways (localized neuraxial vasodilatory effect, reduced catecholamine release). **Neuraxial analgesia is the analgesia of choice in hypertensive pregnant women as it allows clinicians to avoid the possibility of difficult intubation and the severe hypotension that accompanies endotracheal intubation.**

**Disadvantages**
Patients with gestational hypertension, preeclampsia, and eclampsia are at increased risk for hemodynamic instability during both labor and surgical anesthesia. Neuraxial techniques can be used safely with increased vigilance for maternal hypotension. In one focused review, marked hypotension after spinal in women with preeclampsia did not occur, lending support to the safety of spinal anesthesia in these women [43]. Women with preeclampsia may have thrombocytopenia, which increases the risk of neuraxial hematoma formation. Most anesthesia providers consider a platelet count above 75,000/mm³ to be adequate for the administration of neuraxial anesthesia.

**Cautions**
Caution must be taken in fluid management in this patient population as there is altered vascular leaking, decreased oncostatic pressure, and a higher incidence of pulmonary edema. Also, there can be an exaggerated hypertensive response to ephedrine and phenylephrine. The prevention, rather than treatment, of hypotension has been associated with better outcomes for the fetus. Women with severe preeclampsia who must undergo general anesthesia are at risk for an exaggerated hypertensive response to intubation and often benefit from pretreatment with an antihypertensive such as labetalol immediately prior to induction. Treatment with magnesium sulfate for preeclampsia/eclampsia can potentiate neuromuscular blockade in patients receiving general anesthesia, so care must be taken when using nondepolarizing muscle relaxants.
Maternal Cardiac Disease
Heart disease in the parturient is the leading cause of maternal mortality outside of obstetric complications. Understandably, the risk increases with severity of maternal disease. The normal changes in maternal cardiac physiology resulting from pregnancy can either unmask subclinical or worsen clinical cardiac disease. Every effort should be made to care for these patients in facilities equipped to provide the multidisciplinary approach that is required to ensure the best possible outcome for mother and neonate. Goals of anesthetic management for parturients with heart disease are: (1) maintain a normal heart rate, sinus rhythm, and adequate SVR; (2) maintenance of intravascular volume and venous return; (3) avoidance of aortocaval compression; and (4) avoidance of myocardial depression during general anesthesia.

Valvular Heart Disease
Patients with acquired valvular disease (rheumatic fever, mitral valve prolapse, artificial valves, and endocarditis) are at increased risk for arrhythmias, pulmonary edema, and cardiac ischemia from the increased heart rate, cardiac output, metabolic demand, and decreased oxygen reserve associated with pregnancy and the pain of labor. Patients with arrhythmias or artificial valves may also be on heparin or low-molecular weight heparin.

Advantages of neuraxial analgesia. Epidurals and CSE block the pain and stress of contractions therefore reducing tachycardia and increased cardiac output. Ablation of the bearing down reflex can be advantageous in patients with aortic or mitral regurgitation.

Disadvantages. Hypotension is the largest disadvantage with neuraxial analgesia; even transient hypotension can lead to coronary hypoperfusion, ischemia, arrhythmias, or arrest. This is especially dangerous in patients with moderate to severe aortic stenosis. However, parturients with aortic stenosis have safely undergone both neuraxial labor analgesia and anesthesia for CD. Meticulous anesthetic technique and allowing adequate time to slowly administer the requisite medication is the key to safe provision of neuraxial anesthesia in these patients. Inadvertent IV injection of local anesthetics may drink clear liquids for up to 2 hours before induction of anesthesia for planned CD [3].

Congenital Heart Disease
Women with congenital heart disease (e.g., tetralogy of Fallot, hypoplastic left ventricle, transposition of the great vessels, and septal defects) are now surviving to childbearing years. Depending on the adequacy of their surgical repair, pregnancy may or may not severely impact these patients with underlying cyanotic heart disease. The increased cardiac output, oxygen consumption, changes in systemic and pulmonary resistance, and aortocaval compression can exacerbate preexisting right to left shunts increasing the risk of maternal cyanosis and death.

Advantages of epidural analgesia. Although the hypotension of large-dose spinal anesthesia can be associated with risk of shunting and cyanosis, slowly administered epidural, low-dose spinal, or continuous spinal analgesia are advantageous to these patients by reducing catecholamine levels and preventing maternal expulsive reflexes. Additionally, if an instrumented delivery or CD is required, a surgical anesthetic level can be slowly produced, avoiding the risks of general anesthesia in these patients.

Disadvantages. The largest disadvantage of neuraxial analgesia is the risk of hypotension. Hemodynamic management of these patients should be aggressive and tailored to the underlying cardiac defect.

Previous Lumbar Surgery
Previous lumbar surgery (e.g., discectomy, placement of Harrington rods) is not a contraindication for lumbar epidural or spinal analgesia or anesthesia. One case series found that successful block can be achieved, although at a lower rate (55%) than in the control population. There were no cases of spine infection, low back pain, or headaches [44].

Maternal Obesity
The incidence of maternal obesity has been rapidly increasing worldwide. Obese parturients have higher rates of hypertension, diabetes, preeclampsia, chorioamnionitis and cesarean section compared with normal weight women. Obese patients are also more at risk for hyperventilation and apnea after administration of opioids, complicating postpartum and postoperative pain control. Relative immobility also increases the risk of thromboembolic events [45]. Most importantly, obesity increases the risk for death during pregnancy. The report of Confidential Enquiries into Maternal Deaths in the United Kingdom for the 2006–2008 triennium showed that 49% of maternal deaths were overweight or obese [46]. The risks of low Apgar scores, neonatal intensive care unit admission, fetal death, and perinatal death are increased [47]. Respiratory, gastrointestinal, and anatomical changes of pregnancy significantly increase the risk of a failed intubation and aspiration, which is as high as 15%–33% in obese pregnant women. Venous access can be difficult increasing the need for central venous access. Bony landmarks and increased adipose and soft tissue can complicate placement of neuraxial analgesia and anesthesia with initial epidural failure as high as 42%. Despite this increase in technical difficulty, neuraxial anesthesia is the technique of choice in the obese parturient.

ANESTHESIA FOR CESAREAN DELIVERY
Current guidelines recommend a fasting period for solids of 6–8 hours prior to scheduled CD. Parturients without complications may drink clear liquids for up to 2 hours before induction of anesthesia for planned CD [3].

Epidural
Epidural anesthesia can be used for CD, and for most urgent or emergent deliveries if a functioning catheter is in place. Patients with existing labor epidurals can have their block extended to an adequate level for surgery using a large volume of high-concentration local anesthetic solution containing opioids and epinephrine (which is thought to produce analgesia via α2 receptors in the spinal cord).

Advantages
There are numerous advantages to using epidural anesthesia for CD, including avoiding instrumentation of the maternal airway, the ability to redose the epidural in the event of a delayed or prolonged surgery, the ability to administer long-acting epidural opioids to augment postoperative pain control and the avoidance of a dural puncture. Opioids administered via neuraxial approach are associated with a 35%–55% incidence of maternal pruritus severe enough to require treatment [48]. An additional advantage is that the patient is awake and can experience the birth. The epidural catheter should be...
removed after 24 hours to reduce urinary retention, pruritus, and infection risks.

Disadvantages
Disadvantages of epidural for CD include longer onset time for surgical block compared with spinal anesthesia and the possibility of incomplete or patchy block, making epidural anesthesia a less attractive option than spinal anesthesia in emergent situations (when an epidural catheter is not already in place). The higher doses of local anesthetics used in epidural blocks (compared with spinal) increase the risk of maternal systemic toxicity.

Spinal
Spinal anesthesia can be used for both planned cesarean deliveries and in most emergencies. Intrathecal opioids can be added to augment postoperative pain control.

Advantages
Spinal anesthesia is a reliable form of anesthesia that is technically easier to perform and produces adequate anesthesia significantly faster than epidural anesthesia [49]. Other advantages are its simplicity, lower drug doses, and superior abdominal muscle relaxation. Compared with epidural, spinal technique is associated with similar failure rate, need for additional intraoperative analgesia, need for conversion to general anesthesia intraoperatively, maternal satisfaction, need for postoperative pain relief, and neonatal intervention [49]. Compared with epidural, spinal anesthesia for cesarean section is associated with reduced time, by about 8 minutes, from start of the anesthetic to start of the operation.

Compared with general anesthesia, women with neuraxial anesthesia have less intraoperative blood loss but significantly more nausea. In a review of randomized clinical trials, no differences in umbilical cord arterial blood pH were found among general and neuraxial anesthetic techniques [50].

Disadvantages
Hypotension, possibly profound, is increased in incidence 23% with spinal versus epidural anesthesia [49]. A number of strategies can be used to mitigate hypotension. No intervention reliably prevents hypotension due to spinal anesthesia for CD, but five interventions have been found to reduce the incidence of hypotension: (1) crystalloid preload, (2) preemptive colloid administration, (3) ephedrine, (4) phenylephrine, and (5) lower limb compression [35,51,52]. No differences were detected for different doses, rates or methods of administering colloids or crystalloids. High doses of ephedrine may increase the incidence of hypertension and tachycardia and is associated with fetal acidosis of uncertain clinical significance. Patients who receive intrathecal bupivacaine with prophylactic IV phenylephrine infusion have less hypotension than those without phenylephrine [51]. Newer studies have shown that phenylephrine is as safe as ephedrine; in fact, fetal pH is higher and the incidence of maternal nausea is lower with phenylephrine [53]. Given the efficacy of phenylephrine and better umbilical cord pH many anesthesia providers now use phenylephrine as a first line agent for the treatment and prevention of maternal hypotension. Different methods of compression appeared to vary in their effectiveness. In summary, interventions such as crystalloids, colloids, ephedrine, phenylephrine, or lower-leg compression can reduce the incidence of hypotension, but none have been shown to eliminate maternal hypotension during spinal anesthesia for CD.

Limited duration is another disadvantage of spinal anesthesia. CSEs and spinal catheters combine the advantages of the rapid onset of spinal anesthesia with the ability to redose in the case of prolonged surgical time.

The rate of PDPH after spinal anesthesia ranges between 1.5% and 11.2%. To reduce this risk, this technique should be performed using a small-gauge (24 g or smaller), pencil point spinal needle when possible.

In summary, both spinal and epidural techniques are shown to provide effective anesthesia for cesarean section. Both techniques are associated with moderate degrees of maternal satisfaction. Spinal anesthesia has a shorter onset time, but treatment for hypotension is more likely if spinal anesthesia is used. Because of its safety and effectiveness, spinal anesthesia has evolved as the regional technique of choice for CD due in particular to rapidity of anesthetic onset, quality of anesthesia, and ease of performance of block.

General Anesthesia
While neuraxial techniques are clearly the preferred anesthetic for CD, general anesthesia is indicated in the case of failed regional anesthesia, an obstetric emergency preventing placement of a neuraxial technique, a contraindication for regional anesthesia, or objection by the patient to regional anesthesia.

Precautions
If a general anesthetic is chosen, patients must receive aspiration prophylaxis that may include a nonparticulate oral antacid within 30 minutes of surgery and/or metoclopramide. Time permitting, an H2 blocker can be given 30–50 minutes before induction of anesthesia, to confer additional protection. Airway protection with endotracheal intubation is mandatory.

There is insufficient evidence to assess prevention of aspiration during general anesthesia. When compared with no treatment or placebo, there is a significant reduction in the risk of intragastric pH < 2.5 with antacids, H2 antagonists, and proton pump antagonists [54]. H2 antagonists are associated with a reduced risk of intragastric pH < 2.5 at intubation when compared with proton pump antagonists, but compared with antacids the findings were unclear. The combined use of antacids plus H2 antagonists is associated with a significant reduction in the risk of intragastric pH < 2.5 at intubation when compared with placebo or compared with antacids alone (RR 0.12, 95% CI 0.02–0.92, 1 trial, 119 women). In general, the quality of the evidence is insufficient to make a recommendation. None of the studies assessed potential adverse effects or substantive clinical outcomes [54].

Advantages
There are few advantages to general anesthesia in the absence of a contraindication to a neuraxial approach. One possible advantage is the relaxation properties halogenated anesthetics have on uterine muscle. This property can be useful in the management of uterine inversion, fetal entrapment, or retained placenta. IV nitroglycerine and terbutaline are other options in these situations.

Disadvantages
Compared with neuraxial anesthesia, general anesthesia is associated with a threefold risk of maternal death [55]. The greatest risk is from the inability to intubate or ventilate the patient. Parturients have increased upper airway edema, decreased pulmonary functional residual capacity, increased metabolic oxygen consumption, decreased lower esophageal
sphincter tone, and delayed gastric emptying. These conditions increase the risk of both hypoxemia and aspiration. Airway edema can also make anesthetizing the airway for awake fiberoptic intubation more difficult. The incidence of failed intubation in obstetric population is approximately 1 in 300, nearly eight times that of the general population [55,56].

CD is considered a high-risk procedure for intraoperative recall. Concern about neonatal depression and uterine atony has led to minimal use of benzodiazepines and low dose intraoperative halogenated anesthetics intraoperatively. Compared with nonobstetric surgery, the risk of maternal awareness under general anesthesia is increased (0.4% vs. 0.2% for nonobstetric surgery) [57]. Although benzodiazepines and opioids can depress the fetus, they are pharmacologically reversible and administration should be considered if their use would benefit the mother. For instance, the judicious use of preintubation opioids can be considered to blunt the sympathetic response to labor. However, following major surgery, such as cesarean hysterectomy, the effectiveness of continuous epidural anesthesia provides effective镇静和镇痛作用. However, following major surgery, such as cesarean hysterectomy, the effectiveness of continuous epidural anesthesia provides effective pain relief in the first 12–24 hours [5]. An alternative is PCEA, which can be associated with increased maternal motor weakness. However, following major surgery, such as cesarean hysterectomy, the effectiveness of continuous epidural anesthesia may justify the potential for increased maternal motor weakness.

IV patient-controlled opioids are another reasonable alternative, using morphine, hydromorphone hydrochloride, or fentanyl.

Oral analgesia with oxycodone-acetaminophen 5/325 mg two tablets every 3 hours for 12 hours and then one or two tablets every 4 hours as needed is associated with superior pain control and fewer side effects compared with morphine patient-controlled IV analgesia in one trial [59].

The transversus abdominis plane (TAP) block is performed by injection of local anesthetic between the internal oblique and transversus abdominis muscles to block the plexus of nerves supplying the anterior abdominal wall. In a randomized controlled trial comparing intrathecal opioid to TAP block after CD, intrathecal opioid was superior and patients receiving TAP block had increased opioid consumption in the immediate postoperative period [60]. The TAP block is a reasonable alternative in a patient with a contraindication to a neuraxial block [61].

After 24 Hours
Nonsteroidal anti-inflammatory drugs reduce maternal opioid consumption after CD, even in the first 24 hours, and should be the main intervention for pain control after the first 24 hours. Use of oral narcotics should be quickly weaned.

ANESTHETIC EMERGENCIES
An anesthetic emergency in the obstetric patient necessitates clear and precise communication and cooperation between the anesthesia and obstetric teams. The goal is to stabilize the mother while, if necessary, safely and quickly delivering the neonate.

Total Spinal
A total spinal occurs with cephalad spread of local anesthetic to the breathing centers of the brainstem. This can result from unintentional intrathecal placement of an epidural dose of local anesthetic or from subdural catheter placement with subsequent migration of the catheter. Agitation, difficulty speaking, and profound hypotension are signs of a total spinal. Control of the airway with endotracheal intubation, blood pressure support with fluid, vasopressors and left uterine displacement should be performed immediately. Once the airway has been secured, assessment of the fetus should be facilitated. If the fetus is stable, delivery can safely await maternal recovery.

Local Anesthetic Systemic Toxicity (LAST)
IV injection of local anesthetic can lead to systemic toxicity including seizures and cardiovascular collapse. The mother's airway should be controlled immediately and delivery of the fetus is often indicated because of maternal instability. Seizures can be quickly terminated with administration of diazepam. The pharmacologic treatment of LAST is different from other cardiac arrest scenarios. Management of cardiac arrest includes treatment according to the American Heart Association (AHA)/Advanced Cardiac Life Support (ACLS) guidelines with adjustment of medications and possibly prolonged effort as per American Society of Regional Anesthesia and Pain Medicine (ASRA) guidelines [62]. Administration of intralipid, a 20% fat emulsion, has been shown to increase the survival rate of patients who experience cardiac arrest secondary to local anesthetic systemic toxicity [63].

Failed Intubation
The risk of failed intubation is increased in the parturient at approximately 1 in 300 nearly 8 times that of the general population (1:2330) [55,56]. Increased edema in the upper airway, increased breast size, and increased friability of the mucosa increase chance of failure. In patients where intubation was difficult, it is important to note that emergence is an equally high-risk event. This is emphasized by the number of anesthetic deaths in parturients involving loss of airway, the majority of which occurred during emergence from anesthesia or in the postanesthesia period [64]. Open communication between the obstetric team and anesthesia team is crucial and all decisions should incorporate multidisciplinary communication and cooperation.
Maternal Hemorrhage, Resuscitation, and Massive Transfusion

Maternal hemorrhage can lead to exsanguination and the need for massive transfusion, defined as the need for 10 or more units of packed red blood cells (PRBC) in 24 hours. Volume resuscitation with crystalloid, nonbiologically active colloid, and PRBCs can lead to a dilutional coagulopathy necessitating the transfusion of clotting factors and platelets. Recent retrospective studies from both military and civilian trauma have shown improved outcomes using an empiric ratio of FFP:RBC (fresh frozen plasma:red blood cells) 1:1 in settings that require massive transfusion. Cryoprecipitate may also be needed in patients with low fibrinogen levels. Parturients may acquire a dilutional thrombocytopenia, thus transfusion of platelets may also be warranted; however, the optimal ratio of platelet units to other factors is not known. Risks of massive transfusion include transfusion reactions, viral infection, fluid overload, pulmonary edema, and transfusion related acute lung injury (TRALI) [65]. Intraoperative cell salvage is an addition to the above armamentarium and is gaining acceptance. The ASA Practice Guidelines on Obstetric Anesthesia recommend “in case of intractable hemorrhage when banked blood is not available or the patient refuses banked blood, intraoperative cell salvage should be considered if available” [3].

Cardiopulmonary Resuscitation in the Pregnant Patient

Cardiac arrest during late pregnancy occurs in approximately 1 in 30,000 pregnancies. ACLS protocols apply to pregnant women with a few important adjustments. Pregnant women should be intubated promptly to facilitate oxygenation and protect the airway from aspiration. Left uterine displacement is essential to relieve aortocaval compression. The AHA states the resuscitation team leader should consider delivery of the fetus after 4 minutes to improve cardiopulmonary resuscitation of the mother by relieving aortocaval compression [66]. The AHA further notes improved survival for infants occurs when the delivery occurs no more than 5 minutes after maternal cardiac arrest [66].

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Operative vaginal delivery

Adeeb Khalifeh

KEY POINTS

- Vacuum- and forceps-assisted deliveries have the same indications. There are no circumstances where operative vaginal delivery (OVD) is definitely indicated. Alternatives, including allowing the patient to labor longer, oxytocin augmentation, and cesarean delivery (CD), should always be considered.
- When used by experienced operators, OVD is safe for both mother and baby and effective in obtaining vaginal delivery, with forceps having slightly higher success rates.
- Forceps achieve a vaginal delivery more often than vacuum, 91% versus 86%, respectively. Complication rates differ between vacuum and forceps, with the predominant differences being that maternal third- and fourth-degree perineal (14% vs. 7.5%) and vaginal wall (26% vs. 8%) injuries are more common with forceps-assisted delivery. Neonatal facial injury is uncommon with operative delivery, 1.7% with forceps, and 0.2% with vacuum. The choice of instrument is decided after appropriate counseling and depends also on operator experience.
- There is insufficient evidence to compare different types of forceps.
- Soft vacuum cups fail at attaining vaginal delivery more often than by rigid cups but have a lower rate of significant fetal scalp trauma. Rigid cups may be better for occiput posterior and other more difficult deliveries, while soft cups may be better suited for less complicated, routine deliveries.
- If attempted, delivery with a vacuum should ideally be achieved within 5 minutes from vacuum application and, in general, should be discontinued if the vacuum cup pops off the fetal head three times.
- Attempting to use a different extraction instrument after failing with one should be avoided due to increased incidence of fetal injury.

HISTORICAL PERSPECTIVES

OVD has been practiced for centuries. Its initial function was fetal extraction during prolonged dysfunctional labor in an attempt to preserve the life of the laboring women. The invention of modern forceps can be traced back to the Chamberlain family in Europe during the sixteenth century. Vacuum extraction was first described by Dr James Yonge in 1705. The modern evolution of vacuum delivery (also called ventouse) can be attributed to Malmström’s metal cup vacuum system developed in 1954. OVD has evolved significantly and today implies a mechanism for facilitating vaginal delivery of a healthy infant while minimizing maternal risk.

In modern obstetrics, OVD, whether forceps or vacuum, is used to expedite safe vaginal delivery for maternal or fetal indications. Despite wide variations in different countries, OVD has a crucial role in safely avoiding primary CD. Emphasis should be placed on indications, contraindications and prerequisites for an OVD with instrument choice based on clinical circumstances and operator skill to minimize maternal and neonatal morbidity.

INCIDENCE

Rates of OVD (forceps and vacuum) have been declining in the United States. The rate of OVD decreased from 9.01% in 1992 to 3.30% in 2013. In 2013, forceps-assisted deliveries accounted for only 0.59% of live births [2].

INDICATIONS

Both forceps and vacuum have the same indications. Use should depend mostly on proper evaluation of the patient’s labor, risk factors, clinical pelvimetry, estimated fetal weight, and operator experience. There are no circumstances where OVD is definitely indicated. Alternatives, including allowing the patient to labor longer, oxytocin augmentation, and CD, should always be considered. OVD is usually considered for the following [3]:

- Maternal:
  - Prolonged second stage: At least 3 hours in nulliparous women and at least 2 hours in multiparous women (longer duration on individualized basis, e.g., with the use of epidural analgesia) [4].
  - Shortening of the second stage of labor for maternal benefit (e.g., underlying medical condition precluding pushing).
  - Inefficient maternal effort (e.g., exhaustion or underlying medical condition precluding pushing).
- Fetal:
  - Suspicion of fetal compromise.

“Elective” forceps delivery, that is, without an indication, is associated with increased maternal perineal trauma, and given the other potential maternal and neonatal complications, should not be preferred to spontaneous vaginal delivery [5].

CONTRAINDICATIONS TO OVD

Contraindications to OVD are listed in Table 12.1. Severe scalp trauma and unexplained active bleeding may be relative contraindications in individual cases. OVD should be used with extreme caution in women with maternal diabetes, prolonged labor, and fetal macrosomia, with appropriate preparations due to an increased risk of shoulder dystocia.

RISKS FOR FAILED OVD

Risks for failed vacuum or forceps vaginal delivery include increased maternal age, increased body mass index, diabetes, polyhydramnios, African-American race, induction of labor, occiput posterior (also increases rates of third- and fourth-degree perineal lacerations), dysfunctional labor, and prolonged labor [6].
Table 12.1 Contraindication to Operative Vaginal Delivery

- Nonvertex presentation
- Unengaged fetal head
- Unknown fetal head position
- Fetal prematurity such as <34 weeks (vacuum)
- Known fetal coagulation disorders (e.g., hemophilia and NAIT)
- Known fetal bone demineralization conditions (e.g., osteogenesis imperfecta)

Abbreviation: NAIT, neonatal alloimmune thrombocytopenia.

CLASSIFICATION OF OVD

- Outlet: Scalp is visible at introitus without separating the labia, fetal skull has reached pelvic floor, sagittal suture is anteroposterior (AP) diameter or right or left occiput anterior or posterior position, fetal head is at or on perineum, and rotation ≤45° [3].
- Low: Leading point of the fetal skull is at station ≥+2 cm and not on the pelvis floor, rotation is ≤45° (left or right occiput anterior to occiput anterior or left or right occiput posterior to occiput posterior), or rotation is >45°.
- Mid: Station is above +2 cm but head is engaged.

Originally devised for forceps, this classification is valid for any OVD, including vacuum [7].

TYPES OF FORCEPS

There are many different designs for forceps, but all consist of two separate halves that each have the same four basic components: blade, shank, lock, and handle. There is insufficient evidence to compare different types of forceps and it is recognized that the choice is often subjective. In the only small trial performed, severe facial abrasion was decreased from 4.1% to 1.9% from the regular forceps compared with soft forceps, but unfortunately successful delivery rates for the two different forceps were not reported, and the soft forceps were self-made. Given the paucity of data, choice of forceps type is operator dependent.

- Classical forceps: These have cephalic and pelvic curvatures. Usually indicated when no rotation of the fetal head is necessary before delivery. Common types include the following: Simpson forceps (fenestrated blades and nonoverlapping shanks), Tucker–McLane forceps (nonfenestrated blades and overlapping shanks), and Elliot forceps (fenestrated blades, overlapping shanks, and largest cephalic curvature). Many of these forceps have been modified with a Luikart pseuddafenestration of the blade.
- Rotational forceps: These have cephalic curvature but lack a pelvic curvature. Also have a sliding lock to allow forceps to slide to correct asymclitism of the fetal head if present. After rotation of the fetal head is accomplished, classical forceps should be used to complete the delivery. Types include Kielland, Luikart, Barton, and Salinas forceps.
- Forceps for breech delivery: These are indicated to help with the aftercoming head in a breech delivery. These forceps lack a pelvic curvature and have blades that are beneath the plane of the shank. Types include Piper and Laufe forceps.

TYPES OF VACUUM EXTRACTORS

Vacuum extractors were originally designed with a rigid metal cup. Subsequently, soft cups have been developed. Several types of rigid (metal or plastic) and soft (silicone plastic or rubber) vacuums are in clinical use [9–11]. Among different types of vacuums, the metal cup is more likely to result in a successful vaginal birth than the soft cup (9% vs. 17%), with more cases of scalp injury (41% vs. 30%) and cephalohematoma (14% vs. 8%). The handheld ventouse is associated with more failures than the metal ventouse, and a trend to fewer than the soft ventouse [9–12].

Rigid cups may be better for occiput posterior and other more difficult deliveries, while soft cups are better suited for less complicated, routine deliveries [9]. Maternal injury, low Apgar scores at 1 or 5 minutes, umbilical artery pH < 7.20, hyperbilirubinemia/phototherapy, retinal/intracranial hemorrhage, and perinatal death do not differ between soft and rigid vacuum cups [9,12]. Soft vacuum cups have largely replaced the rigid cup in routine clinical practice. For a comprehensive review of vacuum delivery see Vacca, 2009 [13].

Possible Complications of Operative Vaginal Deliveries

1. Maternal
   - Forceps use is associated with a sixfold increase in third- and fourth-degree perineal tears compared with a spontaneous vaginal delivery [14].
   - Vacuum use is associated with a twofold increase in third- and fourth-degree lacerations compared with spontaneous vaginal deliveries [14].
   - Forceps delivery has an increased risk of anal sphincter injury compared with vacuum delivery [9].
   - Urinary, flatus, liquid and solid incontinence is similar at 1 year in women who had OVD compared with women who had a second stage CD [15].
   - Pelvic floor and sexual function do not differ at 1 year postpartum compared with women who had CD [16].
   - In the absence of anal sphincter injury, anal incontinence rates at 5–10 years are similar to those in women who had a spontaneous vaginal delivery [17].

2. Neonatal
   - Intracranial hemorrhage rate is increased in OVD, but the absolute risk is low [18].
   - The rates of intracranial hemorrhage and neonatal encephalopathy compared with second stage CD are similar [18,19].
   - Cephalohematoma, fetal scalp lacerations, retinal hemorrhages, subgaleal hematoma, and intracranial hemorrhages have been reported in vacuum deliveries.
   - Facial lacerations, facial nerve palsy, and corneal abrasion are more common with forceps delivery.
   - Long-term cognitive outcomes are similar to spontaneous vaginal deliveries [20,21].

COMPARISON OF FORCEPS VERSUS VACUUM-ASSISTED DELIVERY

Safety/Complications

Maternal
- There is a trend for less regional (37% vs. 40%) and significantly less general anesthesia (1% vs. 10%) with vacuum compared with forceps-assisted deliveries. We do not use general anesthesia for operative delivery [9].
• Third- and fourth-degree perineal [14% vs. 7.5%; relative risk (RR) 1.89, 95% confidence interval (CI) 1.51–2.37] and vaginal wall lacerations (26% vs. 8%; RR 2.48, 95% CI 1.59–3.87) (maternal trauma) are significantly increased with forceps compared with vacuum.

• Severe perineal pain at 24 hours is decreased (9% vs. 15%) with vacuum compared with forceps-assisted deliveries.

• Flatus incontinence or altered continence is more common with forceps compared with vacuum in a small trial (59% vs. 33%; RR 1.77, 95% CI 1.19–2.62). In one randomized controlled trial (RCT) comparing forceps and vacuum delivery there was no difference in either bowel or urinary dysfunction 5 years postpartum [22].

• Rates of moderate/severe pain at delivery, endoanal ultrasound abnormalities [22], and other maternal outcomes are similar.

• Rotational forceps does not increase adverse maternal outcome compared with rotational vacuum delivery [23].

Fetal/Neonatal

• Facial injury is more likely with forceps (1.7% vs. 0.2%; RR 5.10, 95% CI 1.12–23.25).

• Using a random effects model because of heterogeneity between studies, there was a trend toward fewer cases of cephalohematoma with forceps (5% vs. 9%; RR 0.64, 95% CI 0.37–1.11). Rates of cephalohematomas occur in about 10% versus 4% in vacuum and forceps, respectively [9]. Cephalohematomas have an overall rate of 2.5% in the general population [24]. However, the diagnosis of cephalohematoma can be falsely positive in up to 75% of cases [25].

• Retinal hemorrhages trended to be less common in forceps than vacuum deliveries (5% vs. 8%; RR 0.68, 95% CI 0.43–1.06) [11]. One study found retinal hemorrhages occurred in 18% in spontaneous vaginal deliveries [26]. The clinical significance of retinal hemorrhages remains unclear [24].

• Rates of scalp/face injury other than cephalohematoma, use of phototherapy, perinatal death, readmission to hospital, and hearing or vision disability are similar between forceps and vacuum-assisted vaginal deliveries [9,27].

• Other possible uncommon fetal complications associated with OVD include facial nerve injury, corneal abrasions, facial bruising, and lacerations. Very rare findings include facial nerve palsy, skull fractures, cervical spine injury, and intracranial hemorrhage. With vacuum-assisted delivery, life-threatening neonatal injuries include subgaleal (subaponeurotic) hematoma (0%–4%). Intracranial hemorrhage is a rare complication of OVD (0%–2.5%).

Efficacy

Both vacuum- and forceps-assisted delivery have high delivery success rates (with vacuum from 83% to 94% and with forceps from 78% to 92%) [9,25]. Forceps are less likely than the vacuum to fail to achieve a vaginal birth with the allocated instrument (9% vs. 14%; RR 0.65, 95% CI 0.45–0.94) [9].

CD occurred in 4.5% of forceps, and 2.6% of vacuum deliveries, as in these studies unfortunately failed vacuum would at times be followed by forceps, which, in general, should not be done [9].

Alternatives to Operative Vaginal Delivery

Alternative management should always be discussed with the patient. This includes continued expectant management for prolonged second stage in the presence of a reassuring fetal status, oxytocin augmentation or proceeding with a CD. However, the appropriate use of OVD is encouraged by American Congress of Obstetricians and Gynecologists (ACOG) to safely prevent the primary CD [4].

Summary

There is a recognized place for forceps and all types of vacuum in clinical practice [9]. The role of operator training with any choice of instrument must be emphasized. The increasing risks of failed delivery with the chosen instrument from forceps to metal cup to handheld to soft cup vacuum, and trade-offs between risks of maternal and neonatal trauma (see Table 12.2) need to be considered when choosing an instrument. Overall forceps or the metal cup appears to be most effective at achieving a vaginal birth, but with increased risk of maternal trauma with forceps and neonatal trauma with the metal cup [9].

MANAGEMENT

Preoperative Assessment

Counseling

Review with the patient the indication, risks (possible complications) and benefits, and type of instrument to be used for OVD. The option of CD, including its risks and benefits, should also be reviewed. Obtain verbal or written informed consent prior to OVD.

Preparation/Documentation

Maternal. Sufficient analgesia, clinical assessment of pelvis, lithotomy position, ± empty bladder (Table 12.3).

Fetal. Vertex presentation, head engaged (lower part of bony vertex—not caput—at or lower than level of ischial spines), knowledge of the position of the head, asynclitism, and estimated fetal weight. Suboptimal instrument placement increases maternal and neonatal morbidity [28]. In a recent RCT, the use of ultrasound significantly decreased the incidence of incorrect diagnosis of fetal head position [29].

Table 12.2  Comparison of Forceps vs. Vacuum for Operative Vaginal Delivery

<table>
<thead>
<tr>
<th></th>
<th>Fetal</th>
<th>Maternal</th>
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<tbody>
<tr>
<td>Favors Forceps</td>
<td>Favors Vacuum</td>
<td>Favors Forceps</td>
</tr>
<tr>
<td></td>
<td>Less third- and fourth-degree perineal tears (75% vs. 14%) and vaginal wall (8% vs. 26%) lacerations</td>
<td>Less likely to fail to deliver the baby vaginally (9% vs. 14%)</td>
</tr>
<tr>
<td>Less facial injury (0.2% vs. 1.7%)</td>
<td>Less severe perineal pain postpartum (9% vs. 15%)</td>
<td></td>
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Uteroplacental: Cervix completely dilated, ruptured membranes, absence of placenta previa, or other contraindications.

Other: Alert nursing, anesthesia, and neonatology of plan for OVD. Be prepared for possible shoulder dystocia. Willingness to discontinue the procedure if it does not proceed as planned [2].

Episiotomy
There is insufficient evidence (no trials) to assess the benefits and risks of episiotomy in operative deliveries. Lateral episiotomy has been shown to be protective against anal sphincter injuries in vacuum deliveries, compared with mediolateral and median episiotomies, in a meta-analysis [30]. In a large observational study, mediolateral episiotomy protected significantly for anal sphincter damage in both vacuum extraction (odds ratio [OR] 0.11, 95% CI 0.09–0.13) and forceps delivery (OR 0.08, 95% CI 0.07–0.11), compared with no episiotomy. The number of mediolateral episiotomies needed to prevent one sphincter injury in vacuum extractions was 12, whereas five mediolateral episiotomies could prevent one sphincter injury in forceps deliveries [31,32]. Episiotomy should not be routinely performed, as it is associated with perineal lacerations in nonoperative vaginal deliveries. However, episiotomy should not be used as an obstetric quality measure in assisted vaginal deliveries, as this could decrease further the use of OVD and increase CD rate [33].

Antibiotic Prophylaxis
Antibiotic prophylaxis cannot be recommended solely for the indication “OVD.” Two grams cefotetan IV at the time of vacuum or forceps delivery are associated with a nonsignificant decrease (0% vs. 3.5%) in endometritis [34].

Vacuum Application
Vacuum application, if performed, should begin with low suction and be slowly increased to vacuum of about 0.7–0.8 kg/cm² (500–600 mmHg). Compared with stepwise negative pressure for vacuum delivery, rapid negative pressure application is associated with reduced duration (by 6 minutes) of vacuum procedure, but no other maternal or perinatal effects, in a small trial [23]. No torque or rocking motions should be applied to the vacuum. Traction should only be in the direct line of the vaginal canal. The risk of cephalohematoma increases as the time of vacuum application increases. There is no evidence that reducing pressure between contractions decreases risk of fetal injuries [35]. Most deliveries are achieved with one to three pulls with increased neonatal trauma (45%) after three pulls [36]. The risk of cephalohematoma is increased after 5 minutes of vacuum application [37].

Failed Operative Delivery
Attempting to use a different extraction instrument after failing with one should be avoided, as cephalopelvic disproportion may be present, and the highest incidence of neonatal intracranial hemorrhage, as well as other neonatal injuries, is highest among infants delivered using forceps and vacuum sequentially [27,38–40]. At that point, cesarean is usually offered and performed.

Postpartum
It is essential to examine carefully both the fetus and the maternal perineum after OVD.

REFERENCES
Cesarean delivery

A. Dhanya Mackeen

See also specific chapters on Anesthesia (Chapter 11), Trial of Labor after Cesarean Section (TOLAR) (Chapter 14), and Postpartum Care (Chapter 30).

KEY POINTS

- A cesarean delivery on maternal request (CDMR) or a planned repeat CD without other indication should not be performed before 39 weeks.
- Prophylactic antibiotics should be administered before every CD. The evidence suggests that a single dose of cefazolin or ampicillin intravenous (IV) be given 30–60 minutes prior to skin incision.
- All women undergoing CD should receive mechanical venous thromboembolism (VTE) prophylaxis with either pneumatic compression devices or compression stockings. These should be applied preoperatively and continued until full ambulation.
- Left lateral tilt is associated with a lower incidence of hypotension.
- Skin should be cleansed with chlorhexidine-alcohol immediately prior to skin incision.
- Compared with no scrub, vaginal irrigation with povidone-iodine (p-i) immediately before CD significantly reduces the incidence of postcesarean endometritis.
- Adhesive drapes for CD are associated with a higher incidence of wound infection.
- A transverse skin incision with either the Joel–Cohen or Pfannenstiel technique is the preferred one for CD.
- Routine development of a bladder flap may not be necessary.
- The uterine incision should be performed with the scalpel in a transverse fashion and expanded cephalo-caudal bluntly with fingers.
- Tranexamic acid (TA) should be used for prevention of postpartum hemorrhage.
- Carbetocin, when available, is superior to oxytocin for prevention of postpartum hemorrhage.
- Uterine massage, associated with cord traction, is associated with less blood loss.
- Spontaneous placental removal should be preferred to manual removal given the significant decrease in blood loss and endometritis.
- The uterus should be repaired with sutures with full-thickness bites, in a continuous fashion.
- Compared with two (double) layer closure, one (single) layer of suture for uterine incision repair is associated with a statistically significant reduction in mean blood loss; duration of the operative procedure; and presence of postoperative pain; but also with poorer healing and thinner residual myometrium. It might be reasonable to omit the second layer if the woman is planning no more pregnancies (e.g., receives tubal ligation). For women planning future pregnancies, the uterus can be closed in two layers.
- There is no evidence to justify the time taken and the cost of peritoneal closure, so avoid it.
- The evidence supports routine subcutaneous suture closure in women with a subcutaneous tissue depth ≥2 cm.
- Suture closure of the skin incision is associated with a significant decrease in wound complications, especially separation, compared to staple closure; it is therefore the preferred closure method for the transverse skin incision.
- Gum chewing three times/day for at least 30 minutes each time after CD is associated with earlier return of bowel sounds (by 5 hours), earlier passage of flatus (by about 5 hours), and less (12% vs. 21%) symptoms of mild ileus.
- There is insufficient evidence for a strong recommendation, but early oral fluids and even food after CD seem to be safe and possibly beneficial.
- Reclosure of the disrupted laparotomy wound of CD is associated with success in >80% of women, faster healing times, and fewer office visits.

HISTORIC NOTES

The word cesarean is probably derived either from the “Lex Regia,” later called “Cesarea,” which allowed the postmortem abdominal delivery of the child in ancient Rome, or from the Latin “caesare,” which means “to cut.” Until the late 1800s, most CDs were done after maternal death, for attempt at fetal salvage. In 1882, the era of modern CD began when Saenger advocated closing all uterine incisions immediately after surgery. The lower uterine segment incision was introduced by Kronig in 1912 and popularized in the United States by DeLee in 1922. The transverse uterine incision was described by Munro-Kerr in 1926 [1]. CD has been associated with relatively low maternal mortality for about 100 years. Safety has improved in the last 50 years, as the above techniques have become more widely used, and antibiotics have been introduced.

DIAGNOSIS/DEFINITION

Birth via abdominal route by laparotomy.

EPIDEMIOLOGY/INCIDENCE

CD is now the most common surgical procedure in the United States with over 1 million performed each year. Its incidence has increased to 33% of deliveries in the United States in 2013 [2]. This increase has been fueled at least in part by the increased incidence of multiple gestations and decreased incidences of vaginal births after CD and vaginal breech deliveries. Women’s demand for scheduled CD (CDMR) has increased as cesarean complications diminish, women have fewer children, and fear or concerns about vaginal delivery do not abate.
INDICATIONS

Common accepted indications for CD are failure to progress (aka failure to dilate, failure to descend, suspected cephalopelvic disproportion [CPD], dystocia, etc.), nonreassuring fetal heart rate tracing (NRFHT), nonvertex presentation, etc. (Figure 13.1). See Chapters 7, 8, 10, and 24.

There is insufficient evidence (lack of any trial) to assess the benefits and risks of a policy of CDMR (only indication: woman's desire; also called elective CD, a term which should be avoided) compared with trial of labor in women with term singleton gestations in cephalic presentation. The most common reason for a request for CDMR is fear of labor pain. CDMR should not be motivated by the unavailability of effective pain medication. There is also no randomized controlled trial (RCT) to evaluate the obstetrician's recommendation for CD without indication. As there are no such trials, there is insufficient evidence to assess the long-term maternal and neonatal morbidity and mortality of planned CD versus trial of labor. There is insufficient evidence on the impact of counseling during pregnancy with the aim of reducing the incidence of CDMR, but counseling reduces anxiety and concerns related to pregnancy preparation classes and labor care in low-risk patients may help decrease CD. There is insufficient evidence that prenatal education and support programs, computer-based patient decision aids, decision-aid booklets, and intensive group therapy are effective. There is insufﬁcient evidence (lack of any trial) to assess the benefits and risks of a policy of CDMR (only indication: woman's desire; also called elective CD, a term which should be avoided) compared with trial of labor in women with term singleton gestations in cephalic presentation. The most common reason for a request for CDMR is fear of labor pain. CDMR should not be motivated by the unavailability of effective pain medication. There is also no randomized controlled trial (RCT) to evaluate the obstetrician's recommendation for CD without indication. As there are no such trials, there is insufficient evidence to assess the long-term maternal and neonatal morbidity and mortality of planned CD versus trial of labor. There is insufficient evidence on the impact of counseling during pregnancy with the aim of reducing the incidence of CDMR, but counseling reduces anxiety and concerns related to pregnancy and birth [5]. The incidence of women (without prior CD) preferring CDMR in a systematic review of studies is about 10%, and even less in low-income countries [6]. A practitioner is not obligated to perform a CDMR, but should appropriately refer the woman as necessary.

A CDMR or a planned repeat CD without other indication should not be performed before 39 weeks.

OPTIMAL CD RATE

The optimal CD rate is unknown. Per the World Health Organization (WHO), there were no reductions in maternal and neonatal mortality with CD rates higher than 10% [7]. However, it is important to note that this is based on an ecological analysis at the population level and does not provide information that can be used at an institutional or individual physician level [7]. Maternal and neonatal morbidity and mortality are the important outcomes, not CD rate per se. What is most important is that a woman/fetus who needs a cesarean is able to have one. A recent study on the relationship between CD rate and maternal and neonatal mortality in 194 WHO member states revealed that about 19% was the CD rate associated with lower maternal and neonatal mortality [8]. CD rates of less that 15%-20% have been associated instead with higher neonatal mortality rates [9].

In centers which have excessive CD rates, strategies to decrease the CD rate should focus on the three main indications for CD: arrest (or dystocia, CPD); NRFHT; and malpresentation. These strategies include using 6 cm for active labor, not performing CD for arrest before 6 cm, not performing a CD for prolonged latent phase, reserving consideration for CD for arrest in the first stage of labor only for women ≥6 cm dilation with rupture of membranes who failed to progress despite ≥4 hours of adequate uterine activity, or ≥6 hours of inadequate uterine activity and no cervical change, calling a failed induction in the latent phase one which requires ≥18–24 hours of oxytocin after membrane rupture, using secondary means (e.g., scalp stimulation) to assure normal fetal status in cases of NRFHT, routine use of external cephalic version for nonvertex presentations, use of operative vaginal delivery as appropriate, offering trial of labor after CD, and others [4,10].

There are limited data of nonclinical interventions aimed at decreasing the unnecessary CD rate [11]. Implementation of guidelines with mandatory second opinion can lead to a small reduction in CD rates, predominately in intrapartum CD [12]. Peer review [13], including precesarean consultation, mandatory secondary opinion, postcesarean surveillance, and audit [13] can lead to a reduction in repeat cesarean delivery rates. Guidelines disseminated with endorsement and support from local opinion leaders may increase the proportion of women with previous CD being offered a trial of labor in certain settings (see also Chapter 14). Nurse-led relaxation classes and birth preparation classes may reduce CD rates in low-risk pregnancies [11]. Primary midwifery antenatal [14] and labor [15] care in low-risk patients may help decrease CD. There is insufficient evidence that prenatal education and support programs, computer-based patient decision aids, decision-aid booklets, and intensive group therapy are effective. There

![Figure 13.1](https://example.com/f13.1.png)  
**Figure 13.1**  Primary cesarean delivery indications. (Adapted from American College of Obstetricians and Gynecologists, Society for Maternal-Fetal Medicine, *Obstet Gynecol*, 123(3), 693–711, 2014; Barber, E. L. et al., *Obstet Gynecol*, 118(1), 29–38, 2011.)
is insufficient evidence that training of public health nurses, insurance reform, and legislative changes are effective.

**TIMING OF DELIVERY**

In a study of 23,794 women, planned repeat CD at 37 or 38 weeks gestation has a significantly increased risk of adverse neonatal outcomes when compared with planned repeat CD at 39 or 40 weeks or expectant management [16]. A planned repeat CD should in general be performed at 39 0/7–39 6/7 weeks, unless there is a medical indication to perform it earlier (Table 56.9, Chapter 56, Antenatal Testing chapter in *Maternal-Fetal Evidence Based Guidelines*).

**PREOPERATIVE CONSIDERATIONS**

See Tables 13.1 and 13.2 for summary of recommendations on how to perform a CD [17].

Consent should always be obtained after counseling. Counseling should include a discussion of indications, benefits, risks including possible complications, and alternatives.

A checklist, including possibly pre-, intra-, and postoperative steps aimed at preventing complications, has been associated with benefits in general surgery [18] and should probably be implemented at CD [19] (Table 13.2).

**Prophylactic Antibiotics**

Who to Give Them to

Prophylactic antibiotics should be administered before every CD [20,21]. They are associated with decreased incidence of endometritis by 62%, wound infection by 60%, fever by 55%, and serious maternal infectious complications by 69%. Urinary tract infections (UTI) are also markedly decreased [21]. These results are similar for elective, scheduled CDs.

Which Antibiotics to Use and How

Comparing which antibiotic to give, the efficacy of a first-generation cephalosporin such as cefazolin (Ancef) appears equivalent to that of ampicillin [22–25]; however, the former is preferred [26]. These are the recommended agents to use unless there are drug allergies to them. A seemingly equally efficacious alternative is penicillin [27], though sample size may have been too small to show a difference. Later-generation (e.g., second or third), or more expensive broad-spectrum agents, do not improve efficacy further [27]. A multiple-dose regimen for prophylaxis appears to offer no added benefit over a single-dose regimen [21,24,28]. Systemic administration after cord clamping versus lavage routes of antibiotic administration seems to have similar efficacy to each other [21,28]. If ampicillin or a first-generation cephalosporin has already been given in labor (e.g., for chorioamnionitis), there may be no need for additional prophylactic antibiotics at CD [26]. If preoperative antibiotics were not given, prophylaxis should include an extended-spectrum regimen, involving azithromycin or metronidazole in the setting of chorioamnionitis [29]. Obese women may benefit from higher doses, e.g., Ancef 2–4 g) [26,30–34] (see Chapter 3 in *Maternal-Fetal Evidence Based Guidelines*).

When to Give the Antibiotics

Compared with after cord clamp, administration of antibiotics within 1 hour (optimally about 30 minutes) before skin incision is associated with a lower incidence of endometritis and wound infection [20,35–39]. In a meta-analysis of 10 trials, antibiotics administered preoperatively as compared with after neonatal cord-clamp are associated with a 46% decreased incidence of endometritis and 41% decreased incidence of wound infection [20]. Pharmacokinetic studies demonstrate that adequate cefazolin tissue concentration is attained 30 minutes after administration [20,40,41].

**Summary**

This evidence suggests the use of a single dose of IV cefazolin or ampicillin given within 1 hour (about 30 minutes) prior to skin incision.

<table>
<thead>
<tr>
<th>Table 13.1</th>
<th>Standard Recommendation Language and Quality of Evidence</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Recommendation:</strong></td>
<td></td>
</tr>
<tr>
<td>A. Strongly recommend that clinicians provide (the service) to eligible patients. There is good evidence that (the service) improves important health outcomes and concludes that benefits substantially outweigh harms.</td>
<td></td>
</tr>
<tr>
<td>B. Recommend that clinicians provide (this service) to eligible patients. There is at least fair evidence that (the service) improves important health outcomes and concludes that benefits outweigh harms.</td>
<td></td>
</tr>
<tr>
<td>C. No recommendation for or against routine provision of (the service). There is at least fair evidence that (the service) can improve health outcomes but concludes that the balance of benefits and harms is too close to justify a general recommendation.</td>
<td></td>
</tr>
<tr>
<td>D. Recommend against routinely providing (the service) to asymptomatic patients. There is at least fair evidence that (the service) is ineffective or that harms outweigh benefits.</td>
<td></td>
</tr>
<tr>
<td>E. Evidence is insufficient to recommend for or against routinely providing (the service). Evidence that the (service) is effective is lacking, of poor quality, or conflicting and the balance of benefits and harms cannot be determined.</td>
<td></td>
</tr>
<tr>
<td><strong>Quality of evidence:</strong></td>
<td></td>
</tr>
<tr>
<td>Good: Evidence includes consistent results from well-designed, well-conducted studies in representative populations that directly assess effects on health outcomes.</td>
<td></td>
</tr>
<tr>
<td>Fair: Evidence is sufficient to determine effects on health outcomes, but the strength of the evidence is limited by the number, quality, or consistency of the individual studies, generalizability to routine practice, or indirect nature of the evidence on health outcomes.</td>
<td></td>
</tr>
<tr>
<td>Poor: Evidence is insufficient to assess the effects on health outcomes because of limited number or power of studies, important flaws in their design or conduct, gaps in the chain of evidence, or lack of information on important health outcomes.</td>
<td></td>
</tr>
</tbody>
</table>

Source: Adapted from the Method Outlined by the U.S. Preventive Services Task Force (USPSTF) U.S. Preventive Services Task Force, Agency for Health Care Research and Quality, Available at www.ahcpr.gov, Updated 2015.
### Table 13.2 Evidence-Based Recommendations for CD

<table>
<thead>
<tr>
<th>CD Technical Aspect</th>
<th>Recommendation&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Quality&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Prophylactic antibiotics</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes or no</td>
<td>A</td>
<td>Good</td>
<td>Yes for all CD</td>
</tr>
<tr>
<td><strong>Antibiotic type</strong></td>
<td>A</td>
<td>Good</td>
<td>First generation cephalosporin or ampicillin preoperatively</td>
</tr>
<tr>
<td><strong>Route of administration</strong></td>
<td>A</td>
<td>Good</td>
<td>IV</td>
</tr>
<tr>
<td><strong>Single dose</strong></td>
<td>B</td>
<td>Good</td>
<td>b</td>
</tr>
<tr>
<td><strong>Timing</strong></td>
<td>A</td>
<td>Good</td>
<td>Administer 30–60 minutes before skin incision</td>
</tr>
<tr>
<td><strong>Prophylaxis for VTE</strong></td>
<td>C</td>
<td>Poor</td>
<td>Mechanical prophylaxis with graduated compression stockings or a pneumatic compression device during and after CD</td>
</tr>
<tr>
<td><strong>Lateral tilt</strong></td>
<td>B</td>
<td>Fair</td>
<td>15° to left</td>
</tr>
<tr>
<td><strong>No indwelling bladder catheterization</strong></td>
<td>C</td>
<td>Poor</td>
<td>No complications with its avoidance</td>
</tr>
<tr>
<td><strong>Hair removal</strong></td>
<td>B</td>
<td>Fair</td>
<td>Hair does not need to be removed. If the decision is made to remove hair, use hair clippers (not razors) on the morning of the surgery</td>
</tr>
<tr>
<td><strong>Skin cleansing</strong></td>
<td>B</td>
<td>Fair</td>
<td>Use chlorhexidine-alcohol&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td><strong>Vaginal irrigation</strong></td>
<td>C</td>
<td>Fair</td>
<td>Povidone-iodine&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td><strong>Adhesive drapes</strong></td>
<td>B</td>
<td>Fair</td>
<td>Avoid</td>
</tr>
<tr>
<td><strong>Skin incision</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Type</strong></td>
<td>C</td>
<td>Fair</td>
<td>Transverse, Pfannenstiel or Joel–Cohen</td>
</tr>
<tr>
<td><strong>Length</strong></td>
<td>I</td>
<td>Poor</td>
<td>15 cm</td>
</tr>
<tr>
<td><strong>Changing to a second knife</strong></td>
<td>C</td>
<td>Fair</td>
<td>No need to change knife after skin incision&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td><strong>Subcutaneous dissection</strong></td>
<td>I</td>
<td>Poor</td>
<td>we prefer bluntly&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td><strong>Fascial incision</strong></td>
<td>I</td>
<td>Poor</td>
<td>b</td>
</tr>
<tr>
<td><strong>No rectus muscle cutting</strong></td>
<td>B</td>
<td>Fair</td>
<td>Avoid</td>
</tr>
<tr>
<td><strong>Dissection of fascia off rectus</strong></td>
<td>I</td>
<td>Poor</td>
<td>b</td>
</tr>
<tr>
<td><strong>Opening of peritoneum</strong></td>
<td>I</td>
<td>Poor</td>
<td>b</td>
</tr>
<tr>
<td><strong>Bladder flap</strong></td>
<td>C</td>
<td>Fair</td>
<td>Routine use not necessary for term deliveries</td>
</tr>
<tr>
<td><strong>Uterine incision</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Type</strong></td>
<td>B</td>
<td>Fair</td>
<td>Low transverse uterine incision preferred</td>
</tr>
<tr>
<td><strong>Stapling device</strong></td>
<td>B</td>
<td>Fair</td>
<td>Not recommended</td>
</tr>
<tr>
<td><strong>Expansion of uterine incision</strong></td>
<td>A</td>
<td>Good</td>
<td>Use blunt uterine incision expansion, with cephalad-caudad traction</td>
</tr>
<tr>
<td><strong>Fetal delivery</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Delayed cord clamping</strong></td>
<td>B</td>
<td>Good</td>
<td>Delay cord clamping for 30–120 seconds for all infants &lt;37 weeks&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td><strong>Prevention of uterine atony</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Oxytocin</strong></td>
<td>I</td>
<td>Poor</td>
<td>IV or IM&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td><strong>Oxytocin infusion rate</strong></td>
<td>C</td>
<td>Poor</td>
<td>Optimal rate unclear&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td><strong>Carbetocin</strong></td>
<td>B</td>
<td>Fair</td>
<td>Carbetocin superior to oxytocin, should be used if available&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td><strong>Misoprostol</strong></td>
<td>C</td>
<td>Fair</td>
<td>b</td>
</tr>
<tr>
<td><strong>Tranexamic acid</strong></td>
<td>A</td>
<td>Good</td>
<td>Use to prevent postpartum hemorrhage&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td><strong>Uterine massage</strong></td>
<td>B</td>
<td>Fair</td>
<td>Perform with cord traction</td>
</tr>
<tr>
<td><strong>Placental removal</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Spontaneous</strong></td>
<td>A</td>
<td>Good</td>
<td>Avoid manual removal</td>
</tr>
<tr>
<td><strong>Glove change</strong></td>
<td>B</td>
<td>Fair</td>
<td>Not recommended</td>
</tr>
<tr>
<td><strong>Uterine exteriorization</strong></td>
<td>C</td>
<td>Fair</td>
<td>Suggested to facilitate better visualization</td>
</tr>
<tr>
<td><strong>Cleaning of uterus</strong></td>
<td>I</td>
<td>Poor</td>
<td>b</td>
</tr>
<tr>
<td><strong>Cervical dilation</strong></td>
<td>I</td>
<td>Poor</td>
<td>Avoid&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td><strong>Closure of uterine incision</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Two vs. one layers</strong></td>
<td>B</td>
<td>Fair</td>
<td>Double layer generally though single layer can be considered if bilateral tubal ligation performed concurrently&lt;sup&gt;b&lt;/sup&gt;; continuous stitches</td>
</tr>
<tr>
<td><strong>Sharp vs. blunt needles</strong></td>
<td>I</td>
<td>Poor</td>
<td>b</td>
</tr>
<tr>
<td><strong>Intra-abdominal irrigation</strong></td>
<td>B</td>
<td>Fair</td>
<td>Not recommended</td>
</tr>
<tr>
<td><strong>Peritoneal closure</strong></td>
<td>A</td>
<td>Good</td>
<td>Not recommended (for both parietal and visceral)</td>
</tr>
<tr>
<td><strong>Reapproximation of rectus muscles</strong></td>
<td>I</td>
<td>Poor</td>
<td>b</td>
</tr>
<tr>
<td><strong>Fascial closure</strong></td>
<td>I</td>
<td>Poor</td>
<td>b</td>
</tr>
<tr>
<td><strong>Subcutaneous tissue closure</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≥2 cm thickness</td>
<td>A</td>
<td>Good</td>
<td>Close subcutaneous space if ≥2 cm</td>
</tr>
<tr>
<td><strong>Subcutaneous tissue drain</strong></td>
<td>A</td>
<td>Good</td>
<td>Routine use not recommended</td>
</tr>
<tr>
<td><strong>Closure of skin</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Staples vs. subcuticular suture</strong></td>
<td>A</td>
<td>Good</td>
<td>Close transverse skin incision with suture&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

<sup>a</sup> Level of evidence was based on the U.S. Preventive Services Task Force recommendations (Table 13.1).

<sup>b</sup> See text for more details.

<sup>c</sup> Based on trials of women with vaginal delivery, not CD.

Abbreviations: CD, cesarean delivery; IV, intravenous; IM, intramuscular; VTE, venous thromboembolism.
Prophylactic Agents to Prevent VTE

There is insufficient evidence to give strong recommendations for thromboprophylaxis during pregnancy and the early postnatal period [42]. The three most commonly used interventions are pneumatic compression devices, compression stockings, and anticoagulants such as unfractionated (UH) or low-molecular-weight heparin (LMWH).

Pneumatic compression devices have been recommended based on retrospective data [43–46]. They appear to be safe and effective.

While there are two RCTs on UH versus placebo, two RCTs of LMWH versus placebo, and four RCTs on LMWH versus UH given antenatally, the RCTs are, in general, small and the data are insufficient to make any recommendation [44,45]. It is not possible to assess the effects of any of these interventions on most outcomes, and especially on rare outcomes such as VTE, death, and osteoporosis, because of small sample sizes and the small number of RCTs making the same comparisons. There was some evidence of side effects associated with thromboprophylaxis. In summary, given the higher risk of VTE at CD compared with vaginal delivery, all women undergoing CD should receive at least mechanical VTE prophylaxis with either pneumatic compression devices or compression stockings [45]. These should be applied preoperatively and continued until full ambulation.

In women undergoing CD with body mass index (BMI) > 50 kg/m², previous VTE, or two or more additional risk factors for VTE (such as smoking, multiple gestation, BMI ≥ 30 kg/m², prolonged immobilization, and infection), pharmacological VTE prophylaxis, with either enoxaparin 40 mg daily or UH 5000 every 12 hours in addition to mechanical prophylaxis, should be considered. This pharmacological prophylaxis can start at 6–12 hours postoperatively, after concerns for hemorrhage have decreased, and can continue until full ambulation [45].

Fetal Heart Monitoring

1. If external monitoring has been employed, it should be continued up until the abdominal prep has begun. This includes the time when regional anesthesia is administered. If continuous fetal monitoring is not possible, reapply the external monitor for 2–3 minutes if feasible after completion of the regional anesthesia to determine the postanesthesia fetal status.

2. If internal monitoring has been employed, the scalp electrode can be kept on until delivery of the fetal head, at which point the lead can be cut and the fetus delivered or the fetus delivered with the electrode attached. The operating room team will be responsible to document on the count sheet the location of the scalp electrode after delivery.

3. If the CD is done for nonreassuring fetal status, all attempts should be made to perform continuous fetal monitoring until the delivery occurs. This may not apply when the CD is done in an emergent manner.

There is no trial regarding optimal time of “decision to incision” for CD. Thirty minutes for CD for NRSFHT and 60 minutes for CD for dystocia and most other indications have been proposed but are not based on trials [47,48]. Additionally, studies have shown that while many cesareans are not performed within this time frame, neonatal outcomes are not adversely affected by longer intervals from decision to delivery [49,50], unless significant neonatal compromise is suspected (such as with fetal bradycardia).

Steroids for Fetal Maturity

If delivery by cesarean is necessary before 37 weeks, beta-methasone 12 mg intramuscular (IM) × 2 doses, 24 hours apart (or dexamethasone 6 mg IM × 4 doses, 12 hours apart) should be given for fetal maturity [51,52]. There is new evidence that steroids are associated with neonatal benefits also ≥34 weeks, as three RCTs have been done on steroids administered between 34 and 36 6/7 weeks [51,53,54] in women at risk of preterm birth (PTB), and two RCTs in women at ≥37 weeks [55,56]. Women who received antenatal corticosteroids ≥34 weeks had a significantly lower incidence of respiratory distress syndrome (RDS) (relative risk [RR] 0.76, 95% confidence interval [CI] 0.62–0.93), mild RDS (RR 0.40, 95% CI 0.23–0.69), moderate RDS (RR 0.39, 95% CI 0.18–0.89), severe RDS (RR 0.66, 95% CI 0.53–0.82), transient tachypnea of the newborn (RR 0.62, 95% CI 0.50–0.77), use of surfactant (RR 0.61, 95% CI 0.38–0.99), mechanical ventilation (RR 0.62, 95% CI 0.41–0.94), significantly less time on oxygen (mean difference [MD] –2.06 hours, 95% CI –2.17 to –1.95), lower maximum inspired oxygen concentration (MD –0.66%, 95% CI –0.69 to –0.63), shorter length of stay (LOS) in neonatal intensive care unit (NICU) (MD –7.64 days, 95% CI –7.65 to –7.64), higher Apgar scores at 1 and at 5 minutes (MD 0.06, 95% CI 0.05–0.07) compared with those who did not [57]. Steroids should be considered to decrease respiratory and other neonatal morbidities in women at high risk of preterm birth between 34 0/7 to 36 6/7 weeks [51].

Betamethasone 12 mg × 2 doses 24 hours apart at 37 weeks or beyond before planned CD has been also shown to reduce the incidence of RDS, to 0.2% from 1.1% in one RCT [58]. However, as this trial was not blinded, or placebo controlled, these data are insufficient for a definite recommendation. Steroids for fetal maturity should in general not be given at ≥39 weeks since the incidence of RDS is small.

Music

Playing music preoperatively significantly increases positive emotions and decreases negative emotions [59]. Playing music during planned CD under regional anesthesia may improve pulse rate and birth satisfaction score. However, the magnitude of these benefits is small and the methodological quality of the one included trial is questionable. Therefore, the clinical significance of music is unclear [60,61].

PREPARATIONS ON THE OPERATIVE TABLE

Maternal Position

There is insufficient evidence to support or clearly disprove the value of the use of tilting or flexing the table, the use of wedges and cushions, or the use of mechanical displacers at CD [62]. Most of the results below are from small single RCTs. The incidence of air embolism is not affected by head up versus horizontal position. Lateral tilt involves tilting the woman toward her left side 10°–15° to avoid vena caval compression by the gravid uterus. There are no changes in hypotensive episodes when comparing left lateral tilt (RR 0.11, 95% CI 0.01–1.94), right lateral tilt (RR 1.25, 95% CI 0.39–3.99), and head down tilt (MD –3.00; 95% CI –8.38 to 2.38) with horizontal positions or when comparing full lateral tilt with 15° tilt (RR 1.20, 95% CI 0.80–1.79) [62]. Hypotensive episodes are
decreased with manual displacers (RR 0.11, 95% CI 0.03–0.45), a right lumbar wedge compared with a right pelvic wedge (RR 1.64, 95% CI 1.07–2.53), and increased right lateral tilt (RR 3.30, 95% CI 1.20–9.08) versus left lateral tilt [62]. Position does not affect systolic blood pressure when comparing left lateral tilt or head down tilt to horizontal positions, or full lateral tilt to 15° tilt. Manual displacers resulted in a decreased fall in mean systolic blood pressure compared with left lateral tilt. Position does not affect diastolic blood pressures when comparing left lateral tilt versus horizontal positions. The mean diastolic pressure is a bit lower in head down tilt when compared with horizontal positions. There are no statistically significant changes in maternal pulse rate, 5-minute Apgars, maternal blood pH, or cord blood pH when comparing different positions [62].

**Indwelling Bladder Catheterization**

Compared with use of indwelling urinary catheters inserted pre-CD and removed ≥12 hours after CD, a systematic review of three trials showed that nonuse is associated with a lower incidence of urinary tract infection (UTI) (RR 0.08, 95% CI 0.01–0.64), lower rate of discomfort at first voiding (RR 0.06, 95% CI 0.03–0.12), less time to first voiding (by 16 minutes), and less time until ambulation (by about 6 minutes) [63]. No differences in intraoperative difficulties, complications (including urinary retention), or operative time were seen [63]. Given that these studies were not powered to assess differences in bladder or ureteral injury, were both performed in developing countries, that the quality of the RCTs were poor, and the use of prophylactic antibiotics and other important confounders were not reported, there is still insufficient evidence to justify the routine use of bladder catheterization, but its avoidance does not seem to be associated with complications.

**Hair Removal**

Based on a meta-analysis of 1343 patients, shaving was associated with twice the number of surgical site infections as compared with electric clipper the morning of the surgery is preferred [64,65] (see Chapter 7).

**Skin Cleansing**

Skin is impossible to sterilize. In nonpregnant adults, there are no differences in wound infection with different types and times of scrubs. Skin cleansing techniques for CD have been insufficiently studied for an evidence-based recommendation. Compared with standard preparation of 7.5% p-i scrub and then p-i 10% solution, the addition of preceding parachlorometaxylenol scrub for 5 minutes, in the women who had received prophylactic antibiotics for CD, is not associated with differences in incidences of endometritis or wound infection in a small, likely underpowered trial [66]. Chlorhexidine-alcohol scrub results in less wound infections than p-i scrub (9.5% vs. 16.1%, RR 0.59; 95% CI 0.34–0.95). No other outcomes were significantly different between the vaginal cleansing and control groups. No adverse effects were reported with the p-i vaginal cleansing [71]. Most, but not all, of these studies used p-i skin cleansing and prophylactic antibiotics so it is difficult to determine if vaginal preparation is necessary in women who receive the current recommendations for preoperative antibiotics; however, as a simple, generally inexpensive intervention, providers should consider implementing preoperative vaginal cleansing with p-i.

**Anesthesia**

For anesthesia, see Chapter 11. Metoclopramide 10 mg IV before spinal and ondansetron 4 mg IV after neonatal cord clamping are associated with less intraoperative and early postoperative nausea and vomiting as compared with placebo [78].

**Surgical Technique**

**Skin Incision**

Skin incision techniques for CD have been evaluated separately from other aspects of CD in limited studies [79,80]. In general, a transverse skin incision is recommended, since this is associated with less postoperative pain and improved cosmesis compared with a vertical incision. The Pfannenstiel (slightly curved, 2–3 cm or two fingerbreadths above the symphysis pubis, with the midportion of the incision lying within the shaved area of the pubic hair) and Joel–Cohen (straight, 3 cm below the line joining the anterior superior iliac spines, and therefore slightly more cephalad than the Pfannenstiel) are the preferred transverse incisions. Most RCTs not only evaluate type of skin incision, but also other technical aspects of CD,
making it often impossible to evaluate the effect of only the type of skin incision [81]. The better designed, larger trial revealed no differences in total operative time (32 vs. 33 minutes), intra- and postoperative complications, and neonatal outcomes, with the extraction time 50 seconds shorter for the Joel–Cohen group [79,82]. Considering the absence of clinical benefits to the mother and fetus, there is no clear indication for preferring either a Pfannenstiel or a Joel–Cohen incision for CD. In contrast, a smaller, less well-designed trial [80] shows significantly shorter operating times, reduced blood loss and postoperative discomfort associated with the Joel–Cohen compared with the Pfannenstiel incision [82].

There are probably no absolute indications for performing a vertical skin incision. Compared with transverse skin incision, vertical skin incision is associated with slightly shortened incision-to-delivery intervals of about 1 minute for primary and about 2 minutes for repeat CD [83].

Skin incision length has not been studied in a trial. Abdominal surgical incision size should probably provide about 15 cm (size of a standard Allis clamp) of exposure to assure optimal outcome of both mother and term fetus [1,84].

Changing to a second scalpel after the first scalpel has been used for skin incision versus no such change has never been evaluated in a trial, or in any obstetrical literature. From general surgery data, one scalpel is probably adequate to use throughout the whole surgical procedure.

Subcutaneous Tissue Opening
There is limited data on whether the subcutaneous tissue should be opened with blunt or sharp technique. We use the scalpel as little as possible, opening layers bluntly from medial to lateral to avoid injury to tissue and the inferior epigastric vessels. Blunt dissection has been associated with shorter operating times. In one RCT, use of diathermy (Bovie) for CD abdominal wall opening from subcutaneous tissue until the peritoneum was associated with lower blood loss, lower skin-to-peritoneum incision time, and lower post-CD pain compared with the use of No. 22 disposable scalpel blade [85].

Fascial Incision
Fascial incision has not been studied separately in a trial. A transverse incision is usually performed with the scalpel, and then extended with scissors. Digital extension can alternatively be accomplished by separating the forefingers in a cephalad-caudad direction after inserting the fingers into a small, midline transverse fascial incision. In an RCT evaluating entry into the abdomen, blunt entry with rectus sheath incision extended manually and parietal peritoneum entered and extended bluntly (manually) was associated with less blood loss, shorter operative time, and less post-CD fever and pain, compared with sharp entry [86].

Rectus Muscle Cutting
Rectus muscle cutting with Maylard technique is not associated with any difference in operative morbidity, difficult deliveries, or postoperative complications compared with Pfannenstiel (no muscle cutting) technique [84,87,88], but abdominal muscle strength at 3 months tends to be better in the Pfannenstiel group [88]. Pain scores also did not differ between the groups, but this may have been due to a sample size of only 97 women [88]. Therefore, rectus muscle cutting is probably not necessary [82].

Dissection of Fascia off the Rectus Muscles
Nondissection of the fascia off the rectus muscles inferiorly may result in less pain and similar blood loss as compared with dissection of the rectus sheath inferiorly during cesarean [89]. There seems to be no necessity of this commonly used technical step of CD [1].

Extraperitoneal versus Transperitoneal CD
There is insufficient evidence to compare maternal and perinatal outcomes between extraperitoneal versus transperitoneal CD, as only one small RCT has been performed [90]. The current standard is to perform CD transperitoneally.

Opening of the Peritoneum
Opening of the peritoneum has not been studied separately in a trial. The peritoneum is usually carefully opened with blunt or sharp dissection, and blunt expansion, high above the bladder, avoiding injury to organs below. As compared with sharp entry, blunt entry and extension of the rectus sheath incision and parietal peritoneum was associated with less blood loss, shorter operative time, and less post-CD fever and pain [86].

Retractors
There is insufficient evidence for comparing different types of retractors in CD.

Bladder Flap
Four RCTs of 581 women have compared development of a bladder flap versus direct uterine incision above the bladder fold [91]. There were no differences in the rate of bladder injury, estimated blood loss or hospitalization. Though the skin incision to delivery was 1.27 minutes longer for bladder flap formation, overall operative time did not differ [91]. However, one study was unpublished, two were judged to be of poor methodological quality, populations were heterogeneous, emergency cesareans were excluded from analysis and the majority of fetuses were >32 weeks gestation [91]. Based on the trial that was of better quality, there is some evidence that omission of the bladder flap shortened incision to delivery time in primary CD by 1 minute, though there was no difference in total operating time [92]. These results may not be able to be extrapolated to preterm and emergency CD since they were typically not included in these RCTs. No long-term effects (e.g., adhesions, bladder function, and fertility) have been evaluated. As bladder injury at CD is an uncommon event (1–3/1000), a sample size over 40,000 women would be required to show a difference in this outcome [93]. Developing a bladder flap at CD may not be necessary at term.

The use of a bladder blade to protect the bladder has not been studied separately in a trial.

Uterine Incision
Uterine incision type has not been studied separately in a trial. The transverse incision of the lower uterine segment is usually recommended because there is less blood loss and it allows for TOLAC in subsequent pregnancies [1,94]. Some experts advocate the classical vertical or at least low vertical incision if the lower uterine segment is not large enough to allow a transverse incision, for example, for the very preterm (<28 weeks) uterus or fibroids, but this has been associated with increased blood loss compared with low transverse incision [95].

Uterine Stapling
Use of uterine stapling (autosuture) device for opening and closing of the uterus has been assessed in two trials of 300
women. There is no difference in febrile morbidity between the groups [96]. One trial showed a nonsignificant increase in the duration of the procedure (by about 3 minutes) [96,97]. There is not enough evidence to justify the routine use of stapling devices to extend the uterine incision of the lower segment, especially since there is a possibility that stapling could cause harm by prolonging the time to deliver the baby.

Expansion of Uterine Incision
Expansion of the uterine incision with fingers (blunt) is associated with significantly decreased blood loss (by about 55 mL) and need for transfusion (RR 0.24; 95% CI 0.09–0.62) [96]. As it is also quicker, and associated with less risk of inadvertently cutting the neonate or cord, blunt should be preferred to sharp expansion of the uterine incision [98].

Compared with transverse expansion, cephalad-caudal expansion of the low transverse uterine incision is associated with significantly lower incidence of blood loss >1500 mL (0.2% vs. 2%) [99], less unintended uterine extensions (3.7% vs. 7.4%) [99] and less blood loss overall [100]. A meta-analysis of these two RCTs showed that women who were randomized in the cephalad-caudal group had lower incidences of postpartum blood loss (MD –67.64 mL, 95% CI –102.85 to –32.43), hemoglobin drop (MD –0.26 g/dL, 95% CI –0.37 to –0.14) and hematocrit drop 24 hours after CD (MD –1.20 g/dL, 95% CI –1.87 to –0.53), unintended extension (4.8% vs. 8.9%; RR 0.51, 95% CI 0.20–1.37), blood loss >1500 mL (0.2% vs. 1.7%; RR 0.12, 95% CI 0.02–0.99) and need for additional stitches (20.3% vs. 29.2%; RR 0.60, 95% CI 0.44–0.82). Therefore, cephalad-caudal uterine incision expansion by fingers should be preferred to transverse expansion [101].

Instrumental Delivery
Instrumental delivery of the fetal head by either vacuum or forceps compared with manual means has been insufficiently evaluated for a firm recommendation in women with cephalic [102] or breech presentation undergoing CD. As instrumentation has been associated with maternal (especially for forceps) or fetal (especially for vacuum) harm in vaginal deliveries, the principle of “primum non nocere” (first do no harm) should be applied in this setting, therefore favoring manual delivery of the fetal head whenever possible until further data are available.

Tocolysis for assisting in delivery of the fetal head at CD has been insufficiently studied [103].

Delivery of the Impacted Fetal Head
Though a strong recommendation cannot be made based on the available evidence, in cases where the fetal vertex is wedged into the maternal pelvis, vaginal displacement of the presenting part upward has been associated with longer operating time, more extension of the uterine incision and postpartum endometritis as compared with reverse breech extraction (“pull” method) in a small RCT [104]. A meta-analysis of RCT and non-RCT data confirmed reductions in uterine incision extension, blood loss and operative time with reverse breech extraction [105].

Family-Oriented CD
One study showed that allowing the parents to directly visualize delivery of the baby’s body (after head is delivered), to cut umbilical cord and to perform early skin to skin contact, improves birth satisfaction without increased blood loss [106].

Skin-to-Skin
Early skin to skin contact is beneficial for both mothers and babies and can be performed at time of CD [106–108]. Professional supervision is warranted to ensure neonatal well-being.

Collection and Drainage of Cord Blood
At CD, drainage of fetal blood from the umbilical cord is associated with less incidence of fetomaternal transfusion (measured by Kleihauer–Betke test) compared with no drainage [109]. The clinical significance of this finding is unknown.

Delayed cord clamping (DCC) for 30–120 seconds (or milking) increases neonatal blood volume by approximately 30% and decreases morbidity including intraventricular hemorrhage in preterm infants [110,111]. In term infants, it is associated with higher hematocrit, but also higher bilirubin levels. Therefore DCC is indicated routinely in preterm infants [112].

Prevention of Uterine Atony and Postpartum Hemorrhage
Prevention of uterine atony and postpartum hemorrhage has not been comprehensively studied for CD, but has been studied extensively for the third stage of labor after vaginal delivery (see Chapter 9).

Oxytocin
In the setting of vaginal delivery, both IV and IM oxytocin effectively reduce postpartum hemorrhage and the need for therapeutic uterotonics by at least 40% compared with placebo or no routine prophylactic agent. Oxytocin is as effective and has fewer side effects than ergot alkaloids.

Regarding oxytocin infusion rates at CD, patients required fewer additional uterotonics (19% vs. 36%) when treated with 80 international units (IU) oxytocin/500 mL infused over 30 minutes as compared with those who received 10 IU/500 mL infused over 30 minutes [113]. One study showed lower rates of excessive blood loss (EBL) >1000 cc, need for uterotonic and blood transfusion in those that received 5 IU oxytocin bolus and 30 IU infusion as compared with those who received 5 IU bolus and placebo [114]. Other lower oxytocin doses have been studied with nonsignificant differences between treatment groups [115]; the optimal infusion rate for oxytocin at CD is still unclear. For CD, oxytocin 20 IU IV is as effective as ergometrine plus oxytocin, with less vomiting, in a small RCT [116].

Carbetocin
For CD, carbetocin as a single 100-g dose is associated with more effective prevention of uterine atony and lower need for additional uterotonics compared with oxytocin 8- or 16-hour infusion [117,118]. Carbetocin (where available) may be recommended over oxytocin for prevention of uterine atony.

Misoprostol
There is insufficient evidence to compare misoprostol to oxytocin for prevention of uterine atony and postpartum hemorrhage at CD, as the seven RCTs compared different regimens of misoprostol and oxytocin. In single RCTs, either sublingual misoprostol or rectal misoprostol were associated with lower post-CD blood loss compared with oxytocin [119,120]. Given that misoprostol and oxytocin appear equally efficacious based on this limited evidence and that side effects such as shivering and pyrexia [119] are more common with misoprostol, oxytocin for now remains preferred [121].
Misoprostol combined with oxytocin (e.g., 400 μg sublingual after cord clamping, or rectal) was associated with less post-CD blood loss, fall in hematocrit and need for additional uterotonic agents when compared with oxytocin alone. In women at high-risk for post-CD hemorrhage, the combination of both misoprostol and oxytocin should be considered.

Tranexamic acid
TA inhibits fibrinolysis that potentiates the clotting system and can be used to prevent bleeding. Its half-life is 2–10 hours and it typically works immediately after IV administration. Side effects include gastrointestinal upset, but additional rare complications have been reported. TA was administered pre-cesarean though the time frame varied among trials (three out of nine studies administered it 10 minutes prior to incision). Dose also varied but was typically 1 g of TA in 20 mL of 5% glucose given IV over 5–10 minutes [122]. There was significantly less postpartum hemorrhage (PPH) (odds ratio [OR] 0.43) and mean blood loss (72 mL) in those treated with TA compared with those that were not [122]. Four trials reported no cases of maternal death or severe morbidity (including thromboembolism, seizure, ICU admission) among 1511 women. The most recent meta-analysis showed that all women in the nine RCTs received standard oxytocin prophylaxis; in addition the TA group received TA 1 gram or 10 mg/kg IV 10–20 minutes before skin incision or spinal anesthesia. Women who received TA experienced less postpartum blood loss (MD -167.50 mL, 95% CI -225.79, -109.20) compared with controls. Women who were randomized to the TA group had a significantly lower incidence of postpartum hemorrhage, i.e., blood loss more than 500 mL (3.9% vs. 41.9%; RR 0.06, 95% CI 0.04–0.10) and of severe postpartum hemorrhage, i.e., blood loss more than 1000 mL (1.3% vs. 3.0%; RR 0.42, 95% CI 0.19–0.92), compared with controls. The number of women who needed additional uterotonic agents was significantly lower in the TA group compared with controls (3.9% vs. 6.6%; RR 0.59, 95% CI 0.38–0.92). Women who received TA had a significantly lower hemoglobin drop as compared with controls (1.1 g/dL vs. 1.8 g/dL; MD -0.61 g/dL, 95% CI -1.04, -0.18). The percentage of women who required blood transfusions at or immediately after CD was significantly lower in the TA group compared with controls (2.1% vs. 5.7%; RR 0.36, 95% CI 0.20–0.64). There was no difference in the incidence of thromboembolic events in the two groups [123]. Therefore, TA should be recommended for preventing PPH in all women undergoing CD [122].

Uterine Massage
Uterine massage, associated with cord traction, is associated with less blood loss compared with no such interventions [124]. Uterine massage has not been studied by itself in an RCT for CD.

Placental Removal
In a meta-analysis of 4694 women, manual removal of the placenta is associated with greater morbidity than spontaneous expulsion with gentle cord traction: increased endometritis (RR 1.64, 95% CI 1.42–1.90); greater blood loss (by 94 mL); increased postpartum hemorrhage (RR 1.81, 95% CI 1.44–2.28); and decreased hematocrit after delivery (by 1.6%) [124]. Blood loss may be increased in manual removal because dilated sinuses in the uterine wall are not closed yet. Bacterial contamination of the lower uterine segment and incision may contaminate the surgeon’s dominant hand, and therefore the upper segment in manual removal, or the glove itself may be contaminated. Therefore, uterine massage with gentle cord traction resulting in spontaneous expulsion should be utilized for delivery of the placenta given the significant decrease in blood loss and endometritis as compared with manual placental removal.

Change of Gloves
Changing the operator’s glove before manual removal of the placenta does not alter the incidence of endometritis [125].

Uterine Exteriorization
Meta-analyses have showed there are no significant differences in blood loss, intraoperative hypotension, nausea/vomiting or pain, blood transfusion, endometritis or wound infection, with uterine exteriorization (extra-abdominal uterine incision repair) versus repair in situ [126,127]. The Cochrane review showed that there was less febrile morbidity (RR 0.41; 95% CI 0.17–0.97) and 0.24 day longer hospital stay with extra-abdominal closure [126]. So the balance of the benefits and harms is too close to justify a strong recommendation, but many obstetricians subjectively prefer to exteriorize the uterus for easier uterine incision repair.

Uterine Cooling
Uterine cooling after uterine exteriorization and during uterine closure is associated with a decrease in blood loss and postpartum hemorrhage in one RCT [128].

Cleaning the Uterus
Cleaning any placental remnants or blood clots from the uterus with a sponge or other means is a technique frequently used after placental removal, but not studied in any trial.

Cervical Dilation
Routine cervical dilation at CD before uterine incision repair has been insufficiently studied, but it is not associated with an effect on infectious morbidity (UTI, wound infection, endometritis) or change in hemoglobin [42,129].

Closure of Uterine Incision
At least one-layer uterine closure is always done, as the uterus should not be left open. Closure of uterine incision involves several decisions. These include use of blunt versus sharp needles; type of suture; full- versus split-thickness repair; continuous versus interrupted sutures; locking versus nonlocking of sutures; and whether or not to imbricate the second layer if it is even closed.

Blunt needles for closure of the uterus, peritoneum, and rectus sheath are associated with similar outcomes compared with sharp needles in one RCT [130]. In another RCT, glove perforation was significantly less with use of blunt compared with sharp needles, especially for the assistant surgeon. However, physicians reported decreased satisfaction performing CD with blunt needles [131]. In summary, there is still limited evidence to recommend blunt versus sharp needles at CD.

In one RCT, placing the sutures with the left hand and pulling the suture in a caudal direction was associated with lower need for additional sutures, shorter operative times, and lower decrease in hemoglobin compared with placing and pulling the suture with the right hand [132].

There is insufficient evidence to compare different sutures at closure at CD (no RCTs). In the only RCT comparing different sutures for uterine incision repair, polygla
tin-910
was associated with generally similar outcomes compared with chronic catgut [86]. Compared with split-thickness repair (avoiding the endometrium), full-thickness uterine incision repair is associated with a lower incidence of incomplete healing (documented by split in uterine muscle seen on transvaginal ultrasound about 40 days after the CD) of the uterine incision after CD [96,133].

Continuous single-layer closure may save operating time and reduce blood loss compared with interrupted single-layer closure [134]. Locking of sutures in the first layer has been insufficiently studied, but associated with poorer healing and possibly thinner residual myometrium, as is single layer closure [135].

Compared with two (double) layers, one (single) layer of suture for low transverse uterine incision repair is associated with no differences in febrile morbidity (13,980 women; RR 0.98). Although there was a reduction in mean blood loss for single layer closure, there were no differences in blood transfusion and heterogeneity was high for included studies [86,96]. Unfortunately, the women followed-up are too few to detect a significant difference in rare but extremely important long-term outcomes such as rates of rupture in the next pregnancy [42,96,134–137], with contradictory results of retrospective studies. Since there is, as of yet, no trial demonstrating benefit from two- versus one-layer uterine closure, it might be reasonable to omit the second layer if the woman is planning no more pregnancies (e.g., receives tubal ligation). For women planning future pregnancies, the uterus can be closed in two layers [138]. Vertical uterine incisions require a double or triple layer closure [139].

**Intra-Abdominal Irrigation**

Intra-abdominal irrigation with 500–1000 mL of normal saline before abdominal wall closure should not be routinely performed since it provides no significant differences in blood loss, intrapartum complications, hospital stay, return of gastrointestinal function, or incidence of infectious complications versus no irrigation [140]. In another RCT, 500–1000 mL of warm normal saline irrigation before the closure of the abdominal wall was associated with increased intraoperative nausea, but similar incidences of post-CD infectious morbidities [141].

**Adhesion Formation Prevention**

There is insufficient evidence to assess if any intervention is effective at adhesion prevention at CD. In one RCT, hyaluronate carboxymethylcellulose (Seprafilm®) adhesion barrier applied at CD did not reduce adhesion formation at the subsequent CD [142]. Evidence from non-CD abdominal surgery shows that oxidized regenerated cellulose (Interceed®) and hyaluronate carboxymethylcellulose (Seprafilm®) safely reduce clinically relevant consequences of adhesions [143].

**Intraoperative Interventions to Reduce Postoperative Pain**

One study of 370 patients who underwent primary CD showed that intraperitoneal instillation of 10 mL of 2% lidocaine significantly decreased persistent pain postoperatively from 21% to 11% parietal peritoneum was closed [144].

**Appendectomy**

Performing a planned appendectomy without indication at CD is not associated with inpatient morbidity in a small RCT [145]. However, no clear benefits were shown. The evidence is insufficient to make a recommendation regarding this nonindicated procedure.

**Intra-Abdominal Drain**

There is insufficient evidence to evaluate the effect of placing a drain in the abdominal cavity at CD. In one RCT, liberal use of a subrectus sheath drain was not associated with any effect compared with the restricted use of such intervention [146].

**Peritoneal Nonclosure**

Observational studies have shown that the peritoneum regenerates in 5–6 days. Compared with closure, peritoneal nonclosure is associated with a reduction in operating time whether both or either visceral or parietal peritoneal layers were not sutured. For nonclosure of both layers, the operating time was reduced by about 6 minutes [147]. While nonclosure of visceral peritoneum versus closure of both peritoneal surfaces showed an increase risk of adhesion formation, one of the two included studies had a high risk of bias. Peritoneal nonclosure is also associated with significantly less postoperative pain and shorter operative time as compared with closure of both layers. Nonclosure of the visceral peritoneum when the parietal peritoneum is closed is associated with decreased urinary symptoms of urgency, frequency and stress incontinence [147].

Long-term follow-up in one trial showed no significant differences. Long-term follow-up [148] after 7 years showed no differences in pain, fertility, urinary symptoms, and adhesions. Long-term studies following CD are limited; there is therefore no definite evidence for nonclosure until long-term data become available [147]. A review of general surgery and gynecological data concluded that "we encourage clinicians not to close both parietal and visceral peritoneum." [149] The hypothetical benefits of closing these layers for anatomic barrier, reduction of wound dehiscence, and minimization of adhesion have not been proven, and in fact have been invalidated by trials. There is no evidence to justify the time taken and the cost of peritoneal closure.

**Reapproximation of Rectus Muscles**

Reapproximation of rectus muscles has not been studied in any trial. Most clinicians agree that they do go back to their original anatomic place, and suturing them together can cause unnecessary pain when the woman starts to move postoperatively.

**Fascial Closure**

Techniques of fascial closure have not been studied in any trial of CD. Most experts suggest continuous nonlocking closure with delayed absorbable suture at about 1cm intervals. Recent non-CD evidence instead has shown that small fascial tissue bites of 5 mm every 5 mm are associated with prevention of incisional hernia in midline incisions and is not associated with a higher rate of adverse events, compared with large fascial bites of 1 cm every 1 cm [150].

**Subcutaneous Tissue**

**Irrigation**

Irrigation of the subcutaneous tissue to minimize wound infections and other complications has not been studied versus no irrigation in a trial of CD. The type of irrigation, with saline or antibiotic solution, has also not been studied in a trial.
**Suture Closure**

**Subcutaneous tissue closure** versus nonclosure by suture should be analyzed by the thickness of the subcutaneous tissue, as results differ according to <2 cm versus ≥2 cm of subcutaneous tissue thickness [151,152]. Most studies used 3-0 Vicryl for suture closure.

Any subcutaneous thickness: **Suture closure of subcutaneous fat in women** with a thickness of >2 cm is overall associated with less wound disruption versus nonclosure, but inclusion of both women with <2 cm and ≥2 cm thickness (which can have differing outcomes), and inability to blind represent a possible source of confounding and bias [152].

**Less than 2 cm subcutaneous thickness:** Routine subcutaneous tissue closure in women with a depth <2 cm has been insufficiently studied. It is not associated with any effects on outcome, and therefore cannot be recommended [151].

**Greater than or equal to 2 cm subcutaneous thickness:** Suture closure of subcutaneous fat in women with ≥2 cm thickness is associated with a significant decrease in wound disruptions, defined as any wound complication that required intervention, and seromas, compared with nonclosure. The evidence supports routine subcutaneous suture closure in women with a subcutaneous tissue depth ≥2 cm [151]. We find that many obstetricians underestimate subcutaneous thickness, so consider measuring this space if close to 2 cm.

**Drainage**

Some RCTs have evaluated drainage of subcutaneous tissue, compared with no drainage, or compared with tissue closure.

In meta-analyses of all RCTs, there is no evidence of a difference in the risk of wound infection, other wound complications, febrile morbidity, or endometritis in women who had wound drains compared with those who did not [153,154]. Drainage of subcutaneous tissue (i.e., wound drainage) in women with any thickness and who did not receive prophylactic antibiotics with a 2-cm corrugated rubber drain, left to drain open, coming out of one end of incision, and removed the following day is associated with a trend toward increased wound infection [153]. Drainage is also not as effective as tissue closure for women with ≥2 cm of subcutaneous fat. Drainage was usually performed with a 7-mm Jackson–Pratt drain [153]. Therefore, routine subcutaneous tissue drainage in women undergoing CD cannot be recommended [149]. These trials do not answer the question of whether wound drainage is of benefit when hemostasis is not felt to be adequate.

**Closure of Skin**

Closure of skin at CD has been most commonly performed with either absorbable sutures or nonabsorbable metal staples. In a meta-analysis of 3112 women, compared with sutures, staple closure is associated with higher rates of wound complications (13.0% vs. 4.8%), separation (9.4% vs. 2.5%), and a shorter duration of surgery by 7 minutes [155–157]. This decrease in wound complications persists even when only examining obese patients as suture was still associated with less complications (6.7% vs. 12.8%) [156]. Though the incidences were small, there were no significant differences in hematoma, seroma, or readmission between groups. Additionally, there were no significant differences between groups with regards to pain perception, patient satisfaction, and incision cosmesis. Therefore, the low transverse cesarean skin incision should be closed with suture.

There is insufficient evidence to compare different sutures for CD skin closure. The suture most commonly used in the RCTs showing superiority of suture compared with staples was poliglecaprone [156]; though the largest of these studies used poliglecaprone or polyglaclin [157]. Neither of these sutures has been shown to be superior to the other. In one small RCT, polyglycolic acid suture was associated with a higher incidence of hypertrophic scarring compared with interrupted nylon suture [158]. In one RCT, barbed suture was associated with similar rates of wound dehiscence, infection and other adverse outcomes, compared with 3-0 polydioxanone suture [159].

If staples are used, they should probably be removed on or after day 7, as early (day 3) removal is associated with a non-significant trend for higher rate of wound dehiscence (15.2% vs. 11.5%) compared with delayed (days 7–10) removal [160]. In one RCT, absorbable staples were associated with similar outcomes compared with metallic staples, except for a longer closure time [161].

There is insufficient evidence to evaluate the effectiveness of a new wound closure device, Leukosan® Skinlink, for skin closure at CD. The one small RCT comparing it to Prolene suture closure showed similar cosmetic results [162].

**POSTOPERATIVE CARE**

**Gum Chewing**

Compared with no gum chewing, gum chewing typically three times/day for at least 30 minutes each time after CD is associated with earlier return of bowel sounds (by 4.4 hours), earlier passage of flatus (by about 7.9 hours) [163] and stool (9.1 hours) [163] and less ileus (OR 0.36) [164,165].

**Early Oral Fluids and Feeding**

A meta-analysis of 11 somewhat heterogeneous studies showed that compared with delayed (usually after 8 hours or upon passage of flatus) oral fluids or food, early oral fluids or food are associated with reduced time (by about 8.8 hours) to return of bowel sounds; reduced time to flatus (73 hours) and decreased time to bowel movement (6.3 hours) [166]. No significant differences were identified with respect to nausea, vomiting, abdominal distention, and mild ileus [166,167]. Typically feeding was initiated within 6–8 hours with water, clear liquids or solid foods (4 studies) [168]. In summary, there is insufficient evidence for a strong recommendation, but early oral fluids and even food within 6–8 hours after CD seem to be safe and possibly beneficial.

**Pain Relief after CD**

**Nonsteroidal anti-inflammatory drugs** (NSAIDs; e.g., ibuprofen) and/or narcotics (e.g., oxycodone) are commonly used in the United States for post-CD pain relief (see also Chapter 11). There is no RCT evaluating oral narcotic use or oral NSAIDs use. Local analgesia wound infiltration and abdominal nerve blocks as adjuncts to regional analgesia and general anesthesia seem to be of benefit in CD by reducing opioid consumption in small RCTs. NSAIDs (even as a wound infiltration) as an adjuvant may confer additional pain relief [169]. In women who had CD performed under regional analgesia, wound infiltration is associated with a decrease in morphine consumption at 24 hours compared with placebo. In women with regional analgesia and also a local anesthetic, NSAID cocktail wound infiltration is associated with less morphine use compared with local anesthetic control. Women who have regional analgesia with
abdominal nerves blocked have decreased opioid consumption. In women under general anesthesia, with CD wound infiltration and peritoneal spraying with local anesthetic, the need for opioid rescue is reduced [169].

DISCHARGE
A study of almost 3000 women who were randomized to be discharged at 24 hours versus 72 hours postcesarean with their newborn showed that those discharged at 24 hours were more likely to report mood swings and less success with breastfeeding. Additionally, although there was no difference in maternal readmission, there were increased neonatal admissions (typically due to jaundice) in those discharged at 24 hours [170]. After a planned CD, discharge on the first as compared with the second day is not associated with any difference in maternal or perinatal immediate or 6-week outcomes [171]. For women who are discharged early, a home health registered nurse (RN) visit is advised.

MANAGEMENT OF COMPLICATIONS
Disrupted (Open) Laparotomy Wound
Compared with healing by secondary intention, reclosure of the disrupted laparotomy wound is associated with success in >80% of women, faster healing times (16–23 vs. 61–72 days), and fewer office visits [172]. No serious morbidity or mortality is associated with either method. There is insufficient evidence to assess optimal timing (probably 4–6 days after disruption if noninfected) and technique (superficial vertical mattress or “en bloc” reclosure of entire wound thickness with absorbable sutures, or adhesive tape) of reclosure, as well as utility of antibiotics. Compared with reclosure using sutures, reclosure using permeable, adhesive tape (Cover-Roll; Biersdorf, Inc., Norwalk, CT) is associated with faster procedure, less pain scores, and similar healing times in a small RCT [173].

Postoperative Counseling
Interval until next pregnancy after a CD should be about 18–23 months, as shorter intervals have been associated with increased risk of uterine rupture [174,175] (see also Chapter 14).

SHORT- AND LONG-TERM OUTCOMES FOR THE BABY
Scheduled CD compared with vaginal delivery (but not compared with unscheduled CD) has been associated with a small absolute increased risk of childhood asthma requiring hospital admission, salbutamol inhaler prescription at age 5 years, and all-cause death by age 21 years (0.40% vs. 0.32%; difference, 0.08% [95% CI, 0.02%–1.00%]; adjusted HR, 1.41 [95% CI, 1.05–1.90]) [176].

SHORT- AND LONG-TERM OUTCOMES FOR THE MOTHER
The dominant maternal risk in subsequent pregnancies is placenta accreta and its associated complications (Chapter 28). Pregnancies following CD also have increased risk for other types of abnormal placentation, reduced fetal growth, preterm birth, and possibly stillbirth. Chronic maternal morbidities associated with CD include pelvic pain and adhesions. Adverse reproductive effects may include decreased fertility and increased risk of spontaneous abortion and ectopic pregnancy [177].

FOR FUTURE RESEARCH
In order to make comparisons of CD rates over time, it is important that everyone use a similar classification scheme. Robson proposed a scheme that is mutually exclusive and totally inclusive and takes into consideration category of pregnancy (singleton cephalic/breech/other lie, multiples), prior obstetric record (nulliparous, multiparous with or without uterine scar), course of labor (spontaneous, induced, CD before labor), and gestational age (based on completed weeks at time of delivery) [178].

REFERENCES


Cracianus L, Sajid MS, Ahmed AS. Chewing gum in preventing postoperative ileus in women undergoing caesarean section: A systematic review and meta-analysis of randomised controlled trials. BJOG. 2014;121(7):793–799. discussion 799. [Meta-analysis, 7 RCTs, n = 1462].


Trial of labor after cesarean delivery

Amen Ness and Amanda Yeaton-Massey

KEY POINTS
• A woman with a prior cesarean delivery (CD) has two options for mode of delivery in the subsequent pregnancy: a planned repeat cesarean delivery (PRCD) or a trial of labor after cesarean (TOLAC) to try to achieve a vaginal birth after cesarean (VBAC).
• There is insufficient evidence (no large randomized controlled trial) to compare the safety, complications, maternal, and fetal/neonatal morbidity and mortality between these two options.
• TOLAC is a “reasonable option” for most women with a single prior low transverse CD with no other contraindications to a vaginal birth.
• Absolute contraindications to TOLAC are as follows:
  • Medical or obstetrical complications that preclude vaginal delivery
  • Inability to perform emergency CD
  • Vertical (classical) uterine scar or fundal or perifundal complete (from endometrium to serosa) uterine scar from other surgery (e.g., myomectomy)
  • Prior uterine rupture
• Successful VBAC rates in the general population of women with previous low transverse uterine incisions vary from 40% to 80%.
• No screening tool is sensitive enough to be clinically useful in predicting an unsuccessful trial of labor. One available for clinical use is at www.bsc.gwu.edu/mfmu/vagbirth.html.
• Rates of maternal and perinatal complications are in general similar and low with both TOLAC and PRCD, except for the risk of uterine rupture, which confers both maternal and perinatal risks.
• Uterine rupture is the main complication associated with TOLAC. Most maternal and perinatal morbidity results from repeat CD after TOLAC.
• The overall risk of uterine rupture during a TOLAC at term is 0.7% after one prior low transverse CD, versus 0.26% after PRCD. The lowest risk (0.4%) of rupture is with spontaneous labor TOLAC. With prior VBAC, TOLAC has also a very low (about 0.5%) risk of rupture. The risk is increased with >1 prior CD, prior vertical scar, prior rupture, induction of labor with no prior vaginal deliveries (especially when using prostaglandin ripening agents), augmentation with higher doses of oxytocin, interval between deliveries <18 months, maternal age >30-year-old, fetal macrosomia, single-layer closure, fever at time of prior CD, etc. (Table 14.2).
• With term uterine rupture, the risks of fetal/neonatal morbidity/mortality are about 33% risk of pH < 7.00, 40% admission to neonatal intensive care unit (NICU), 6% risk of hypoxic–ischemic encephalopathy (HIE), and 1.8% risk of neonatal death (rupture-related risk of neonatal death: 1/10,000 TOLAC) in equipped academic centers.
In other centers, these risks are higher, including risk of neonatal death from rupture up to 10%–25%.
• Compared with PRCD, TOLAC is associated with slightly higher rates of adverse perinatal outcome: cord pH <7.00 (1.5/1000 TOLAC), HIE (5–8/10,000 TOLAC), and perinatal death (13/10,000 with TOLAC vs. 1/10,000 for PRCD) (Table 14.3). The overall risk of adverse perinatal outcome is 1/2000 with TOLAC, slightly higher than with PRCD.
• When compared with women without a previous CD (instead of with those having a planned primary CD), the perinatal mortality for TOLAC is higher than PRCD (10/10,000 vs. 0.4/10,000 births), but this rate is twice as high as that of a non-VBAC multipara in labor and the same as that of a nullipara in labor.
• Appropriate counseling including risks as described should be provided to the woman with a prior CD deciding on subsequent mode of delivery. The ultimate decision regarding TOLAC or PRCD is up to the patient.
• To minimize risks, an experienced obstetrician in addition to anesthesia, nursing, and operating room (OR) personnel in a facility equipped to perform emergency CD must be immediately available throughout the TOLAC.

HISTORICAL PERSPECTIVE
Each year almost 1.5 million childbearing U.S. women have CDs. Most of these are primary CDs but, the largest single indication for CD is prior CD, accounting for over half a million CDs each year [1]. In order to reduce this trend, there is a need to both prevent the first CD and increase the rate of TOLAC in appropriate candidates.

Until the late 1970s, “Once a cesarean always a cesarean” was the general rule among most obstetricians. This phrase did not derive from formal studies and was clearly not evidence based. A classical uterine incision was used until the 1920s when the low transverse incision was first introduced. The low transverse incision was associated with a tenfold decreased rate of uterine rupture in labor than the classical incision. On the basis of studies in the 1970s, when the VBAC rate was very low, the National Institutes of Health (NIH) in 1980 and then the American College of Obstetricians and Gynecologists (ACOG) in 1988 and again in 1994 suggested that a trial of labor (TOL) after a previous low transverse CD is a reasonable option [2]. ACOG encouraged all women with a single prior low transverse cesarean section to consider a TOLAC. In response to these recommendations, the VBAC rate in the United States increased from 3.5% in 1980 to 28.3% in 1996 (Figure 14.1).

As more VBACs were attempted, more ruptures were seen and litigation for complications of a TOL also increased. In 1999, ACOG addressed these risks and added the requirements of a “readily available” physician “throughout labor” and “availability of anesthesia and personnel for emergency delivery” immediately available throughout the TOLAC.
Since that time, after declining until about 1997, CDs have increased globally, 33% in the United States in 2013 and 26% in England in 2013–2014 [4,5]. At the same time, VBAC rates have decreased rapidly to just 9% in the United States in 2011 (Figure 14.1), although they have remained relatively high in the United Kingdom at 33% (range 6%–64%) [6]. The ACOG requirement for “immediately available” personnel for women undergoing a TOLAC, but not for other laboring women, eliminated the option of TOLAC/VBAC in many community hospitals. Since 1996, about one-third of hospitals and one-half of physicians have stopped offering TOLAC/VBAC in the United States [6].

DEFINITIONS

**TOLAC:** Trial of labor after cesarean.

**VBAC:** Vaginal birth after cesarean.

**PRCD:** Planned repeat CD (before labor).

**VBAC rate:** Number of vaginal births after previous CD per 100 live births to all women with a previous CD (Same denominator as the CD rate).

**TOLAC rate:** If the average success rate of a TOLAC is about 70%, then the TOLAC rate is the VBAC rate/0.7.

**Adjusted VBAC rate:** Number of women with prior CD and no contraindications to TOL who had a VBAC per 100 live births to all women with a previous CD.

**Successful VBAC rate:** Percentage of women with prior CD who attempted a TOLAC achieving VBAC.

**Successful adjusted VBAC rate:** Percentage of women with prior CD and no contraindications to TOLAC achieving a VBAC.

**Failed TOLAC (failed VBAC):** TOLAC that results in a repeat CD.

**Uterine dehiscence:** Disruption of the uterine muscle with intact serosa [6]. It can include asymptomatic opening if the uterine scar is from prior surgery, without protrusion of fetus/fetal organs outside the uterus.

**Uterine rupture:** Disruption or tear of the uterine muscle and visceral peritoneum, or separation of the uterine muscle with extension to the bladder or broad ligament [6]. It includes symptomatic gross rupture of the uterine scar from prior surgery, with or without protrusion of fetus/fetal parts outside the uterus.

GENERAL CONSIDERATIONS

A woman with a prior CD has two options for mode of delivery in a subsequent pregnancy: a PRCD or a TOLAC to try to achieve a VBAC. There are two randomized controlled trials examining outcomes for women who underwent PRCD versus TOLAC [7,8]. Only one of these studies reported clinical
outcomes and was quite small with a sample size of 22 [8]. The other, while larger, looked at maternal psychometric outcomes [7]. There is also a small prospective cohort study to compare these two options which also focuses on psychological outcomes, not maternal and fetal safety and complications [9]. Virtually all studies on VBAC, with a few exceptions [10], are retrospective and often use differing criteria for patient selection and differ in their ability to correctly ascertain (make sure all cases are included) and define uterine rupture. Most studies also include women at various gestational ages and may therefore not specifically apply to women at term considering their options of delivery after a prior CD. Studies with <1000 TOLAC cannot adequately assess maternal and fetal/neonatal morbidity and mortality, as these complications are rare, and meta-analyses [11–13] might compound errors from different retrospective studies. Many studies do not differentiate between asymptomatic uterine dehiscence and true acute symptomatic uterine rupture.

The main issues regarding TOLAC are the following: (1) Which women are candidates for TOLAC, and which should instead be recommended a PRCD? (2) Among women who attempt a TOLAC, what is the VBAC rate (successful TOLAC), and which factors that influence it? (3) What are the short- and long-term maternal and perinatal benefits and harms of attempting TOLAC versus PRCD, and what factors influence benefits and harms? Complications and safety are especially related to the risk of uterine rupture.

These issues are important in order to properly counsel women who are considering a TOLAC [6]. This information should be shared with the woman in a way best suited to her understanding. When a TOLAC and a PRCD are medically equivalent options, the woman’s preference should be honored if possible [1].

CANDIDATES FOR TOLAC

The choice of candidates for TOLAC should be based on an acceptable balance between the chance of achieving a VBAC and maternal and fetal risks. The ACOG and the 2010 NIH consensus statement acknowledged that TOLAC is a reasonable option for most women with a single prior low transverse CD with no other contraindications to a vaginal birth [2,6]. Criteria to be a candidate, and absolute and relative contraindications for TOLAC are shown in Table 14.1.

FACTORS FOR SUCCESSFUL TOLAC

Vaginal delivery rates for TOLAC in the general population of women with previous low transverse uterine incisions vary from 60% to 80% [10,13–15]. In tertiary care centers the rates may be higher, about 73%–76% [10,16]. The rate is highly dependent on demographic and obstetric factors (Table 14.2). Based on these factors, women with a probability for VBAC of at least 60%–70% have similar or less maternal and perinatal morbidity with a TOLAC than a PRCD [20,33]. Conversely, women with less than a 60% probability of VBAC have a greater likelihood of morbidity with TOLAC than if they had a PRCD. One study demonstrated that composite neonatal morbidity is similar between TOLAC and PRCD for the women with the greatest probability of achieving VBAC [2,20]. Factors that influence the likelihood of VBAC after TOLAC, and their effect of success rates, are shown in Table 14.2, and described below.

Table 14.1 Candidates, and Absolute and Relative Contraindications for TOLAC

<table>
<thead>
<tr>
<th>Contraindications</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Singleton gestation</td>
</tr>
<tr>
<td>• Clinically adequate pelvis</td>
</tr>
<tr>
<td>• One (or two) prior low transverse CD(s)</td>
</tr>
<tr>
<td>• No other uterine scars or previous rupture</td>
</tr>
</tbody>
</table>

Limited evidence

• A twin pregnancy who is a candidate for VBAC
• Unknown uterine scar (low clinical suspicion of prior classical uterine incision)

Absolute contraindications

• Medical or obstetrical complications that preclude vaginal delivery
• Inability to perform emergency CD
• Vertical (classical) uterine scar
• Fundal or perifundal complete (from endometrium to serosa) uterine scar from other surgery (e.g., myomectomy)
• Prior uterine rupture

Relative contraindications

• Multiple uterine scars (e.g., ≥ three prior CDs)
• Any other factor associated with a risk of rupture of >1% (see text)

Abbreviations: CDs, cesarean deliveries; TOLAC, trial of labor after CD; VBAC, vaginal birth after cesarean.

Maternal Demographics

Age

There is inconsistent evidence regarding the effect of age on successful VBAC after a TOLAC. Recent prospective studies did not show any relationship although overall there appears to be a small inverse association between maternal age and the likelihood of vaginal delivery [1].

Race/Ethnicity

Race and ethnicity are the strongest demographic predictors of VBAC. Hispanic and African-American women have lower rates of VBAC than non-Hispanic white women.

Obesity

Obese women attempting a VBAC have lower vaginal delivery rates. This appears to be true whether measured at first prenatal visit (per body mass index [BMI] unit) or at delivery (BMI > 30) (adjusted ORs [aORs] 0.94, 95% confidence interval [CI] 0.93–0.95 and 0.55, 95% CI 0.51–0.60, respectively) [17,34]. Obese women, weighing ≥300 lb, had rates of VBAC of only 15%, while women weighing 200–300 lb had rates of 56% [35]. Greater maternal height and BMI < 30 kg/m2 are associated with an increased likelihood of VBAC.

Others

Single marital status and less than 12 years of education also have been associated with lower rates of VBAC.

Obstetric History

Prior Vaginal Delivery

A prior vaginal delivery, either before or after a prior CD, is the strongest predictor of a VBAC after a TOLAC. A previous successful VBAC is the most predictive [28]. The VBAC rate is >85% for women with a prior VBAC, compared with 65% for women without a prior vaginal delivery [36,37]. The rate of VBAC increases with each prior VBAC [38]. Women with zero, one, two, three, and four or more prior VBACs have likelihoods of VBAC of 63%, 88%, 91%, 91%, and 91% (p < .001), respectively [38].
Prior Indication for Cesarean

Breech presentation or other nonrecurring indication (e.g., nonreassuring fetal monitoring) for the prior CD significantly increase the chances for a vaginal delivery (85%) compared with CD done for a recurring indication such as dystocia or failure to progress. These rates are similar to vaginal delivery rates in nulliparous women. Nevertheless, about 60%–70% of women undergoing a TOLAC for dystocia (failure to progress) deliver vaginally [1]. In addition, most studies show no reduction in the rates for a successful TOLAC after a prior CD done for failure to progress in the second stage of labor (75%–80%), with no increased risk of operative vaginal delivery [1,39–41].

Uterine Scar Type

Vaginal delivery rates appear to be similar for low transverse, low vertical, and for unknown incision types [12,34]. Most women with unknown scar types have had low transverse incisions.

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Table 14.2 Factors Associated with Success and Risk of Uterine Rupture Associated with TOLAC

<table>
<thead>
<tr>
<th>Factor</th>
<th>Effect on Success Rates</th>
<th>Effect on Risk of Uterine Rupture with TOLAC (% risk)</th>
<th>Comment (References)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maternal Demographics</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Older age</td>
<td>↓</td>
<td>↑</td>
<td></td>
</tr>
<tr>
<td>Obesity</td>
<td>↓</td>
<td>NA</td>
<td></td>
</tr>
<tr>
<td>Prior vaginal delivery</td>
<td>↑</td>
<td>↓ (0.2%)</td>
<td>Especially prior VBAC. Recommend TOLAC [5,17,18]</td>
</tr>
<tr>
<td>No prior uterine scars</td>
<td>NA</td>
<td>NA (&lt;0.01%)</td>
<td>For example, prior CD for malpresentation</td>
</tr>
<tr>
<td>Prior nonrecurring indication for CD</td>
<td>↑</td>
<td>NA</td>
<td></td>
</tr>
<tr>
<td>Labor before primary CD</td>
<td>↓</td>
<td>(see below)</td>
<td>10%–15% lower success rate per CD</td>
</tr>
<tr>
<td>More than one prior CDs</td>
<td>↓</td>
<td>↑</td>
<td></td>
</tr>
<tr>
<td>1 Prior LT CD (no prior VBAC)</td>
<td>(0.7%, range 0.5–1.0)b</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 Prior LT CD (prior VBAC)</td>
<td>↓ (&lt;0.5%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2 Prior LT CD (no prior VBAC)</td>
<td>↑ (1.8%, 1%–3.7%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2 Prior LT CD (prior VBAC)</td>
<td>↓ (0.5%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Prior vertical (classical) CD</td>
<td>↑ (4%–10%)</td>
<td>Avoid TOLAC [6]</td>
<td></td>
</tr>
<tr>
<td>Prior low-vertical CD</td>
<td>↑ (1%–2%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Prior &quot;unknown uterine scar&quot; CD</td>
<td>(0.5%–2%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1-Layer uterine closure</td>
<td>↑ (1%–2%)</td>
<td>[9,48,49]</td>
<td></td>
</tr>
<tr>
<td>Fever at prior CD</td>
<td>↑</td>
<td>If both intra- and postpartum fever</td>
<td></td>
</tr>
<tr>
<td>Prior preterm CD</td>
<td>↑ (1%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Prior uterine rupture</td>
<td>↑ (6% if lower segment rupture; 32% if upper segment rupture)</td>
<td>Avoid TOLAC [6]</td>
<td></td>
</tr>
<tr>
<td>Short interpregnancy interval</td>
<td>↑ (2%–5%)</td>
<td>&lt;18 months; avoid TOLAC [61,62]</td>
<td></td>
</tr>
<tr>
<td>Current Labor Factors</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PRCD</td>
<td>NA</td>
<td>(0%–0.15% in labor) (0.5 risk of dehiscence)</td>
<td>[5,72]</td>
</tr>
<tr>
<td>Favorable cervical status</td>
<td>↑↑</td>
<td>↓</td>
<td>Bishop score &gt;8, or CL &lt;15 mm</td>
</tr>
<tr>
<td>Postdates</td>
<td>↓</td>
<td>–</td>
<td>Uterine rupture rate increased if induced/ augmented</td>
</tr>
<tr>
<td>Fetal macrosomia</td>
<td>↓</td>
<td>↑</td>
<td>Uterine rupture rate increased by relative risk 2.3 [22]</td>
</tr>
<tr>
<td>(e.g., BW &gt;4000–4500 g)</td>
<td>↓</td>
<td></td>
<td>Avoid misoprostol; avoid PGE2 followed by oxytocin; avoid oxytocin &gt;20 mU/minute. See below [5,51–55]</td>
</tr>
<tr>
<td>Induction/augmentation of labor</td>
<td>↓ (mostly 1%–2%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Misoprostol induction</td>
<td>↑↑ (&gt;5%)</td>
<td></td>
<td>Avoid</td>
</tr>
<tr>
<td>PGE2 only induction</td>
<td>↑ (1%)</td>
<td>Cannot predict if PGE2 will be sufficient</td>
<td></td>
</tr>
<tr>
<td>PGE2 then oxytocin induction</td>
<td>↑ (1%–3%)</td>
<td></td>
<td>Probably avoid</td>
</tr>
<tr>
<td>Foley induction</td>
<td>–</td>
<td></td>
<td>Probably safe</td>
</tr>
<tr>
<td>Oxytocin only induction</td>
<td>↑ (1.1%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oxytocin augmentation</td>
<td>↑ (0.9%)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: BW, birth weight; CL, cervical length; LT, low transverse; NA, not applicable; PGE2, prostaglandin E2; PRCD, planned repeat CD.

*a Chance of achieving vaginal birth after CD.

*b Used as reference.

*c See Chapter 13.
Gestational Age

previous studies had reported that 75%–80% of women attempting a VBAC with a single prior cesarean will deliver vaginally versus about 60%–70% with more than one prior cesarean. Recent studies have shown that the greater the number of prior CDs, the lower the chances for a vaginal delivery (about 10%–15% lower per CD) [12]. But a large retrospective multicenter study found similar rates of vaginal delivery after TOL in women with two versus one prior CD (75%) [42], while a recent meta-analysis of six studies that compared outcomes between women having a TOL after one or two CD found similar vaginal delivery rates of 77% and 72%, respectively [43].

Interdelivery Interval

Interdelivery interval of <18 months may be associated with similar success rates of TOL after CD compared with longer intervals [12], but in one study, rates were lower with induction [44].

Current Labor Factors

Cervical Status

Labor factors associated with a higher VBAC rate include greater cervical dilation at admission or at rupture of membranes. The more favorable the cervix, the greater the odds for a vaginal delivery. Fetal head engagement and a lower station are considered important predictors of successful VBAC [1].

Gestational Age

Preterm patients with prior CD have a slightly higher VBAC success rate than term patients (82% vs. 74%) [45].

Gestational age greater than 40 weeks is associated with a decreased rate of VBAC. There is conflicting data regarding the chance of VBAC after TOL in postterm pregnancy. Successful VBAC rates of 65%–82% have been reported for women past 40 weeks, with the higher rates in women with prior vaginal deliveries [1]. In the MFMU cohort, women >41 weeks had a lower chance for vaginal delivery than women less than 41 weeks (aOR 0.61, 95% CI 0.55–0.68) [34]. If VBAC is still desired after 40 weeks, awaiting the onset of spontaneous labor may be a better option than induction before 40 weeks for women planning a VBAC given the lower success rates with induction in this population, particularly for women with an unfavorable cervix [34,46].

Induction/Augmentation of Labor

The overall estimated rate of vaginal birth with a TOLAC after any method of labor induction is 63% [1]. Women with a prior CD who are induced have lower rates of successful TOLAC (about 10%) than those who labor spontaneously [11,12,47]. Studies demonstrate that the rate of VBAC ranges from 54% for induction of labor with mechanical (transcervical balloon catheter) to 69% for induction with pharmacologic methods. There are no high-quality studies comparing VBAC rates with different induction methods [1].

Fetal Macrosomia

The most consistent infant factor associated with an increased likelihood of VBAC is birth weight less than 4000 g. As expected, large for gestational age/macroismia, especially birth weight >4000 g, decreases the odds of a VBAC after TOLAC [48]. The success rates for VBAC in women with a previous CD and no other births is about 60% in the >4000 g group and 71% in the ≤4000 g group [49]. There is a progressive reduction in VBAC success rates as birth weight increases. With a prior VBAC or a vaginal delivery, there is no success rate below 63% for any of the birth weight strata. But with no previous vaginal delivery, VBAC success rates can drop below 50% as neonatal weight exceeds 4000 g [36,48]. This success rate can decrease further if the indication for the previous CD is cephalopelvic disproportion or failure to progress. There is limited data regarding prenatal estimated fetal birth weight and VBAC success. Even so, VBAC rates after TOL were noted to be 60%–70% in fetuses suspected to be macrosomic [36]. This is likely due to the limitations of estimating birth weight by ultrasound. Thus, suspected macrosomia alone should not rule out a TOLAC [2].

Multiple Gestations

Vaginal delivery rates of both twins after prior CD generally range from 65% to 84% [12,50,51]. In most series, rates do not seem to differ from success rates in singletons, without increased risk for maternal morbidity or uterine rupture [50,51].

Prediction Tools for Vaginal Delivery

Given the associations above, several different scoring systems have been proposed to predict the likelihood of vaginal delivery or cesarean in women undergoing a TOLAC [17,36,43,52,53]. The best studies used scoring systems that were validated in separate data sets and in external studies [1]. These scoring systems performed as well in the validation testing as they did in the original data set. Unfortunately, although these models are accurate at predicting which women will have a VBAC, they are limited in their ability to predict who will have a repeat cesarean following a TOLAC [54,18]. Half of women with unfavorable risk factors had a vaginal delivery.

One available tool can be found at http://www.bsc.gwu.edu/mfmu/vagbirth.html but this tool only includes factors known at the beginning of pregnancy [33]. This model has been validated [55].

The authors of the NIH-sponsored review of prediction tools recommend clinicians use a sequential approach to screening [18] (Table 14.2). Initially, prenatal predictors such as previous vaginal delivery and previous indication for CD and possibly patient demographics and estimated fetal weight should be considered. This can then be modified by intrapartum predictors such as onset of spontaneous labor or the need for induction, and cervical status can be used to reevaluate the likelihood of vaginal delivery [17,54]. A variable that can be added to prediction of successful TOLAC is transvaginal ultrasound cervical length (CL) done at 19–23 weeks. Women with a CL <45mm have a 81% VBAC rate, while those with a CL ≥45mm have a 43% success rate [56].

Nevertheless, none of these screening tools is sensitive enough to be clinically helpful in predicting an unsuccessful trial of labor, and none have been validated prospectively to improve outcomes.

UTERINE RUPTURE

Complications and safety of TOLAC are especially related to the risk of uterine rupture. Uterine rupture is the most serious complication associated with TOLAC. It is defined as a complete separation through the entire thickness of the uterine wall (including serosa). At all gestational ages, pooled data from one review indicate that uterine rupture occurs in approximately 470/100,000 (0.47%, 95% CI 0.28–0.47) of women undergoing TOLAC and in 26/100,000 (0.026%, 95%
CI 0.009–0.082) of women undergoing a PRCD (RR 20.74, 95% CI 9.77–44.02; p < 0.001) [1]. At term, the overall risk for rupture is 778/100,000 (0.78%, 95% CI 0.62–0.96). This means that overall there are about 5 to 6 additional ruptures per 1000 in women undergoing a TOLAC [1]. No maternal deaths due to uterine rupture have been reported [1,57]. Perinatal death occurs in about 6% of women with uterine rupture, which translates to an overall rate of intrapartum fetal death of about 20/100,000, and a risk of perinatal death at term of about 0%–2.8% in women undergoing a TOLAC [1].

**Risk Factors for Uterine Rupture**
The risk for rupture is modified by a number of important factors [21] listed in Table 14.2 and described below. Prior vaginal delivery and prior VBAC reduce the risk of uterine rupture, while number of prior CDs, vertical scars, single-layer closure, induction or augmentation, greater maternal age, fever, prior preterm CD, and TOLAC at ≥40 weeks are associated with higher rupture rates.

**Maternal Age**
Maternal age (≥30 years old) is associated with an increased risk (1.4%) of uterine rupture [22].

**Prior Vaginal Delivery**
Prior vaginal delivery significantly reduces the risk for rupture during TOLAC from 1.1%, in women with no prior vaginal deliveries, to 0.2%. After controlling for maternal demographics and labor characteristics, prior vaginal delivery reduced the risk for rupture fivefold (OR 0.2, 95% CI 0.04–0.08) [23]. This protective effect was also confirmed by recent large retrospective (OR 0.30, 95% CI 0.23–0.62) [24] and prospective (OR 0.66, 95% CI 0.45–0.95) studies [10].

**Number of Prior Cesareans**
Women with 2 prior low transverse CD may be at increased risk of uterine rupture compared with women with one prior CD, but the absolute risk is small (1.36%), while the success rate is high (71%) [43]. Current ACOG recommendations allow this option regardless of whether there is a prior vaginal delivery [2]. Most ruptures occur in women undergoing induction or augmentation and in those without a prior vaginal delivery [42]. Earlier data had indicated an overall relative increased risk of rupture (about two- to threefold) in women having a TOL after two low transverse CD compared with one [29,57,58]. A meta-analysis confirmed these findings (0.9% one prior CD vs. 1.8% two prior CD) [42]. A follow-up case-control analysis of the study by Macones et al. [58] showed no increased risk (OR 1.46, 95% CI 0.87–2.44), and in the MFMU cohort [21] there was no difference in the incidence of rupture after two CDs compared with one (0.9% vs. 0.7%). Furthermore, the rupture rate in women with a prior vaginal delivery and two prior CD is only 0.5%, no greater than a VBAC with only one prior CD [58].

**Prior CD Characteristics**
**Labor Before Primary CD**
Labor before the primary CD has been associated with a lower risk of uterine rupture in a future TOLAC compared with no labor before the primary CD [19].

**Direction of Scar**
Classical and low vertical uterine scars have higher rates of rupture compared with low transverse uterine incisions (4%–10%, and 1%–2%, respectively) [2]. Records regarding the prior CD(s) should be obtained, with special care in documentation of direction of scar. If a woman has had a prior vertical (classical) CD, PRCD at 36 to 37 weeks is recommended [2,30].

**Layers of Closure**
There is insufficient evidence to assess whether the numbers of layers performed at prior uterine closure affects the outcomes, specifically the rate of uterine rupture, for future pregnancies. Randomized trials have insufficient follow-up numbers (see Chapter 13). Compared with women who had double-layer closure, women with a single-layer closure have been reported to have anywhere from similar to a fourfold increased risk of uterine rupture compared with those with a double-layer closure [31,32,59]. One large multi-center case-controlled study that examined operative reports found a fourfold increase in uterine rupture for women with a single layer uterine closure compared with those with a double-layer closure [OR 3.95; 95% CI, 1.35–11.49] [31]. However, a more recent study showed no difference in uterine rupture by layers of closure (OR 1.17; 95% CI 0.78–1.76) [32].

**Fever**
The presence of both intrapartum and postpartum fever at CD, but not either alone, may increase the risk of uterine rupture in a subsequent pregnancy [60].

**Previous Preterm CD**
The Maternal-Fetal Medicine Units Network (MFMU Network) data showed a higher risk of rupture in women with a prior preterm CD compared with in those with a prior term CD (1% vs. 0.68%; aOR 1.6, 95% CI 1.01–2.50) [61]. No difference in rupture rates were found in a secondary analysis of the recent retrospective, multicenter cohort study comparing prior CD before and after 34 weeks (aOR 1.5, 95% CI 0.7–3.5) [62].

**Prior CD for Twins**
Compared with women with prior CD for singletons, women with prior CD for twins have similar risk of uterine rupture and TOLAC success [63].

**Prior Uterine Rupture**
As prior uterine rupture is associated with high rates (6%–32%) of recurrent rupture with TOLAC, these pregnancies should have a PRCD before labor, possibly around 36 to 37 weeks [30].

**Interval between Deliveries**
Interval between deliveries of <18 months is associated with an increased risk of uterine rupture (2.3%–4.8%) compared with longer intervals (1.1%–1.3%) in a number of studies [25,26].

**Post Dates**
The risk of uterine rupture does not increase substantially after 40 weeks, but is increased with induction of labor regardless of gestational age [64,65].

**Preterm**
In one study there was a trend toward a lower uterine rupture rate in preterm patients who attempted a VBAC [45].
In the large MFMU cohort the rates of uterine rupture and dehiscence were lower in preterm TOL versus term; 0.34% versus 0.74%, \( p = 0.03 \) and 0.26% versus 0.67%, \( p = 0.02 \) respectively [66]. On the other hand women with a very preterm delivery (<26 weeks) with a prior low transverse CD may have an increased rate of rupture with subsequent TOL compared with women with a prior term low transverse CD, 1.8% versus 0.4%, respectively [67].

Macrosomia >4000 g
Macrosomia >4000 g is associated with a slightly increased risk of rupture [48]. A recent multicenter case-control study showed that birth weight >3500 g had twice the risk for rupture (OR 2.03, 95% CI 1.21–3.38) [48].

Twins
The risk of rupture or maternal mortality is not increased with prior CD and subsequent TOL with twins, with uncommon perinatal morbidity at ≥34 weeks [50,68].

Induction/Augmentation
The rate of uterine rupture in women with one prior low transverse CD undergoing TOLAC with spontaneous labor is about 0.4%. Almost all studies compare women undergoing induction with those in spontaneous labor instead of expectant management. These studies (including the large prospective MFMU observational trial) reported higher rates of uterine rupture (up to 1%–2%) with induction at any gestational age and of any kind [10,67,69]. There are no randomized controlled trials (RCTs) comparing these two groups but a recent meta-analysis of 8 studies comparing women with prior CD undergoing induction to those in spontaneous labor showed lower rates of vaginal delivery and higher rates of rupture among women undergoing induction but no differences in overall maternal or neonatal morbidity [70].

At term, the risk of rupture is not statistically significantly greater than women with spontaneous labor (OR 1.42, 95% CI 0.57–3.52). But women induced at >40 weeks have higher risk of rupture than those induced between 37 and 40 weeks (3.2% vs. 1.5%) [10]. In the MFMU cohort, induction increased the risk of rupture only in women without a prior vaginal delivery (induced 1.5%, vs. spontaneous 0.8%). In women with a prior vaginal delivery, the incidence of rupture was similar (0.6% vs. 0.4%) [46]. A recent case control study investigating 111 cases of uterine rupture did not find an increase in rupture compared with spontaneous labor even after controlling for labor duration. But the risk was still higher with an unfavorable cervix (<4cm) [71].

Recently two studies compared induction to expectant management instead of spontaneous labor, a more clinically appropriate comparison. One was a secondary analysis of the MFMU data and showed that induction had higher rates of vaginal delivery (73.8% vs. 61.3%) but also had higher rates of rupture (1.4% vs. 0.5%) [72]. The other was a secondary analysis of the Consortium on Safe Labor and found lower rates of vaginal delivery at 37–39 weeks but not at 40 weeks and no difference in uterine rupture [47].

Type of Induction
The risks of uterine rupture with induction in women who had one prior CD may also depend on the type of induction. Risk of rupture may be as high as 1.4%–2.5% with induction with prostaglandin (with or without oxytocin) [10,27], and about 1.1% with oxytocin alone [73]. The MFMU data, which included all prostaglandins, and Grobman et al. [46], which excluded misoprostol, did not find an increased risk for rupture when prostaglandins were used alone, but they did note a threefold increased risk with sequential use of prostaglandins and oxytocin versus spontaneous labor. The recent Cochrane review concluded that there is insufficient evidence from RCTs regarding method of induction (only two small RCTs comparing two different types of prostaglandins to oxytocin). One study was stopped due to two ruptures in the misoprostol group (2/17) [74].

ACOG therefore advises that induction with sequential use of prostaglandins and oxytocin be avoided [2].

ACOG considers the use of a Foley bulb for cervical ripening an acceptable method of labor induction as limited data appears to show no increased rates for rupture [2]. There are no randomized trials comparing induction to elective repeat cesarean delivery (ERCD). Women with prior CD should be made aware of these higher risks of rupture associated with induction.

Second Trimester Induction of Labor
Induction of labor in the second trimester with misoprostol is associated with 0.4% risk of uterine rupture (with 0% risk of hysterectomy and 0.2% risk of transfusion) after one prior low transverse CD, and about 50% risk with a prior classical CD [75]. There is insufficient evidence to assess the safety of mifepristone induction in women with a prior CD.

Augmentation of Labor
Augmentation may be associated with an increased risk of rupture, up to about 1%, but the risk is mainly related to higher doses of oxytocin [10,74,76]. A secondary analysis of a large retrospective cohort and a follow-up nested case-control study reported a dose-response relationship between the maximum oxytocin dose and increased risk for uterine rupture (fourfold increased risk with 21–30 mU/minute vs. no oxytocin; attributable risk of 2.9%–3.6%) [76,77]. Oxytocin should be used judiciously in women undergoing TOLAC, with a possible maximum dose of 20 mU/minute.

**Benefits and Harms Associated with TOLAC**
The majority of benefits and the least morbidity of a TOLAC are dependent on having a VBAC, while the potential harms of a TOLAC are associated with an unsuccessful TOLAC resulting in CD. Studies comparing TOLAC to ERCD usually include both VBAC and TOLAC ending in CD. Thus, the outcomes for women who have a TOLAC reflect both of these possibilities. These data are therefore appropriate for women with a previous CD who are deciding on the mode of delivery for the current pregnancy. Most studies show rates of all maternal and perinatal complications except rupture are infrequent and similar with both TOL and ERCD but these are small studies with few events [15,16,78,79].

One study of 2345 women with one prior CD compared outcomes among those with a planned ERCD and planned TOL and noted lower rates of serious perinatal complications (0.9% vs. 2.4%) and major maternal hemorrhage (0.8% vs. 2.3%) in the ERCD group. The vaginal delivery rate was 56.8% in the TOL group with similar rates of uterine rupture in both groups (0.1% vs. 0.2%) [8].
Overall rates of surgical injury are low, but may be slightly higher with TOLAC compared with PRCD. This is primarily due to those women who have a TOLAC but deliver with a repeat CD in labor [8–10,20,80,81]. There are no studies that specifically evaluated the risks of complications due to subsequent surgery after a TOLAC or PRCD. Nevertheless, increasing number of abdominal surgeries is commonly associated with increased risks of adhesions, injury to bowel and bladder, and other complications at the time of subsequent CD or nonpregnancy-related hysterectomy (Table 14.3).

Hysterectomy
The risk of hysterectomy is about 1–2/1000 TOLAC, and similar to PRCD [9,10,13,16,27,79,82].

Transfusion
The risk of transfusion is very low, and in most studies not significantly different for TOLAC or PRCD (0.9% vs. 1.2%) [1,10]. It has been estimated at about 2 units packed red blood cells (pRBC)/1000 TOLAC [13]. The risk is higher when labor is induced with no prior vaginal deliveries, and it is related to the need for CD following a TOLAC [10].

Venous Thromboembolism
The risk is about 4/10,000 with TOLAC, similar to PRCD (1/1000) [10].

Endometritis
The risk is very similar for TOLAC or PRCD with a 6% versus 7%–8% rate of fever [11]; or 3% versus 2% in academic centers [10]. The overall absolute risk for any fever with TOLAC is 6.5% [1]. Morbid obesity, cesarean after TOLAC, and increased number of CDs increase infection rates.

Hospitalization
Overall, because most women deliver vaginally, a TOLAC results in a shorter hospitalization compared with PRCD. This benefit does not apply to morbidly obese women.

Maternal Mortality
The risk of maternal mortality is lower overall for women having a TOLAC (3–4/100,000) compared with PRCD (13.4/100,000). This compares to the overall U.S. maternal mortality rate of 11–15/100,000. At term, maternal mortality is lower: 1.9 for TOLAC versus 9.6 PRCD per 100,000 live births [1]. Based on limited evidence, mortality is lower for TOLAC in high-volume hospitals (more than 500 deliveries per year) [7].

Short-Term Satisfaction
Anxiety, depression, psychological well-being, and satisfaction scores are similar between women who choose TOLAC and those who elect for PRCD [9].

Long-Term Maternal Risks
There is a clear association between the number of CDs and abnormal placentation as well as other surgical complications [80]. This complication is an important consideration for the 28% of women who have more than two births. For each CD, the risk for placenta previa significantly increases, occurring in 900, 1,700, and 3,000 per 100,000 women who have one, two, and three or more prior CDs. The overall risk for placenta accreta also increases (0.2% in the first CD to 0.3%, 0.6%, 2.1%, 2.3%, and 6.7% for second, third, fourth, fifth, and more than or equal to sixth CD). In the presence of placenta previa, the risk for abnormal placentation is markedly increased for each additional CD (3.3%,

Table 14.3 Maternal and Perinatal Risks of TOLAC versus PRCD

<table>
<thead>
<tr>
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<th>Favors</th>
<th>PRCD</th>
<th>TOLAC 1 prior CD</th>
<th>≥2 prior CDs</th>
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</thead>
<tbody>
<tr>
<td><strong>Maternal complications</strong></td>
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<tr>
<td>Uterine rupture</td>
<td>PRCD</td>
<td>1/200–250</td>
<td>1/111–140</td>
<td>1/55–140</td>
</tr>
<tr>
<td>Operative injury</td>
<td>PRCD</td>
<td>1/166–200</td>
<td>1/250</td>
<td>1/250</td>
</tr>
<tr>
<td>Hysterectomy</td>
<td>–</td>
<td>0–1/250</td>
<td>1/200–500</td>
<td>1/166</td>
</tr>
<tr>
<td>Transfusion</td>
<td>–</td>
<td>1/71–100</td>
<td>1/58–140</td>
<td>1/31</td>
</tr>
<tr>
<td>VTE</td>
<td>–</td>
<td>1/1,000</td>
<td>4/10,000</td>
<td>NA</td>
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<tr>
<td>Endometritis</td>
<td>–</td>
<td>1/47–66</td>
<td>1/34</td>
<td>1/32</td>
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<tr>
<td>Hospitalization</td>
<td>TOlAC</td>
<td>–</td>
<td>1/2,500–5,000</td>
<td>1/5,000</td>
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<tr>
<td>Death</td>
<td>–</td>
<td>1/2,500–5,000</td>
<td>1/5,000</td>
<td>NA</td>
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<td>Satisfaction</td>
<td>–</td>
<td>–</td>
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<td>Long-term risks</td>
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| **Perinatal complications** |        |            |                 |             |
| Stillbirth             | PRCD   | 1/1,000    | 2–4/1,000       | NA          |
| Low cord pH            | NA     | NA         | 1.5/1,000       | NA          |
| Neonatal death         | –      | 5/10,000   | 8/10,000        | NA          |
| Perinatal death        | PRCD   | 1/10,000   | 13/10,000       | NA          |
| Respiratory morbidity   | TOLAC  | 1–5/100    | 0.1–1.8/100     | NA          |
| Hyperbilirubinemia     | TOLAC  | 5.8/100    | 2.2/100         | NA          |
| HIE                   | PRCD   | 1/10,000   | 5–8/10,000      | NA          |


Abbreviations: HIE, hypoxic–ischemic encephalopathy; VTE, venous thromboembolism; NA, not available; PRCD, planned repeat cesarean delivery; CD, cesarean delivery.

Note: Some numbers may contradict each other as they are from different sources.

*See text.
11%, 40%, and >60% for the first, second, third, and more than or equal to fourth CDs, respectively [83]. Even without a placenta previa, the incidence of abnormal placentation, although much lower, increases with the number of CDs (see Chapter 25). The major benefit of TOLAC is the about >70% chance of a VBAC and avoidance of multiple CDs.

In addition, as noted above, women who undergo multiple CDs are at increased risk for many complications unrelated to placentation, including hysterectomy, adhesions, injury to bowel and bladder, transfusion of >4 units of pRBCs, need for postoperative ventilation, ICU admission, and increased operative time and hospital stay [83]. There is insufficient evidence regarding long-term pelvic floor function comparing TOLAC with PRCD. PRCD should not be used as a way to prevent pelvic floor disorders [1].

A decision analysis evaluating both the immediate and subsequent risks of the delivery decision for women with a prior CD suggested that for women planning only one additional pregnancy, PRCD results in fewer hysterectomies than TOLAC, while for women planning two or more additional pregnancies, a TOLAC provides the lowest risk [84].

Therefore, when counseling women about risks and benefits of a TOLAC compared with PRCD, the discussion should address her plans for future pregnancies.

Perinatal
The overall risk of serious perinatal complications is about 1 in 2000 TOLAC, which is slightly greater than that of PRCD [10]. Combining all poor perinatal outcomes, more than 600 PRCD would need to be performed to prevent one poor perinatal outcome. Although a woman with a TOLAC is at higher risk of uterine rupture than any other group, the risk of perinatal death is similar to that of any nulliparous woman in labor [16]. The most serious fetal risk in women with prior CD is from uterine rupture during TOLAC. Risks of fetal/neonatal morbidity/mortality with term uterine rupture are 33% risk of pH < 7.00, 40% admission to NICU, 6% risk of HIE, and 1.8% risk of neonatal death. The rupture-related risk of neonatal death is 1/10,000 in equipped academic centers [10]. In other centers, these risks are higher, including risk of neonatal death from rupture up to 10%–25%.

Stillbirth
The rates of stillbirths are 2 to 6/1000 for TOLAC versus 1 to 2/1000 for PRCD. At 39 weeks or more, the rate of antepartum stillbirth in the MMFU cohort was 1/1000 with TOLAC and 3/10,000 with PRCD [10,79]. This difference might be due to stillbirths occurring at <39 weeks with TOLAC, and/or to encouragement for TOLAC with diagnosis of stillbirth [10].

Low Cord pH
The risk of cord pH < 7.00 is about 1.5/1000 TOLAC [13].

Neonatal Death
Neonatal death rates are similar between the TOLAC and PRCD (0.08% vs. 0.05%, respectively) in academic centers, with the neonatal death rate associated with a rupture at term of about 1.8% [10].

Perinatal Death
Perinatal death rates at term (excluding malformations) are 1.3/1000 with TOLAC versus 0.5/1000 for PRCD [11]. Overall risk of adverse perinatal outcome is 1/2000 with TOLAC, slightly higher than with PRCD [10]. In one study of women at term with vertex presentations, the risk of delivery-related perinatal mortality with a TOLAC was 11 times that of PRCD (OR 11.6, 95% CI 1.6–86.7). When compared with women without a previous CD (instead of to those having a PRCD), the perinatal mortality for TOLAC was the same as that of a non-TOLAC multipara in labor [16].

Respiratory Complications
Rates of respiratory complications, transient tachypnea of the newborn, and need for oxygen and ventilator support may be slightly higher in infants delivered by PRCD versus those delivered by VBAC [1,85]. It is unclear if there is a difference in respiratory outcomes in infants born by PRCD versus those born by repeat CD after a TOLAC.

Hypoxic–Ischemic Encephalopathy
Aside from perinatal death, HIE is the most serious adverse outcome of uterine rupture and is one of the primary concerns regarding the decision to have a TOLAC. The incidence is 46/100,000 live births for TOLAC, versus none in the PRCD [10]. The overall risk of rupture-related HIE is 1 in 2,500 TOLAC. In term infants, the overall incidence for HIE is less than 10/100,000 live births.

MANAGEMENT
Patients with contraindications (Table 14.1) to TOLAC should receive a PRCD at 39 weeks, or earlier if labor starts or in certain cases (e.g., prior early uterine rupture). TOLAC can be offered if the risk of uterine rupture is estimated to be <2%, possibly <1% (Table 14.2), and if spontaneous labor starts by 40 weeks. If the patient reaches her EDC with no labor and an unfavorable cervix, a PRCD should be recommended.

Counseling
TOLAC can be offered to most women with a single or even two prior low transverse CDs, but several safety and success factors should be considered and discussed with the woman (Tables 14.1 through 14.3).

The composite of maternal complications is slightly higher with TOLAC compared with PRCD group primarily due to the risk of rupture, and the increased risks of a CD in labor. These estimates do not take into account the long-term increased risks of repeat CD and the associated risks of placenta previa and accreta. This is why counseling should take into account how many future pregnancies are planned.

The overall risk of serious perinatal complications is about 1 in 2000 TOLAC, which is slightly greater than that of PRCD [10]. Combining all poor perinatal outcomes, more than 600 PRCD would need to be performed to prevent one poor perinatal outcome. Although a woman with a TOLAC is at higher risk of uterine rupture than any other group, the risk of perinatal death is similar to that of any nulliparous woman in labor [16].

For the approximately 60%–80% of women having TOLAC who will deliver vaginally, the maternal and perinatal morbidity and mortality are lower than PRCD.

All women with a single prior low transverse CD without other indications for a CD are candidates for a TOLAC.

Women with two prior CD can also be considered for TOLAC, but induction should be avoided.

For women in whom the chance of having a vaginal delivery with a TOLAC is over 60%, the maternal and perinatal morbidity and mortality are lower than PRCD. About 50% of women with prior CD have a 70% or higher chance of
successful VBAC (http://www.bsc.gwu.edu/mfmuf/vagbirth.html), and they have no higher risk of maternal or perinatal morbidity and mortality compared with those who have PRCD. PRCD is safer than a TOLAC that results in a CD [15].

Although the overall risks of TOLAC for all women are higher than PRCD, the absolute risks are small and comparable to other potential complications of labor. Efforts to reduce the frequency of the first CD reduce the need for a TOLAC or repeat CD. Women at lowest risk for adverse outcomes and highest chance for a vaginal delivery include those with a prior vaginal delivery (especially a prior VBAC), in spontaneous labor, with a favorable cervix (Table 14.2). Other factors include younger age, normal BMI, prior CD for reason other than dystocia, and smaller fetus.

TOLAC should be approached with caution in those with the lowest chance of vaginal delivery and highest risk of rupture: for example, induction of labor in obese women over age 40 with an unfavorable cervix and no prior vaginal deliveries.

The ultimate decision regarding whether to have a TOLAC or PRCD should be based on the patient choice, after appropriate counseling and the availability of adequate resources and personnel to respond to obstetric emergencies. Most women should begin the decision process before term, and their decision should be documented in the medical record. The decision can then be modified at term in women to assess if spontaneous labor and/or favorable cervix make their chances of complications lower and of success higher. There is no evidence that examining the adequacy of the pelvis benefits outcomes.

Prenatal Education
Women who have had a prior CD need information and guidance to help them decide whether to have a TOL or an ERCD. Individualized prenatal education directed toward avoidance of a PRCD does not increase the rate of VBAC [86]. The Cochrane review evaluated three RCTs of decision support tools to help assist women in choosing ERCD or TOLAC and found no differences in the planned mode of birth or the percentage of women who remained unsure about their final decision [87]. A subsequent RCT showed that interactive or written evidenced based decision tools helped reduce conflict around the birth decision compared with baseline [88]. A recent study surveying a woman’s level of knowledge prior to TOLAC or ERCD showed most women had a poor understanding of the risks, benefits and likelihood of success of either option; they also were most likely to choose the mode of delivery favored by their provider [89].

Consent
Specific consent for TOL after CD or PRCD should be signed by the woman after appropriate counseling.

Checklist

Nonvertex Presentation
External cephalic version (ECV) can safely be performed in women with a prior CD. The success rate for ECV is similar or higher in women with a prior CD compared with controls without a prior CD (82% vs. 61%) [90]. Women with a successful version have successful VBAC rates of 65%–76% [90,91].

Ultrasound of Lower Uterine Segment
Due to the uncommon nature of rupture, several thousand women need to be studied to assess whether measuring the thickness of the lower uterine segment predicts complications in women with a prior CD who elect TOLAC, and therefore there is insufficient evidence to assess the clinical utility of this screening test. No women with a lower uterine segment thickness of ≥ 4.5 mm in the late third trimester seem to have dehiscence or rupture, while the proportion of these complications rises as this thickness decreases, with women with large defects, or lower uterine thickness < 3.5 mm (especially < 2.0 mm), or myometrial layer < 2.0 mm (especially < 1.4 mm) in the third trimester, possibly benefiting from PRCD [29,73]. A recent meta-analysis of 21 studies looking at data for 2776 women provided support for using ultrasound measurement of the lower uterine segment in assessing for the presence of a uterine defect [92].

Requirements to Minimize Risks [1]
To minimize risks, the following must be immediately available throughout TOLAC:
- Experienced obstetrician
- Anesthesia
- Nursing and OR personnel
- Ability to perform emergency CD

Labor and delivery units with >500 to 1000 births per year have lower risks of uterine rupture and complications compared with units with less volume [9,88]. If centers cannot provide the above resources, this does not mean TOLAC should not be offered. Referral should be made to adequate facilities early during prenatal care so that TOLAC is safely available.

Detecting and Managing Rupture Intrapartum
Fetal heart rate (FHR) disturbances are the most common (but not universal) sign of uterine rupture (55%–85%) [93]. The most commonly reported FHR disturbance is repetitive progressively severe variable decelerations and prolonged bradycardia, although in most cases they are not caused by rupture. Nevertheless, in women with a prior CD, in the presence of such FHR disturbances, uterine rupture must be considered. Abdominal pain over the area of the prior uterine scar is a poor predictor of uterine rupture. Epidural usually does not mask rupture. Epidural should not be withheld in women attempting TOL after prior CD. Intratuerine pressure catheter (IUPC) monitoring has not been shown to be helpful [94]. Significant loss of fetal station especially in the second stage may occur with rupture, but is of limited predictive value. There is insufficient data to assess the utility of uterine exploration after a successful VBAC.

Neonates delivered within 18 minutes after a suspected uterine rupture have the best outcome, with all normal umbilical pH levels and 5-minute Apgar scores in a recent series, while those with a decision-to-delivery time >30 minutes have poor outcomes [95].

Cost-Effectiveness
Multiple cost-effectiveness analyses have been performed to examine the relative cost of TOLAC versus PRCD. For women with at least a 47% likelihood of successful vaginal delivery TOLAC appears to be more cost-effective than ERCD [96,97]. In a recent analysis, TOLAC has similar cost-effectiveness for the first TOL but becomes less costly and more effective with subsequent deliveries [98].


84. Pare E, Quiñones JN, Macones GA. Vaginal birth after cesarean section versus elective repeat caesarean section: Assessment of maternal downstream health outcomes. *BJOG*. 2006;113(1):75–85. [II-2]
Early pregnancy loss
Lisa K. Perriera, Beatrice A. Chen, and Aileen M. Gariepy

KEY POINTS
• The diagnosis of early (e.g., first trimester) pregnancy loss may be suspected based on symptoms, but is usually made by transvaginal ultrasound, and/or serial beta human chorionic gonadotrophin (BHCG) levels.
• Early pregnancy loss (EPL) is an inclusive term that comprises the following: incomplete, inevitable, or complete spontaneous abortion (SAB); anembryonic gestation (blighted ovum); and embryonic/fetal demise (missed abortion). Early pregnancy failure (EPF) is a term that should be avoided as it contributes to internalization of blame for patients.
• There are three options for management of EPL: expectant, medical, and surgical management. Choice of management for EPL does not affect future fertility.
• Patient preference should guide treatment choice.
• Successful management of EPL consists of complete evacuation of the uterus. The success of each management option depends on several factors, for example, the type of loss (e.g., with or without symptoms).
• Threatened pregnancy loss can be defined as vaginal bleeding in pregnancy before 20 weeks of gestation. Several interventions have been studied, but none has been confirmed to be beneficial.

Medical management
• Medical management is a safe and effective alternative to expectant management or surgical curettage for EPL.
• Medical management of EPL is more effective than expectant management.
• Misoprostol 800 µg vaginally, with a repeat dose on day 3 if complete evacuation is not confirmed, has a success rate of 93% with incomplete or inevitable abortion, 88% with embryonic or fetal death, and 81% with anembryonic gestation in women at<13 weeks of gestation.
• Misoprostol 800 µg vaginally is the most studied regimen for medical management of EPL. Success of medical management of EPL increases with multiple dose regimens. Whether there is added benefit from adding mifepristone to misoprostol is still uncertain.
• Women choosing medical management of EPL report an average of 12 days of bleeding. Hemorrhage after medical management of EPL is rare.
• Follow-up: Transvaginal ultrasound after medical management of EPL can be used to confirm successful expulsion of the gestational sac. Measurement of endometrial thickness is not predictive of success.
• Advantages of medical management of EPL: Avoidance of surgery and anesthesia, perception of more natural treatment, increased privacy, and increased control.

Surgical management
• Surgical management has a high (>97%) success rate.
• Endometritis or hemorrhage rates are <1%.
• Maternal safety is highest with vacuum aspiration, when regional or general anesthesia can be avoided.

DEFINITIONS
• Pregnancy loss (PL): Spontaneous loss of pregnancy from conception to <20 weeks.
• Miscarriage: Lay term signifying PL.
• Early pregnancy loss: Inclusive medical term describing inevitable abortion, incomplete abortion, anembryonic pregnancy, and embryonic/fetal demise at <14 weeks [1]. It can also be called “first-trimester” PL. Early first-trimester PL is a loss of pregnancy between conception and 9 6/7 weeks. Late first-trimester PL is a loss of pregnancy between 10 and 13 6/7 weeks. The terms EPL, miscarriage, and SAB are often used interchangeably in the first trimester [2].
• Spontaneous abortion (aka loss): The term spontaneous abortion is often used as an equivalent term for EPL but should be avoided since women may associate negative feelings with this term. This guideline does not discuss voluntary (elective) termination (induced abortion).
• Complete abortion: Clinical definition describing an EPL that is characterized by a history of a positive pregnancy test, vaginal bleeding with passage of tissue, and a closed cervical os at the time of diagnosis. Transvaginal ultrasound examination shows absence of a gestational sac.
• Incomplete abortion: Clinical definition describing a history of positive pregnancy test, vaginal bleeding, and a cervical os that is open. Transvaginal ultrasound examination shows heterogeneous tissue distorting the endometrial canal with or without a gestation sac. There is no agreement on a measurement of endometrial thickness that can distinguish incomplete from complete abortion [3].
• Inevitable abortion: Clinical definition describing an EPL that is characterized by a history of a positive pregnancy test, vaginal bleeding without passage of tissue, gestational sac in the uterus, and an open cervical os.
• Anembryonic pregnancy: Previously described as “blighted ovum.” This is a gestational sac without a visible yolk sac and/or embryo (with no heart motion) in relationship to the mean sac diameter (MSD) size (Table 15.1). It occurs when the embryonic disk has failed to develop or has already been resorbed [5].
• **Embryonic/fetal demise**: Previously described as “missed abortion.” It is defined by the failure of a previously identified embryo to grow and/or retain cardiac activity over time (Table 15.1).

• **Expectant management**: No intervention. Awaiting natural passage of tissue.

• **Medical management**: The use of medications to expel the products of conception.

• **Surgical management**: The mechanical removal of the products of conception.

**DIAGNOSIS**

**Transvaginal Ultrasound**

To ensure 100% specificity to confirm EPL, the diagnostic criteria shown in Table 15.1 were adopted by the Society of Radiologists in Ultrasound Multispecialty Panel on Early First Trimester Diagnosis of Miscarriage and Exclusion of a Viable Intrauterine Pregnancy [4].

It is important to recognize that not all patients may desire 100% certainty of PL [2]. The wishes of the patient must be considered when determining the level of diagnostic certainty that will be used. Table 15.2 lists the sensitivity and specificity of different diagnostic cutoffs that can be used to counsel patients [4,6].

**Beta Human Chorionic Gonadotropin**

In a clinically stable patient with a highly desired pregnancy, a single BHCG level ≥ 3000 mIU/mL cannot differentiate between an ectopic pregnancy or EPL if a gestational sac is not visualized in the uterus [4]. If the pregnancy is desired, ectopic precautions should be given and intervention should be avoided until additional testing is performed.

**SYMPTOMS**

Symptoms of EPL include vaginal bleeding, lower abdominal cramping, and dilation of cervix. Women with EPL may also be asymptomatic.

**EPIDEMIOLOGY/INCIDENCE**

Fifteen to twenty percent of clinically recognized pregnancies end in EPL [7]. It is estimated that up to 60% of conceptions become an EPL, and most are not clinically recognized (e.g., “late cycle”). Human reproduction is relatively inefficient. Only 30% of fertilized eggs result in a viable pregnancy. Sporadic early PL is very common in humans. At least 15%–20% of clinically identified pregnancies (implanted) physiologically end with early PL, and only 50%–60% of all conceptions advance to >20 weeks [8]. Most PLs represent failure of implantation and are difficult to recognize clinically. Oocyte quality and normal karyotype are most important for normal implantation, a lot more than uterine factors. The prognosis after one uncomplicated early PL in a healthy young woman is for >70%–80% chance of a viable pregnancy in the successive pregnancy. Therefore, no workup or therapy is usually indicated after one PL. For women with 2 or more EPLs, see Chapter 16, “Recurrent Pregnancy Loss.”

**ETIOLOGY**

Chromosomal abnormalities are responsible for >50% of all spontaneous EPL, most commonly aneuploidy. When tissue from EPLs <13 weeks (81% < 9 weeks) obtained by chorionic villus sampling (CVS) and manual vacuum aspiration (MVA) is analyzed, about 72% of cases revealed aneuploidy, with trisomy being the most common [9]. Many spontaneous losses may be secondary to other genetic defects that are impossible to discern by simple karyotype. Many other factors are also associated with spontaneous losses (see Chapter 16).

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**Table 15.1** Diagnostic Criteria for EPL

<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>Ultrasound Findings</th>
</tr>
</thead>
</table>
| **Anembryonic pregnancy** (any of the three) | • A gestational sac ≥ 25 mm MSD without an embryo  
|                                    | • Absence of an embryo with cardiac activity ≥ 11 days after an ultrasound showed a gestational sac with a yolk sac  
|                                    | • Absence of an embryo with cardiac activity ≥ 2 weeks after an ultrasound showing a gestational sac without a yolk sac  
| **Embryonic/fetal demise**         | • Absence of cardiac motion in an embryo measuring ≥ 7 mm |

**Table 15.2** Sensitivity, Specificity, and False Positive Rates When Diagnosing EPL

<table>
<thead>
<tr>
<th></th>
<th>Sensitivity (95% CI)</th>
<th>Specificity (95% CI)</th>
<th>FPR (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CRL &gt; 5 mm without cardiac activity</td>
<td>0.50 (0.12–0.88)</td>
<td>1.00 (0.90–1.00)</td>
<td>0</td>
</tr>
<tr>
<td>CRL &gt; 6 mm without cardiac activity</td>
<td>0.50 (0.07–0.93)</td>
<td>1.00 (0.87–1.00)</td>
<td>0</td>
</tr>
<tr>
<td>CRL &gt; 7 mm without cardiac activity</td>
<td>NA</td>
<td>1.00 (NA)</td>
<td>NA</td>
</tr>
<tr>
<td>MSD ≥ 13 mm without yolk sac</td>
<td>0.96 (0.92–0.99)</td>
<td>1.00 (0.69–1.00)</td>
<td>0</td>
</tr>
<tr>
<td>MSD ≥ 16 mm without yolk sac</td>
<td>0.50 (NA)</td>
<td>1.00 (0.88–1.00)</td>
<td>NA</td>
</tr>
<tr>
<td>MSD ≥ 20 mm without yolk sac</td>
<td>0.41 (0.30–0.52)</td>
<td>1.00 (0.96–1.00)</td>
<td>0</td>
</tr>
<tr>
<td>MSD ≥ 25 mm without yolk sac</td>
<td>NA</td>
<td>1.00 (NA)</td>
<td>NA</td>
</tr>
</tbody>
</table>

**Abbreviations:** CI, confidence interval; CRL, crown rump length; FPR, false positive rate; NA, not available.
RISK FACTORS
The risk of EPL increases with maternal age, ranging from 9% at 22 years old, to 84% at 48 years old [10]. Other risk factors include smoking, alcohol, excessive caffeine intake, African racial origin, previous EPL, previous stillbirth, medical complications such as diabetes, ART, and vaginal bleeding [11]. A recent meta-analysis showed that caffeine and coffee consumption, especially more than three servings per day, during pregnancy are significantly associated with PL [12].

THREATENED PL
Threatened PL can be defined as vaginal bleeding in pregnancy before 20 weeks of gestation. Several interventions have been studied, but none has been confirmed to be beneficial.

Complications
- Vaginal bleeding in the first trimester has been associated with several complications in pregnancy, including antepartum hemorrhage (odds ratio [OR] 2.47), preterm premature rupture of membranes (PPROM) (OR 1.78), preterm birth (PTB) (OR 2.05), intrauterine growth restriction (IUGR) (OR 1.54), low birth weight (LBW) (OR 1.83), and perinatal mortality (OR 2.15) [13].
- The presence of a subchorionic hematoma detected on ultrasound (usually between 5 and 20 weeks) is associated with increased risks of SAB (OR 2.18), abruptio (OR 5.71), intrauterine fetal demise (IUFD) (OR 2.09), PPROM (OR 1.64), and PTB (OR 1.40) [14].

Prevention
- Lifestyle modifications
  - Epidemiologic studies suggest that lifestyle modifications can increase fertility potential, although these have not been definitively tested in randomized controlled trials (RCTs). These modifications include eliminating use of tobacco products, alcohol, and caffeine and reduction in body mass index (BMI) [15].
- Multivitamins
  - In non-high-risk women, multivitamin supplementation before 20 weeks is associated with similar total risks of SAB (OR 2.18), abruptio (OR 5.71), intrauterine fetal demise (IUFD) (OR 2.09), PPROM (OR 1.64), and PTB (OR 1.40) [14].
- Progesterone
  - There is insufficient evidence to support the routine use of progesterone (synthetic or natural) to prevent miscarriage in early to midpregnancy. The meta-analysis of all women, regardless of gravidity and number of previous miscarriages, showed no statistically significant difference in the risk of miscarriage between progesterone and placebo or no treatment groups (OR 0.99; 95% confidence interval [CI] 0.78–1.24) and no statistically significant difference in the incidence of adverse effects in either mother or baby [17].
  - However, in a subgroup analysis of four trials involving women who had recurrent miscarriages (3+ consecutive miscarriages; four trials, 225 women), progesterone treatment showed a statistically significant decrease in the miscarriage rate compared with placebo or no treatment (OR 0.39; 95% CI 0.21–0.72) though these studies were of poorer quality.
  - While progesterone supplementation cannot be recommended for prevention of miscarriage, there may be benefit in women with a history of recurrent miscarriage. Treatment for these women may be warranted (see Chapter 16).

Therapy
- Bed rest
  - There is insufficient evidence of high quality that supports a policy of bed rest to prevent miscarriage in women with confirmed fetal viability and vaginal bleeding in the first half of pregnancy. There is no statistically significant difference in the risk of miscarriage in the bed rest group versus the no bed rest group (placebo or other treatment) (relative risk [RR] 1.54, 95% CI 0.92–2.58). Neither bed rest in a hospital nor bed rest at home shows a significant difference in the prevention of miscarriage. There is a higher risk of miscarriage in those women in the bed rest group than in those in the hCG therapy group with no bed rest (RR 2.50, 95% CI 1.22–5.11). The small number of participants included in these studies makes these analyses inconclusive [18].
  - Bed rest cannot be recommended for prevention of miscarriage. In fact, it might be harmful, given the higher rates of venous thromboembolism (VTE), muscle atrophy, and other detriments associated with bed rest.

- Progesterone
  - There is insufficient evidence to assess the effect of progesterone supplementation in women with threatened miscarriage. There was no evidence of effectiveness with the use of vaginal progesterone compared with placebo in reducing the risk of miscarriage (RR 0.47, 95% CI 0.17–1.30) [19].
  - Human chorionic gonadotropin
    - The current evidence does not support the routine use of hCG in the treatment of threatened miscarriage. There is no statistically significant difference in the incidence of miscarriage between hCG and “no hCG” (placebo or no treatment) groups (RR 0.66, 95% CI 0.42–1.05). There were no reported adverse effects of hCG on the patient or fetus [20].

- Muscle relaxant
  - There is insufficient evidence to support the use of uterine muscle relaxant drugs for women with threatened miscarriage, and therefore they should not be used. In one poor-quality RCT, compared with placebo, buphenine (a β-agonist) was associated with a lower risk of intrauterine death (RR 0.25, 95% CI 0.12–0.51). PTB was the only other outcome reported (RR 1.67, 95% CI 0.63–4.38) [9] [21].

MANAGEMENT OF EPL
General Principles
- There are three main options for the woman with EPL (unless the spontaneous loss is complete): expectant, medical, and surgical management.
- Successful management of EPL entails complete evacuation of the uterus. The success of each management option depends on several factors, for example, the type of loss (e.g., with or without symptoms).
• Failure of expectant or medical management results in the need for surgical evacuation.
• There are several types of medical management approaches and several surgical approaches.

Principles of Surgical Management
• Procedure: Surgical management of EPL can be accomplished via vacuum aspiration or sharp curettage.
• When comparing vacuum aspiration versus sharp curettage, vacuum aspiration is preferred, as it is associated with [22]
  • Less blood loss (mean difference [MD] -17.10 mL, 95% CI -24.05 to -10.15 mL).
  • Less pain during the procedure (RR 0.74, 95% CI 0.61-0.90).
  • Shorter duration of the procedure (MD -1.20 minutes, 95% CI -1.53 to -0.87 minutes).
• The small sample sizes of the trials were too small to evaluate rare complications such as uterine perforation and other morbidity.
• Vacuum aspiration can be accomplished with electric vacuum aspiration (EVA) or MVA. Both EVA and MVA are types of dilation and curettage (D&C).
• Location: Vacuum aspiration can be accomplished in the operating room or in the office.
• MVA is a safe alternative for gestations 6–12 weeks with EPL, and can be performed in the office under local anesthesia [23]. This should be strongly preferred at these gestational ages instead of an operating room procedure necessitating general anesthesia [24].
• There is no significant difference in the success or complication rates for MVA versus EVA [25].

Principles of Medical Management
Medications used
• Misoprostol is a prostaglandin E1 analog. It is a uterotonic that results in cervical softening and contractions that expel the products of conception. Routes of administration include vaginal, oral, buccal, or sublingual. Side effects vary, based on route of administration [26].
• Mifepristone is an antiprogestin that results in weakening of the uterine attachment of a pregnancy. This results in capillary breakdown and synthesis of prostaglandins [27].
• Methotrexate (intramuscularly or oral) antagonizes folic acid, a cofactor needed for synthesis of nucleic acids. It is toxic to the rapidly dividing cells of the trophoblast. There is no role for methotrexate in the treatment of EPL [28].
• Success of medical management is determined by the absence of significant symptoms and absence of the gestational sac on transvaginal ultrasound. Studies that use an endometrial thickness of >15 mm to define failure of medical management may underestimate success rates of expectant and medical management [4].
• Success of medical management is higher in patients with symptoms, such as cramping and bleeding [29].

Contraindications
The contraindications listed in Table 15.3 apply to medical management but can also apply to expectant management [25,30,31].

Complications
• Complications are rare. The incidence of gynecologic infection after surgical, expectant, or medical management of EPL is low (2%–3%). There is no evidence to show a differential risk of infection by management choice [32].

Table 15.3 Contraindications to Medical (or Expectant) Management

<table>
<thead>
<tr>
<th>Contraindications to Medical Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Hemodynamically or medically unstable patients</td>
</tr>
<tr>
<td>• Signs of pelvic infection and/or sepsis</td>
</tr>
<tr>
<td>• Suspected molar or ectopic pregnancy</td>
</tr>
<tr>
<td>• Allergy to prostaglandins*</td>
</tr>
</tbody>
</table>


*Specific to medical management.

Principles of Expectant Management
Expectant management of EPL is an option for women who would like to avoid surgical or medical treatment of EPL; however, time until resolution of EPL is unpredictable and may take as long as 6 weeks. Success of expectant management can range from 25% to 83%, depending on length of time of follow-up, definition of “failed” expectant management, and inclusion criteria [33–35].

Medical versus Surgical Management
• Cochrane review [36]
  • Twelve RCTs were included in this Cochrane review of incomplete abortion before 13 weeks with N = 2894.
  • Misoprostol routes varied, including six studies via oral, four studies via vaginal, one study via sub-lingual, and one study via both vaginal and oral.
  • There was a slightly lower incidence of complete miscarriage in the misoprostol group (RR 0.97, 95% CI 0.95–0.99), but high success in both groups.
  • Women using misoprostol had fewer surgical procedures (RR 0.06, 95% CI 0.02–0.13).
  • Risk of unplanned procedure was higher with misoprostol (RR 5.82, 95% CI 2.93–11.56).
  • Deaths and “serious complications” with either management are too rare to compare.

• Largest multicenter RCT [25]
  • N = 652; 491 treated with 800 mg vaginal misoprostol versus 161 by vacuum aspiration
  • Participant characteristics:
    • 58% embryonic/fetal demise
    • 36% anembryonic gestation
    • 6% incomplete/inevitable abortion
  • Medical regimen: 800 mg vaginal misoprostol, repeated on day 3 if incomplete expulsion (diagnosed by persistence of gestational sac or endometrial lining greater than 30 mm on transvaginal ultrasound), vacuum aspiration on day 8 if still incomplete
Medical versus Expectant Management

Medical management commonly uses vaginal misoprostol [28]. There are several additional studies looking at mifepristone followed by vaginal or oral misoprostol [37–39].

- Cochrane review [28]
  - Twenty-four RCTs, n = 1888, of embryonic/fetal demise or anembryonic pregnancy.
  - Vaginal misoprostol compared with expectant management:
    - Shortens the time to achieve complete uterine evacuation:
      - At less than 24 hours after treatment (RR 4.73, 95% CI 2.70–8.28)
      - At least 48 hours after treatment (RR 5.74, 95% CI 2.70–12.19)
    - Results in less need for uterine curettage
    - Does not show a significant difference in need for blood transfusion (RR 0.2, 95% CI 0.01–4.0)
    - Does not have a significant increase in nausea (RR 1.38, 95% CI 0.43–4.40) or diarrhea (RR 2.21, 95% CI 0.35–14.06)
  - Dosage of vaginal misoprostol: When compared with lower dosages, 800 mg vaginal misoprostol is more effective at completing uterine emptying (RR 0.85, 95% CI 0.72–1.00) with similar incidence of nausea.
  - No advantage of “wet” versus “dry” preparation of vaginal misoprostol or of adding methotrexate.
  - Oral misoprostol is less effective than vaginal misoprostol in emptying the uterus (RR 0.90, 95% CI 0.82–0.99).
  - Sublingual misoprostol is equivalent to vaginal misoprostol in inducing complete uterine emptying but was associated with more frequent diarrhea.

- Mifepristone
  - Efficacy of mifepristone followed by misoprostol for treatment of miscarriage ranges from 65.5% to 80% [37–39].
  - In a Cochrane review published in 2006, two trials of mifepristone added to misoprostol show conflicting results [28].
  - One small RCT (n = 115) found that mifepristone in addition to misoprostol did not improve efficacy [39], though one retrospective study did find increased expulsion rates with mifepristone plus misoprostol versus misoprostol alone [40].

Conclusion: Medical management with 800 mg of vaginal misoprostol is significantly more effective than expectant management. Mifepristone and misoprostol appear to have similar success rates to misoprostol alone though data are conflicting, thus based on available evidence, it is uncertain whether there is any advantage to adding mifepristone to misoprostol for medical management.

Expectant versus Surgical Management

In a Cochrane review of seven RCTs, n = 1521: [41]

- Expectant management has a higher incidence of the following:
  - Incomplete miscarriage by 2 weeks (RR 3.98, 95% CI 2.94–5.38) or by 6–8 weeks (RR 2.56, 95% CI 1.15–5.69).
  - Need for unplanned or additional surgical emptying of the uterus (RR 7.35, 95% CI 5.04–10.72).
  - Surgical management is required in 28% in expectant group, while only 4% in surgical group require additional surgery.
  - More days of bleeding (weighted mean difference [WMD] 1.59, 95% CI 0.74–2.45).
  - Need for blood transfusion (RR 6.45, 95% CI 1.21–34.42).
  - There was no difference in infection risk.
  - Cost was lower in the expectant management group.

Conclusion: After counseling regarding the data above, patient preference should guide decision making.

Misoprostol: Route, Dose, and Safety

There is published literature on a wide range of therapeutic regimens [4,30,42]. Optimal doses and routes of administration of misoprostol have not been determined by randomized trials.

- Misoprostol 800 µg per vagina is the most studied regimen for medical management of EPL. Success of medical management of EPL increases with multiple-dose regimens [25].
- A single RCT showed that there is equivalent efficacy between a multidose regimen of vaginal misoprostol 400 and 800 µg with a lower incidence of fever/rigors and higher satisfaction in the lower dose group [43].
- An international panel of experts recommends a single oral dose of 600 µg misoprostol for medical management of incomplete abortion [30] and a single vaginal dose of 800 µg misoprostol for medical management of anembryonic pregnancy and embryonic/fetal demise [44]. Misoprostol 600 µg sublingual is an alternative regimen [44].
- Overall, misoprostol is safe and well tolerated. Side effects of prostaglandins include diarrhea, nausea, and vomiting. These side effects are increased when misoprostol is given orally. Patients receiving misoprostol vaginally have decreased gastrointestinal side effects and prolonged duration of action when compared with oral administration [26].

Antibiotics

There is insufficient evidence to recommend or to abandon prophylactic antibiotics for surgical evacuation in women with an incomplete abortion. Clinical judgment is recommended [30,45].
There is insufficient evidence to recommend prophylactic antibiotics for women undergoing surgical evacuation of the uterus for embryonic/fetal demise or anembryonic gestation but the risk of infection is thought to be similar to risk of infection for induced termination of pregnancy. American Congress of Obstetricians and Gynecologists (ACOG) recommends the use of a single preoperative dose of doxycycline in cases of EPL, but acknowledges the lack of data [2]. If provided, doxycycline 100 mg 1 hour before the procedure and 200 mg after the procedure is a low cost regimen [46].

Rh Negative
Women who are Rh(D) negative and unsensitized should receive Rh(D)-immune globulin within 72 hours of the EPL [2].

PATIENT COUNSELING
Patients choosing medical management of EPL should have appropriate counseling regarding expected symptoms.

- Bleeding with medical management is heavier and longer in duration than with surgical management, and rarely requires intervention [47].
- Women experienced approximately 12 days of bleeding after medical management of EPL [47]. Bleeding will most likely be heavy for about 3–4 days, followed by light bleeding or spotting for several weeks [31].
- Patients should be counseled to contact their physician if they experience heavy vaginal bleeding (soaking through more than two extra large sanitary pads per hour for 2 consecutive hours) or signs of infection [31].
- Some women experience fever and/or chills during the first 24 hours after misoprostol use. Patients should call their doctor and be evaluated for infection if fever and/or chills persist beyond 24 hours after using misoprostol [31].
- Nausea and vomiting may occur with use of misoprostol and will usually resolve 2–6 hours after taking misoprostol [31].
- Pain should be expected with medical or expectant management and patients should be given a narcotic pain medication and non-steroidal anti-inflammatory drugs to treat pain [2].

PATIENT ACCEPTABILITY
- In one study, the majority of women would prefer medical management of EPL with misoprostol to surgical management if its efficacy is >65% [48].
- In a large RCT comparing medical versus surgical management of EPL, women receiving medical management had significantly higher reports of treatment-related symptoms (cramping and bleeding), but overall quality of life and treatment acceptability were similar [49].
- In a large RCT comparing medical versus surgical management of EPL, 83% of women with EPL randomized to medical management with misoprostol would recommend medical management to others and 78% would probably/absolutely use medical management again [25].

PATIENT PREFERENCE
Most women have strong preferences regarding management of EPL [50]. Preferences are diverse and different women place different values on the advantages and disadvantages of avoiding the OR or of miscarrying at home, for example. Patient preference will depend on individual circumstances, expectations, and awareness of the advantages and disadvantages of each management option. Women have greater satisfaction when treated according to their preferences.

Due to the comparable safety and efficacy of all current treatment options for EPL, patient preference should be the guiding force deciding management of EPL.

COST
Expectant management has a lower cost when compared with surgical management [41]. In a decision analysis comparing surgical and expectant management to an expanded care option (expectant management, surgical management in the office or OR, and medical management), there was a cost savings of $241.39 per case in the expanded care option [51].

Follow-Up
There are no RCTs to assess optimal management of follow-up after EPL. Transvaginal ultrasound is the most common follow-up after medical or expectant management of EPL, typically done within 7–14 days [2]. Absence of the gestational sac indicates success [2]. Endometrial thickness after medical or expectant management is not predictive of retained products of conception and/or need for surgical evacuation [52].

If chorionic villi or a gestational sac is obtained at D&C, there is usually no need for BHCG follow-up. After expectant or medical management of a known intrauterine pregnancy, BHCG levels, in general, do not need to be followed. Ultrasound can be used to confirm expulsion of the gestational sac. BHCG can be used in patients with limited access to ultrasound [2].

FUTURE FERTILITY
There is no need for a work-up after one EPL. If >1 EPL has occurred, see Chapter 16, Recurrent PL. There is insufficient data to suggest an optimal interpregnancy interval between an EPL and the next conception [2]. There seems to be no improvement in outcome associated with waiting 3 months of more, as previously recommended [53]. There are no contraindications to the placement of an intrauterine device immediately after surgical (or other) management of EPL, as long as septic abortion is ruled out [54]. Choice of management for EPL does not affect future fertility. In long-term follow-up of women participating in an RCT of expectant, medical, or surgical management of EPL, there was no significant difference in the live birth rate 5 years after the index miscarriage [55].

- Expectant management: 177/224 (79%, 95% CI 73%–84%)
- Medical management: 181/230 (79%, 95% CI 73%–84%)
- Surgical management: 192/235 (82%, 95% CI 76%–86%)

REFERENCES


Recurrent pregnancy loss
Reshama Navathe and Michela Villani

KEY POINTS
- Diagnosis of recurrent pregnancy loss (RPL) is more than or equal to two consecutive losses or three nonconsecutive losses of pregnancy <20 weeks.
- Workup includes uterine study, antiphospholipid antibodies (APAs), and parental karyotypes, as well as karyotype of products of conception (POC) (if available).
- Prognosis with negative workup is for a 60%–70% subsequent successful pregnancy in women <35 years old and 40%–50% in women ≥35 years old.
- Women with RPL and APA should be treated with low-dose aspirin and heparin in subsequent pregnancy (see Chapter 26 in Maternal-Fetal Evidence Based Guidelines). Women with unexplained RPL should not receive anticoagulant therapy.
- Women with RPL and uterine septum, synechiae, or submucous myomata can consider hysteroscopic resection of these abnormalities.
- Couples with abnormal parental karyotype can be offered genetic counseling, prenatal diagnosis, and/or gamete donation.
- There is insufficient evidence for universal screening for diabetes mellitus (DM), thyroid disease, progesterone deficiency (luteal phase defect [LPD]), infections, thrombophilia, etc.
- Women should not be tested for alloimmunization or receive any of the immune therapies, since they are ineffective and at times detrimental.
- Women with RPL, especially more than or equal to three losses, should be offered progesterone until 10 weeks gestation in subsequent pregnancies.
- There is insufficient evidence to support human chorionic gonadotropin (hCG), aspirin, and vitamins as interventions.

DIAGNOSIS/DEFINITIONS
For diagnoses of pregnancy loss, miscarriage, spontaneous and other kind of abortions, anembryonic pregnancy, embryonic demise, etc., please see Chapter 15.

RPL: either two consecutive pregnancy losses, or three nonconsecutive pregnancy losses before the twentieth week of gestation [1], excluding ectopic and molar pregnancies.

The definition of RPL varies in different publications, which makes diagnosing and treating this entity more challenging for physicians and couples. Early pregnancy loss is usually defined as a pregnancy loss <14 weeks, and these make up the majority of RPL.

There are several guiding committees with slightly different variations on this definition (Table 16.1) [2–4].

CLASSIFICATION
Primary RPL: no intervening live births; secondary RPL: intervening live births. The prognosis is better with secondary RPL [5].

INCIDENCE
As seen in Chapter 15, most fertilized eggs do miscarry, often without a recognized clinical pregnancy. Approximately 15% of pregnant women experience loss of a clinically recognized pregnancy. It is estimated that fewer than 5% of women experience two consecutive pregnancy losses, and only 1% experience 3 or more [6].

ETIOLOGY
Etiology of RPL is not established in at least 50% of cases after workup (see below). General categories include: genetic, anatomic, endocrine, immunologic, thrombophilic, and environmental (Table 16.2) [7,8].

RISK FACTORS/ASSOCIATIONS
Advancing maternal age is associated with a higher rate of RPL. Rate of clinically recognized miscarriage is 20% at age 35 years, 40% at age 40 years, and up to 80% at age 45 years [9]. Other risk factors include maternal medical diseases, especially poorly controlled, such as hypertension and diabetes (see Chapter 15).

A previous PL is a risk factor for a subsequent PL (Table 16.3) [6,10]. Women with only unsuccessful pregnancy history have a greater risk of future miscarriage than primigravidae and women with a history of previous successful pregnancy [11].

RECURRENT PREGNANCY LOSS Counseling
The frequency of PL should be reviewed with the patient and partner (see Chapter 15), as well as the prognosis after one or more PLs (Table 16.3) [6,10].

Workup (Screening)
Appropriate diagnostic workup of RPL is essential for choosing the proper intervention (Table 16.4) [12]. Screening tests should not only discover diagnosis (etiology) but also lead to interventions effective in increasing the incidence of
The following women should be offered evaluation:

1. Women with more than or equal to two consecutive miscarriages
2. Women with more than or equal to three nonconsecutive miscarriages

Initial part of workup consists of history (smoking, alcohol and caffeine, illicit drug use, environmental exposures, working conditions, as well as detailed obstetrical and gynecological history) and physical examination (pelvic). Eliciting the gestational age of miscarriage is important, as often RPL occurs at a similar gestational age in subsequent pregnancies, and the most common cause of RPL varies by gestational age. Sometimes, though, obstetrical history is mixed, with early PL, second-trimester PL, preterm birth (PTB), and/or fetal death, so that workup may include other tests (see specific chapters, including Chapter 55 in *Maternal-Fetal Evidence Based Guidelines*).

### Table 16.1 Definitions of RPL

<table>
<thead>
<tr>
<th>Committee</th>
<th>Number of Pregnancies</th>
<th>Consecutive?</th>
<th>Specifics</th>
</tr>
</thead>
<tbody>
<tr>
<td>RCOG 2011 [2]</td>
<td>3</td>
<td>Yes</td>
<td>Not required to be intrauterine</td>
</tr>
<tr>
<td>ASRM 2013 [3]</td>
<td>2</td>
<td>No</td>
<td>Clinical pregnancies confirmed by histology or ultrasonography</td>
</tr>
<tr>
<td>ESHRE 2014 [4]</td>
<td>Not specified</td>
<td>No</td>
<td>Intrauterine, confirmed by histology or ultrasonography</td>
</tr>
</tbody>
</table>

**Abbreviations:** ASRM American Society of Reproductive Medicine; ESHRE: European Society of Human Reproduction and Embryology; RCOG, Royal College of Obstetricians and Gynecologists; RPL, recurrent pregnancy loss.

### Table 16.2 Etiology, Diagnostic Considerations, and Treatment in RPL

<table>
<thead>
<tr>
<th>Etiology</th>
<th>Percentage</th>
<th>Diagnostic Evaluation</th>
<th>Therapy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Genetic</td>
<td>2–5</td>
<td>Karyotype—parental</td>
<td>Genetic counseling</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Karyotype—POC</td>
<td>Genetic counseling, PGD</td>
</tr>
<tr>
<td>Anatomic</td>
<td>12–22</td>
<td>3D ultrasound</td>
<td>Correction of anatomic defect</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Hysterosalpingogram</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>SIS</td>
<td></td>
</tr>
<tr>
<td>Endocrinology</td>
<td>20</td>
<td>TSH</td>
<td>Levothyroxine</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Prolactin</td>
<td>Bromocriptine</td>
</tr>
<tr>
<td></td>
<td></td>
<td>HgbA1c</td>
<td>Diabetic optimization</td>
</tr>
<tr>
<td></td>
<td></td>
<td>APLAS</td>
<td>Aspirin + heparin</td>
</tr>
<tr>
<td>Immunologic/thrombophilic</td>
<td>15–25</td>
<td>Tobacco, EtOH</td>
<td>Eliminate exposures</td>
</tr>
<tr>
<td>Environmental</td>
<td></td>
<td>Exposure</td>
<td></td>
</tr>
<tr>
<td>Unknown</td>
<td>40–50</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>


**Abbreviations:** 3D, three-dimensional; APLAS, antiphospholipid antibody syndrome; EtOH, ethyl alcohol; HgbA1c, hemoglobin A1c; PGD, preimplantation genetic diagnosis; POC, products of conception; SIS, saline infusion sonogram; TSH, thyroid stimulating hormone.

### Table 16.3 Miscarriage Risk after Prior PL

<table>
<thead>
<tr>
<th>Prior PL (n)</th>
<th>Risk of Miscarriage in Subsequent Pregnancy (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>10–15</td>
</tr>
<tr>
<td>1</td>
<td>13–25</td>
</tr>
<tr>
<td>2</td>
<td>17–35</td>
</tr>
<tr>
<td>3</td>
<td>25–45</td>
</tr>
<tr>
<td>4</td>
<td>60–65</td>
</tr>
</tbody>
</table>


**Abbreviation:** PL, pregnancy loss.

### RECOMMENDED SCREENING TESTS

#### Genetic Evaluation

**Parental Karyotype**

Three to four percent of couples with RPL have one parent with balanced translocation, or less commonly a chromosome inversion. Although these couples experience increased reproductive loss rates, most will have successful pregnancies without intervention [13]. Available intervention includes preimplantation genetic screening (PGS), and donor gametes.

**POC Karyotype**

Management of women with more than or equal to two RPLs should also be based on genetic evaluation of POCs (Figure 16.1) [14]. Most sporadic pregnancy losses in the first trimester result from random numeric chromosomal errors. When tissue of anembryonic pregnancies <9 weeks obtained by chorionic villus sampling (CVS) and manual vacuum aspiration (MVA) is analyzed, about 72% of cases revealed aneuploidy, with trisomy being most common [15]. This novel approach in the management of RPL includes CVS before evacuation. When compared with POCs, CVS is more successful in achieving a cytogenetic result, as omitting culturing of cells eliminates the potential for maternal contamination [16]. Additionally, miscarriage associated with aneuploidy increases with increasing maternal age; up to 80% in those over 35 years of age [17]. Aneuploidy is present in >50% of embryos tested preimplantation in women...
with RPL. More than 50% of aneuploidies are trisomies; most common single aneuploidy is 45XO. 46XX karyotype is often associated with maternal cell contamination, so that caution is necessary; microsatellite analysis decreases this confusion. This test can identify probable etiology and decrease further workup; this provides the couple with explanation and has been shown to decrease self-blame [18].

### Anatomic Evaluation

A maternal uterine study (e.g., three-dimensional [3D] or two-dimensional [2D] sonohysterogram [days 8–10 of follicular phase], hysterosalpingogram [HSG], hysteroscopy, or magnetic resonance imaging [MRI]) is recommended. Ten to fifteen percent of women with RPL have uterine anomalies. The most common congenital anomaly associated with RPL is septate uterus (spontaneous abortion [SAB] rate is high, about 65%) [19], followed by didelphys and bicornuate. Arcuate uterus has not been consistently associated with RPL. Uterine synechiae (Asherman's syndrome) and diethylstilbestrol (DES) exposure are associated with RPL. Myomata have been associated with decreased implantation rates in the in vitro fertilization (IVF) literature [20], but have not been consistently associated with RPL. Available intervention is hysteroscopic resection of septum, synechia, or submucous myoma. Surgical correction of these and other uterine anomalies has not been studied in trials.

### Endocrine Evaluation

Endocrine factors may contribute to 8%–12% of RPL. Basic evaluation should consistent of diabetes screening (hemoglobin A1c [HgbA1c], fasting glucose, glucose tolerance testing), screening for thyroid dysfunction (thyroid-stimulating
hormone [TSH] and free thyroxine [T4]); prolactin levels only if clinically suspicious.

**Immunologic Evaluation**

Three to fifteen percent of women with RPL have APAs. Tests should be positive twice, ≥12 weeks apart. See Chapter 26 in Maternal-Fetal Evidence Based Guidelines for tests (lupus anticoagulant, anticardiolipin antibodies [ACAs], and β-2 glycoprotein I) and effective intervention (e.g., low-dose aspirin and prophylactic heparin).

**TESTS THAT CANNOT BE ROUTINELY RECOMMENDED**

**Products of Conception**

- POC molecular genetic abnormalities (e.g., X-chromosome inactivation)
  
  Commercially available tests are not widely available for this testing.

**Mother**

**Endocrine**

- Endometrial biopsy or progesterone levels
  
  **Hypothesis:** The corpus luteum fails to make enough progesterone to sustain early decidua for placentation (LPD). It is normal to have at least two consecutive out-of-phase (≥2 days discrepancy) biopsies (diagnosis of LPD) on endometrial histology (late luteal phase—day 25 or 26—after presumed ovulation) in 50% of menstrual cycles; there is also high interobserver variation on interpretation of endometrial biopsies. There is insufficient evidence that intervention (such as progesterone supplementation—17P, micronized progesterone tablets 100 mg by mouth twice a day [po bid] or Crinone cream [8%] one application per vagina daily [beginning 2 days after ovulation until 10 weeks’ gestation or menses]) improves outcomes specifically in women with RPL and LPD [21,22]. 17P is efficacious in improving pregnancy outcomes in women with IVF, and in decreasing PTB in women with prior PTB (see Chapter 17).
  
  - Thyroid antibodies
    No consistent association and no intervention studies.

**Immunologic**

- Alloimmune tests (includes fecal occult blood [FOB])
  
  No consistent association and no efficacious intervention—see below.

- Antinuclear antibody (ANA)
  
  No consistent association and no intervention studies.

**Both**

**Thrombophilic**

Inherited thrombophilias (factor V Leiden [FVL], prothrombin G20210A gene mutation [PTM], antithrombin III deficiency, protein S, and protein C deficiencies) have not been consistently associated with RPL in the best designed prospective studies. However, a higher frequency of FVL carrier state has recently been shown in women with early RPL [23]. Second-trimester PL has been associated with thrombophilic mutations. There is insufficient evidence regarding any interventions in women with PL and thrombophilias (see Chapter 27 in Maternal-Fetal Evidence Based Guidelines).

**Infectious**

No infectious agent has been proven to cause RPL. Listeria, Toxoplasma gondii, and many viruses have been associated with sporadic early PL. Chlamydia, mycoplasma/ureaplasma...
(proposed diagnosis with endometrial biopsy, with treatment of woman and partner with either doxycycline 100 mg po bid or ciprofloxacin 250 mg po bid), and bacterial vaginosis are associated with sporadic PL, not RPL.

**MANAGEMENT**

**Prevention**

Optimize preconception medical care of all maternal diseases.

**Preconception Care**

Informative and sympathetic counseling should be provided. Workup is best done preconception (Table 16.4). When workup is positive, counsel regarding specific association. If workup is negative (>50% of couples), counseling should include the fact that 60%–70% of couples with unexplained RPL have successful pregnancies in the next gestation (Table 16.3). This percentage decreases to 40%–50% in women ≥40 years old. Offer all women with RPL a support group. UNITE provides the opportunity for emotional support, through parent-to-parent sharing on issues related to grieving. The groups are guided by trained facilitators, but the meetings are not group therapy sessions. UNITE can assist in referral if additional professional support is needed [24].

**Prenatal Care**

See subsection “Preconception Care.”

**Therapy (Specific for Workup)**

**Abnormal Parental Chromosomes**

Offer genetic counseling, prenatal diagnosis, and gamete donation. PGS with IVF is not supported by randomized controlled trials (RCTs) and should therefore not be recommended [12].

**Abnormal Uterine Cavity**

Septum, synechia, and/or submucous myomata can be resected hysteroscopically, but there are no trials regarding this intervention. There is a high likelihood of successful pregnancy in women with un repaired septa, so that some suggest against surgical repair in nulligravid women with uterine septa [12]. Repair of the bicornuate or unicornuate uterus is also usually not suggested in these women, as outcomes are usually good without repair, whereas surgical correction is associated with higher risk of complications. Consider referral to reproductive endocrinology specialist if necessary. There is unfortunately insufficient evidence to give a recommendation for women with fibroids and RPL.

**Medical Condition**

**If a medical condition is identified** (e.g., DM, thyroid disease), treat as indicated (see specific chapters in Maternal-Fetal Evidence Based Guidelines). Metformin has not been shown to reduce the risk of miscarriage in women with RPL and polycystic ovary syndrome [12].

**Antiphospholipid Syndrome**

Heparin and aspirin (see Chapter 26 in Maternal-Fetal Evidence Based Guidelines)

Therapy is usually begun once fetal viability is established. Low-dose aspirin is usually about 75–100 mg daily. For prophylactic unfractionated heparin (UFH): 5,000–7,500 U first trimester, 7,500–10,000 U second trimester, and 10,000 U third-trimester SQ q12h. Heparin used in positive trials against placebo was UFH, and even if low-molecular-weight heparin (LMWH) is associated with fewer side effects in nonpregnant adults, RCTs of LMWH in pregnancy for RPL in antiphospholipid syndrome (APS) women have not shown benefit compared with low-dose aspirin [25,26]. Nonetheless, two small RCTs have directly compared LMWH to UFH, and despite the small number of patients recruited, effectiveness of LMWH appears comparable with that of UFH [27,28], and therefore either UGH or LMWH can probably be used (see Chapter 26 and Table 26.3). UFH combined with aspirin is associated with a significant reduction in PL compared with aspirin alone (risk ratio [RR] 0.46, 95% confidence interval [CI] 0.29–0.71; three trials, n = 140). There is no advantage in high-dose, over low-dose, UFH (one trial, n = 50).

LMWH combined with aspirin compared with aspirin does not significantly reduce PL (odds ratio [OR] 0.70, 95% CI 0.34–1.45; 5 trials, n = 598) [29]. For prophylactic LMWH either enoxaparin (Lovenox) 30–40 mg subcutaneously every 12 hours (SQ q12h) or dalteparin (Fragmin) 5000 U SQ q12h were used in RCTs, and clinicians may adjust prophylaxis in high-risk cases to heparin (anti-Xa) level range of 0.2–0.3.

Three trials of aspirin alone (n = 135) show no significant reduction in PL (RR 1.05, 95% CI 0.66–1.68).

Prednisone and aspirin (three trials, n = 286) result in a significant increase in prematurity when compared with placebo, aspirin, and heparin combined with aspirin, and an increase in gestational diabetes, but no significant benefit.

**Intravenous immunoglobulin (IVIG)** ± UFH and aspirin (two trials, n = 58) is associated with an increased risk of PL or premature birth when compared with UFH or LMWH combined with aspirin (RR 2.51, 95% CI 1.27–4.95). When compared with prednisone and aspirin, IVIG (one trial, n = 82) is not significantly different in outcomes [30].

In summary, therapy with UFH and aspirin can be recommended to the woman with RPL and APS, as it reduces the chance of PL by 54% compared with aspirin alone.

In women with RPL and APA, available guidelines recommend a combined therapy with low-dose aspirin and prophylactic doses of heparin, although the available RCTs include heterogeneous groups of patients [31].

**Inherited Thrombophilia**

As stated above, a workup for inherited thrombophilia is not indicated. If positive inherited thrombophilia is incidentally or previously identified, no intervention has been consistently shown to improve outcomes. Please also refer to Chapter 15, and to Chapter 27 in Maternal-Fetal Evidence Based Guidelines.

**Negative Workup**

**Progesterone**. In women who had more than or equal to three consecutive miscarriages, progesterone treatment is associated with a statistically significant reduction in miscarriage rate (OR 0.39, 95% CI 0.21–0.72) compared with placebo or no treatment in four small trials [32–35]. No statistically significant differences, probably because of small numbers, were found between the route of administration of progesterone (oral, intramuscular, and vaginal) and placebo or no treatment [32–35], so the best route and dose of progesterone for prevention of RPL are still unknown. In summary, progesterone treatment for women
with RPL may be warranted given the reduced rates of miscarriage in the treatment group and the finding of no statistically significant difference between treatment and control groups in rates of adverse effects suffered by either mother or baby [32].

Supportive care. Consider intensive supportive early prenatal care, focusing on antenatal counseling and psychological support. There are no properly controlled trials to assess the effect of this intervention. Three studies (not RCTs) showed improved outcome versus standard or no prenatal care [10,36,37].

Human chorionic gonadotropin. There is not enough evidence to evaluate the use of hCG during pregnancy in order to prevent miscarriage in women with a history of unexplained recurrent spontaneous miscarriage because the trials are small, and have significant (especially two studies) limitations [38–42]. hCG is associated with a 74% reduction in both early (4.1% vs. 8.8%) and late (1.1% vs. 2.3%) miscarriages, but no other effects on other perinatal outcomes [47].

UFH. There is insufficient evidence to assess the effect of UFH in women with unexplained early PLs.

LMWH. There are several RCTs that show no effect of LMWH on prevention of PL in women with unexplained RPL. Compared with low-dose aspirin, enoxaparin (a LMWH) was associated with similar live birth rates, respectively 82% and 84% (RR 0.97, 95% CI 0.81–1.16), in 107 women with consecutive recurrent miscarriage (more than or equal to three first-trimester miscarriages or more than or equal to two consecutive second-trimester miscarriages) without any apparent cause and no hereditary thrombophilia [46]. In 340 women with more than or equal to three unexplained first-trimester RPLs, enoxaparin 20 mg daily from fetal viability until 34 weeks was associated with a slight nonsignificant reduction in both early (4.1% vs. 8.8%) and late (1.1% vs. 2.3%) miscarriages, but no other effects on other perinatal outcomes [47]. Additionally, a more recent multicenter trial with a minimization randomization scheme confirmed similar live birth rates (86% vs. 86.7%) in the intervention (LMWH) and control groups, respectively [48].

No reduction in PL was observed when LMWH and low-dose aspirin were used in combination to treat 297 women with idiopathic recurrent miscarriage (22% in pharmacologic intervention group vs. 20% in surveillance group) [49]. In women with 364 unexplained RPLs, nadroparin, and low-dose aspirin were associated with similar outcomes compared with either low-dose aspirin alone or to placebo [45].

Vitamins

There is no specific adequate trial on multivitamin supplementation of any kind for women with prior RPL.

Diethylstilbestrol

DES should not be used in pregnancy for any indication. Data are mostly from studies of women without risk factors, women with “threatened abortion” in the current pregnancy, or diabetics (most women with RPL). DES use in pregnancy is significantly associated with several harmful consequences for both mother and baby. DES given in the first trimester [50] leads to a 37% increased rate of miscarriage and 61% increased rate of PTB [51–58]. There is also a 48% increase in the numbers of babies weighing less than 2500 g. Stillbirth and neonatal death are not influenced by the intervention (DES) as compared with the control group. Preeclampsia is similar in the two groups. Exposed female offsprings have a nonsignificant trend toward more cancer of the genital tract and cancer other than of the genital tract. Primary infertility, adenosis of the vagina/cervix in female offsprings, and testicular abnormality in male offsprings are significantly higher in those exposed to DES before birth.

The vast use in the 1950s to 1970s of a medication with no benefit proven by evidence-based medicine is the best example of the importance of using data from trials and meta-analyses to guide effective practice.

Immunotherapy

The various forms of immunotherapy did not show significant differences between treatment and control groups in terms of subsequent live births [59]:

- Paternal cell immunization (12 trials, 641 women), OR 1.23, 95% CI 0.89–1.70 [60–70].
- Third-party donor cell immunization (3 trials, 156 women), OR 1.39, 95% CI 0.68–2.82 [50,61,63].
- Trophoblast membrane infusion (one trial, 37 women), OR 0.40, 95% CI 0.11–1.45 [71].
- IVIG (eight trials, 303 women), OR 0.98, 95% CI 0.61–1.58 [72–79]. More recently, another trial confirmed these results [80].

Immunoablation using viable mononuclear cells carries the risk of any blood transfusion such as hepatitis B virus or human immunodeficiency virus (HIV). Reactions have been uncommon but include soreness and redness at the injection site, fever, maternal platelet alloimmunization, blood group sensitization, and one cutaneous graft-versus-host-like reaction. Women who have received lymphocyte immune therapy may have a higher incidence of subsequent miscarriage than women who did not receive such cellular products [54]. The Director of the Office of Therapeutics Research and Review, U.S. Food and Drug Administration (FDA), sent a letter on January 30, 2002, to physicians believed to be using lymphocyte immune therapy to prevent miscarriages. He informed them that the injectable products used in lymphocyte immune therapy do not have the required FDA approval and are considered investigational new drugs that pose several significant safety concerns. Administration of such cells or cellular products in humans can only be performed in the United States as part of clinical investigations, and then only if there is an investigational new drug (IND) application in effect. IVIG therapy is expensive and in relatively short supply.

Immunotherapies should not be offered as treatment for unexplained RPL. Women should be spared the pain and grief associated with false expectations that an ineffective treatment might work.

**ANTEPARTUM TESTING**

No specific testing indicated.

**DELIVERY**

No specific precaution.
ANESTHESIA
No specific precaution.

POSTPARTUM/BREAST-FEEDING
No specific precaution.

FUTURE
Effective treatment of an alleged alloimmune cause of recurrent miscarriage awaits more complete knowledge of the underlying pathophysiology. A specific assay to diagnose immune-mediated early PL and a reliable method to determine which patients might benefit from manipulation of the maternal immune system are urgently needed. It is not presently known exactly how many REPLs are the results of anembryonic or chromosomally abnormal conceptuses, anatomic or structural abnormalities, and how many are embryonic or fetal deaths. It is likely that some unexplained early losses are due to as yet undefined subchromosomal genetic abnormalities impairing early development of the conceptus. New molecular techniques should be directed at understanding the factors responsible for successful pregnancy as well as PL.

REFERENCES
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34. Swyer GM, Daley D. Progesterone implantation in habitual abortion. Br Med J. 1953;1(4819):1073–1077. [RCT, n = 47 (out of 113 total). Women having had two or more consecutive miscarriages before 12 weeks’ gestation. 6 x 25 mg progesterone pellets inserted within the uterine muscle either: (i) as soon as pregnancy was confirmed or (ii) not later than 10th week of gestation, or (iii) not later than the earliest previous miscarriage. Placebo: no but had a no-treatment control group.]


39. Harrison RF. Treatment of habitual abortion with human chorionic gonadotropin: Results of open and placebo-controlled studies. Eur J Obstet Gynecol Reprod Biol. 1985;20(3):159–168. [RCT, n = spontaneous miscarriage of three previous consecutive pregnancies without evidence of a cause, normal investigative profile included: Chromosomal analysis of both partners; no systemic disease; normal bacteriological investigations of semen and cervical secretions; normal HSG, normal serum FSH, LH, estradiol, and prolactin; normal or low progesterone. Initially 10,000 IU hCG by IM injection followed by 5000 IU twice weekly up to 12 weeks, followed by 5000 IU weekly until 20 weeks.]


42. Svigos J. Preliminary experience with the use of human chorionic gonadotrophin therapy in women with repeated abortion. Clin Reprod Fertil. 1982;1(2):131–135. [RCT. Two unexplained (normal genital tract and chromosomes) miscarriages. Treatment group managed in one of two ways according to serum progesterone. A low progesterone precipitated compliance with the intended treatment. A normal progesterone precipitated no treatment (i.e., managed as a control). Groups analyzed on an intention-to-treat basis. 9000 IU IM 3 per week from 6 to 7 weeks until 12 weeks vs. no treatment.]


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56. Reid D. The use of hormones in the management of pregnancy in diabetics. Lancet. 1953;2:833–836. [RCT, n = 147 (High-risk diabetic women). Incremental doses of 50–200 mg DES daily from about 16 weeks to term. Ethisterone, 25 mg/day from 16 weeks, incrementally to 250 mg/day at 32 weeks to term. Ethisterone was given incrementally. Graduated dosing with stilbestrol from 50 mg. 5 weeks or less to 200 mg at 32 weeks or more. Ethisterone from 25 mg/day at 19 weeks or less to 250 mg at 32 weeks or more vs. placebo.]


Preterm birth prevention in asymptomatic women

Anju Suhag

KEY POINTS

• Gestational age (GA) determination is of utmost importance in prevention of preterm birth (PTB) and management of presumed threatened PTB.

• PTB is defined as birth between 20 0/7 and 36 6/7 weeks. It is the number one cause of perinatal morbidity and mortality, and these complications are inversely proportional to GA at birth. Over 1 million babies die of the consequences of PTB every year in the world, 1 every 30 seconds.

• In women with ≥3 prior spontaneous preterm births (SPTBs) or second-trimester pregnancy loss (STLs), history-indicated cerclage (PEIC) is associated with a significant decrease in PTB.

• In women with asymptomatic group B streptococcus (GBS) bacteriuria of >100,000 bacteria/mL, appropriate antibiotics are associated with a significant decrease in PTB.

• In women with asymptomatic cervical dilatation ≥1 cm before 24 weeks, physical examination–indicated cerclage (PEIC) is associated with a significant decrease in PTB.

• Secondary prevention of PTB is based on identification of risk factors for PTB, especially obstetric–gynecological (ob-gyn) history, maternal lifestyle, and prepregnancy weight (Table 17).

• Primary prevention strategies for PTB aimed at the general population include family planning, avoidance of lifestyle risks, and proper nutrition. A reproductive-age woman should avoid (as feasible) extremes of age, of interpregnancy interval (18–23 months is optimal interval between last delivery and next conception), uterine evacuation of pregnancy without ripening, multiple gestations, illegal drugs (e.g., cocaine), physical abuse, sexually transmitted infections (STIs), poverty, poor education, and prepregnancy weight (Table 17).

• Screening with transvaginal ultrasound (TVU) cervical length (CL) at 18–23 6/7 weeks should be offered to all singleton gestations. In singleton gestations without a prior spontaneous preterm birth (SPTB), if CL ≤25 mm develops, cervical dilatation (CI) (formerly called incompetence): recurrent painless dilatation leading to STLs [4]. This chapter reviews evidence-based guidelines for women without symptoms related to PTB. The following two chapters deal with women with symptoms of PTB: first preterm labor (PTL) (Chapter 18), then preterm premature rupture of membranes (PPROM) (Chapter 19).

BACKGROUND

Prevention of PTB is the number one issue in pregnancy, as being born preterm is the number one cause of neonatal mortality and the second leading cause of all under-five childhood mortality in many developed countries, including the United States [1,2]. This chapter reviews evidence-based guidelines for prevention of PTB and management of presumed threatened PTB (see Chapter 4 for best GA determination criteria).

Definitions regarding prematurity vary in different publications, but the ones most commonly accepted and used in trials are the following:

• Preterm birth (PTB): birth between 20 0/7 and 36 6/7 weeks [3]

• Very early PTB: birth between 20 0/7 and 23 6/7 weeks

• Early PTB: birth between 24 0/7 and 33 6/7 weeks

• Late PTB: birth between 34 0/7 and 36 6/7 weeks

• Pregnancy loss (PL): spontaneous loss of pregnancy from conception to <20 weeks. The term spontaneous abortion is equivalent but should be avoided since women associate negative feeling with this term. Miscarriage is a lay term for PL (see also Chapters 15 and 16).

• Second-trimester PL (aka STL): birth between 14 0/7 and 19 6/7 weeks.

Cervical insufficiency (CI): recurrent painless dilatation leading to STLs [4]. A better definition is a CL <25 mm before 24 weeks in women with singleton gestations and prior SPTB <37 weeks.

• Preterm labor (PTL): uterine contractions (≥4/20 minutes or ≥8/hours) and documented cervical change with...
intact membranes at 20–36 6/7 weeks. A better definition is uterine contractions (≥4/20 minutes or ≥8/hour) with TVU CL <20 mm, or 20–29 mm with positive fetal fibronectin (fFN) at 20–36 6/7 weeks (see Chapter 18).

Premature preterm rupture of membranes (PPROM): vaginal pooling, nitrazine, and/or ferning at 16–36 6/7 weeks (see Chapter 19).

• Early PPROM: PPROM between 24 and 33 6/7 weeks
• Very early PPROM: PPROM between 24 and 33 6/7 weeks

EPIDEMIOLOGY/INCIDENCE
Incidence of PTB <37 weeks varies between about 5% and 18% in different countries [5]. The rate of PTB in the United States steadily increased from 9.4% in 1981 and peaked in 2006 at 12.8%. Since then, the rate of PTB in the United States dropped to 9.6% in 2014 (25% decline) [6,7]. The incidence of PTB <32 weeks remains at about 2% in United States; and ≤1% in most other high-income countries. Over 1 million babies die of the consequences of PTB every year in the world, 1 every 30 seconds [1]. The high incidence of PTB in many high-income countries may be due to assisted reproductive technology (ART)-related multiple gestations, older and sicker mothers, earlier GA of registered births and neonatal improvements, better and earlier timing of births (related to ultrasound), worsening socioeconomic factors, and other factors. In addition to the use of progesterone and cerclage in women at high risk for PTB, there are several other demographic factors which lead to reducing national PTB rate. This decline in PTB in the United States appears to be related also to reduced teen birth rate, lower rate of higher order multiple births, smoking cessation counseling and bans, and improved institution/national policies of preventing non-medically indicated PTB <39 weeks [8].

GENETICS
While a genetic predisposition in certain ethnic groups and families has been reported, no clinical genetic studies are yet recommended for prediction/prevention of PTB due to insufficient evidence [9].

ETIOLOGY/BASIC PATHOPHYSIOLOGY
Just like cardiac disease, PTB is a final common manifestation of multifactorial, complex etiology. Several processes leading to PTB are shown in Figure 17.1.

CLASSIFICATION
PTB can be spontaneous, and follow PTL (50%; Chapter 18), PPROM (30%; Chapter 19), or, rarely, cervical insufficiency; or be iatrogenic (20%). Cervical insufficiency may comprise about 1% of spontaneous PTB (SPTB) and/or STLs (see below in the section “Prediction and Prevention of PTB”). Cervical insufficiency represents one extreme of SPTB, as PTB is continuous.
PTB is the final common pathway of many associated possible etiologies. Cl; cervical insufficiency; SPONT.PTL, spontaneous preterm labor; PPROM, preterm premature rupture of membranes; PTB, preterm birth.

**RISK FACTORS/ASSOCIATIONS**

Most women who have a PTB have identifiable risk factors. Risk factors for SPTB (presenting as PTL, PPROM, or cervical insufficiency) are similar (Table 17.1). All these risk factors should be reviewed in details with every pregnant woman at the first prenatal visit. The vast majority of U.S. pregnant women (96% on in one study) has at least one risk factor for PTB [10]. For many of these risks, there are interventions associated with prevention of PTB.

**COMPLICATIONS**

PTB is the number one cause of perinatal mortality. Seventy-five percent of perinatal mortality occurs in preterm babies; more than two-third of perinatal mortality (60% of total) occurs in infants aged <32 weeks. Mortality and morbidities are inversely associated with GA at birth (Table 17.2) [11]. Morbidities include compared with (RDS), bronchopulmonary dysplasia (BPD), intraventricular hemorrhage (IVH), necrotizing enterocolitis (NEC), sepsis, apnea and retinopathy of prematurity [12]. The long-term neonatal morbidities from PTB are inversely associated with GA at birth (Table 17.2) [11].

**PREDICTION AND PREVENTION OF PTB**

A screening test aimed at prediction of PTB is only beneficial if an intervention (prevention) reduces the outcome once the screening test is positive. Several predictive strategies have poor sensitivity and specificity.

Prevention is preferable, rather than treatment once symptoms have been identified. Prevention can be divided into three categories (Table 17.3). Primary and secondary prevention are preferable, rather than tertiary prevention of symptomatic women with PTL or PPROM. This chapter refers to prevention of SPTB mostly in singleton gestations, unless otherwise specified. Prevention of PTB in multiple gestations is also discussed in Chapter 44 in *Maternal-Fetal Evidence Based Guidelines*.

For women at risk for iatrogenic (indicated) PTB, one should aim to keep the pregnant woman as healthy as a nonpregnant adult. Appropriate prevention and therapy of any maternal medical or fetal/congenital anomaly disorder is paramount, as is appropriate prevention and therapy for preeclampsia and fetal growth restriction (FGR) (please refer specific chapters in *Maternal-Fetal Evidence Based Guidelines*) [1].

All pre-pregnancy evaluations of the cervix (e.g., hysterosalpingogram, no. 8 Hegar dilator passage, and catheter traction test) aimed at screening for cervical insufficiency have either been inadequately studied or been shown not to be sufficiently predictive and therefore useful in a prevention program (no trial ever reported).

An accurate history should be taken regarding risk factors for PTB, especially ob-gyn history, maternal lifestyle, and prepregnancy weight (Table 17.1). A detailed history should include whether the prior PTB was spontaneous or indicated (iatrogenic). Spontaneous PTB is defined as PTB prior to 37 weeks due to PTL, PPROM, advanced cervical dilation (ACD) or cervical insufficiency. It is also important to document the sequence of events (antepartum) leading to a PTB, as this could help counsel these women about their risk of recurrent PTB and appropriate PTB prevention strategies could be utilized. For example, women with a history of ACD are at an increased risk of having cervical shortening (50% vs. 14.6% vs. 15.6%, p<0.01), recurrent PTB (55.2% vs. 27.2% vs. 32.2%, p<0.01) and a lower GA at delivery (34.0 vs. 37.2 vs. 37.0 weeks, p<0.01) in a subsequent pregnancy compared with women with prior PTB associated PPROM or PTL, respectively [15].

A Creasy’s score or other similar history-based systems to predict PTB have been associated with a low (10%–30%) positive predictive value (PPV) for PTB, and not clinically useful given negative intervention trials (see below, in the last paragraph in this section and “Intervention: Weekly manual examinations, education”). There are no trials on risk-scoring systems for predicting PTB [16]. There is a need for prospective studies that evaluate the use of a risk-screening tool designed to predict PTB (in combination with appropriate consequent interventions) to prevent PTB, including qualitative and/or quantitative evaluation of their impact on women's well-being.

There are four main risk factors for which there are effective interventions for prevention of PTB: smoking; short TVU CL; prior SPTB; and asymptomatic bacteriuria (Table 17.4). A TVU CL screening at 18–24 weeks is indicated in all women with a singleton gestation (see "Risk: Short Cervix on Ultrasound" in the section “Secondary Prevention”).

Current evidence does not support the use of home uterine activity monitoring (HUAM) [17] or bacterial vaginosis (BV) screening [18] in asymptomatic low-risk women. There are insufficient data to support the use of salivary estriol or fFN in asymptomatic women (even if fFN is one of the best predictive screening tests). Cytokines, matrix metalloproteinases, corticotropin-releasing hormone (CRH), salivary estriol, relaxin, human chorionic gonadotropin (HCG), prothrombin, fetal DNA, and many other tests remain research tools for prediction of PTB and are not yet clinically beneficial, given lack of intervention trials based on these screening tests.

**Primary Prevention**

Primary prevention includes prevention strategies aimed at all asymptomatic pregnant women at risk for PTB (i.e., aimed at all pregnant women). Unfortunately, many primary prevention interventions have been so far either insufficiently studied or found not to be effective.
There are no trials to assess interventions. Avoiding extremes of age, avoiding short interpregnancy interval <6 months (18–23 months is optimal interval between last delivery and next conception) [19], and reducing the rate of multiple gestations (through ART improvements) seem self-evident for efficacy in preventing PTB when feasible. Evidence suggests that risks of PTB, low birth weight (LBW), and small-for-gestational-age (SGA) infants are minimized when interpregnancy intervals are between 18–23 months [19,20]. For appropriate interpregnancy interval, physicians should pay close attention to plans for postpartum contraception. Improvements

<table>
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<tr>
<th>Table 17.2</th>
<th>Preterm Neonatal Morbidity and Mortality</th>
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<tr>
<td>Outcome</td>
<td>Delivery Gestational Age (Weeks)</td>
</tr>
<tr>
<td></td>
<td>All</td>
</tr>
<tr>
<td>Death</td>
<td>1.4</td>
</tr>
<tr>
<td>Major morbidity&lt;sup&gt;a&lt;/sup&gt;</td>
<td>7.9</td>
</tr>
<tr>
<td>Minor morbidity&lt;sup&gt;b&lt;/sup&gt;</td>
<td>37.6</td>
</tr>
<tr>
<td>Survival without any of above morbidities</td>
<td>53.1</td>
</tr>
</tbody>
</table>


Note: Data presented as %; N = 8334.

<sup>a</sup>Includes persistent pulmonary hypertension, intraventricular hemorrhage grade 3/4, seizures, hypoxic–ischemic encephalopathy, necrotizing enterocolitis stage II/III, and bronchopulmonary dysplasia.

<sup>b</sup>Includes intraventricular hemorrhage grade 1/2, necrotizing enterocolitis stage 1, respiratory distress syndrome, hyperbilirubinemia requiring treatment, and hypotension requiring treatment.

<table>
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<tr>
<th>Table 17.3</th>
<th>Definition of the Different Categories of Prevention of PTB</th>
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<tr>
<td>Definition</td>
<td>Examples</td>
</tr>
<tr>
<td>Primary prevention</td>
<td>Prevention strategies aimed at all asymptomatic pregnant women at risk for PTB (i.e., aimed at all pregnant women). Encourage optimal (18–23 months) interpregnancy interval. Limit higher order multiple births (ART). Obtain comprehensive OB history and offer preventive interventions (progesterone, cerclage, CL screening) to appropriate candidates. Smoking cessation programs.</td>
</tr>
<tr>
<td>Secondary prevention</td>
<td>Prevention strategies aimed at identifying asymptomatic women at high risk for PTB through screening. Screen for predictive risk factor (prediction) in asymptomatic women and avoid/treat (preventive intervention). Obtain comprehensive OB history and offer preventive interventions (progesterone, cerclage, CL screening) to appropriate candidates. Smoking cessation programs.</td>
</tr>
<tr>
<td>Tertiary prevention</td>
<td>Prevention strategies aimed at women with active symptoms of PTB.</td>
</tr>
</tbody>
</table>

Abbreviations: OB, obstetrical; PTB, preterm birth; PTL, preterm labor; PPROM, preterm premature rupture of membranes; ART, assisted reproduction technologies; CL, cervical length.

<table>
<thead>
<tr>
<th>Table 17.4</th>
<th>Selected Effective Interventions for Prevention of PTB</th>
</tr>
</thead>
</table>
| Avoid (as feasible) | • Extremes of age  
| | • Interpregnancy interval <6 months  
| | • Induced termination of pregnancy without ripening  
| | • Multiple gestation  
| | • Illegal drugs (e.g., cocaine)  
| | • Physical abuse  
| | • STIs  
| | • Poverty  
| | • Poor education  
| | • Premature pregnancy weight of <50 kg (<120 lb).  
| Risk | Intervention |
| Smoking | Smoking cessation programs |
| Short CL ≤ 20 mm (screen every singleton with TVU CL at 18–23/6/7 weeks)<sup>a</sup> | Vaginal progesterone (e.g., 200 mg daily) |
| Prior spontaneous PTB (SPTB)<sup>b</sup> | 17-OH progesterone caproate, moderate (≥3 meals/week) fish intake |
| Prior SPTB and CL <25 mm between 16 and 23 6/7 weeks<sup>c</sup> | Ultrasound-indicated cerclage |
| Prior ≥3 PTB/STL<sup>d</sup> | History-indicated cerclage |
| Asymptomatic bacteriuria | Appropriate antibiotics |

Abbreviations: STI, sexually transmitted infections; CL, cervical length; TVU, transvaginal ultrasound; SPTB, spontaneous preterm birth; STL, second trimester loss.

<sup>a</sup>For singleton gestations without prior PTB.

<sup>b</sup>For singleton gestations without prior PTB.

<sup>c</sup>For singleton gestations.

Preconception/Early Pregnancy

**Family planning.** There are no trials to assess interventions. Avoiding extremes of age, avoiding short interpregnancy interval <6 months (18–23 months is optimal interval between last delivery and next conception) [19], and reducing the rate of multiple gestations (through ART improvements) seem self-evident for efficacy in preventing PTB when feasible. Evidence suggests that risks of PTB, low birth weight (LBW), and small-for-gestational-age (SGA) infants are minimized when interpregnancy intervals are between 18–23 months [19,20]. For appropriate interpregnancy interval, physicians should pay close attention to plans for postpartum contraception. Improvements
in postpartum contraceptive use have been associated with decreased incidence in PTB. For every month of contraception coverage, the odds of PTB <37 weeks decrease by 1.1% (odds ratio [OR] 0.98, 95% confidence interval [CI] 0.98–0.99) [21].

Induced termination of pregnancy (TOP) is associated with a higher risk of PTB, even if just one early TOP is performed [22]. Recent meta-analyses showed that dilation and curettage (D&C) compared with no D&C increase the risk of PTB <37 weeks (OR 1.29, 95% CI 1.17–1.42) [23,24]. Women who had even just one uterine evacuation for either spontaneous or induced abortion have a higher risk of PTB (OR 1.44, 95% CI 1.09–1.90) [24]. The risk of PTB <37 weeks is noted to be even higher in women with history of multiple D&Cs (OR 1.74, 95% CI 1.10–2.76) [23]. Avoiding TOPs as feasible with effective contraception seems to be an effective preventive strategy to avoid PTB. If TOP is unavoidable, the D&C should be performed with preoperative cervical ripening (e.g., misoprostol, mifepristone, or laminaria), as this is associated with no or very low risk for PTB [22].

Avoidance of lifestyle risks. There are no trials to assess interventions. Avoiding illegal drugs (e.g., cocaine and amphetamines), physical abuse, and STIs (chlamydia, gonorrhea, syphilis, human immunodeficiency virus [HIV], etc.) seem self-evident for efficacy in preventing PTB [25]. Poverty, poor education, lack of women’s empowerment, and several other social stressors (Table 17.1) are all linked profoundly to an increased risk of PTB and should be avoided as possible, especially through political and social changes. Workplace conditions are only weakly related or are not related to adverse pregnancy outcomes. In the third trimester, >42 hours of work/week, >6 hours of standing/day, and pesticide use are associated with PTB [26]. There are no trials on modifying other potential risks, such as physically demanding job, prolonged standing, night work, or others.

Proper nutrition, weight gain. A prepregnancy weight of <50 kg (<120 lb) should be avoided. Recent meta-analysis of 17 randomized controlled trials (RCTs) showed benefit of nutrition education to increase energy and protein intake in decreasing risk of PTB (two trials, 449 women) (relative risk [RR] 0.46, 95% CI 0.21–0.98, low-quality evidence), and LBW (one trial, 300 women) (RR 0.04, 95% CI 0.01–0.14) among undernourished women [27]. A normal BMI at the start of pregnancy, not dieting but instead a balanced diet during pregnancy achieving weight gain of >5 kg by 30 weeks for underweight and normal-weight women, can certainly help to avoid PTB.

The available evidence shows no benefit of balanced protein/energy supplementation on prevention of PTB [28]. High-protein diet (≥25% of total energy content) has no association with PTB prevention, and is not recommended in pregnancy because of increased risk of SGA [29] (see also Chapter 2).

Early Pregnancy

Fish intake. Shark, swordfish, king mackerel, or tilefish contain high levels of mercury and should be avoided or eaten infrequently (≤1/week) in pregnancy. Canned light tuna, salmon, pollock, grouper, mussels, scallops, shrimp, and catfish are common fishes low in mercury, and two portions (6 oz = 1 portion) of these per week can (and probably should) be safely eaten in pregnancy. Albacore (white) tuna has more mercury and should be consumed up to 6 oz/week. In general, for other fishes, smaller fishes have less mercury than larger ones. More information on fish and mercury intake in pregnancy is available at www.cfsan.fda.gov and www.epa.gov/ost/fish. There is limited evidence suggesting potential benefit of maternal fish intake on reduction of PTB. A recent prospective cohort study reported a significant reduction in risk of PTB, in women choosing a “prudent” or a “traditional” dietary pattern, characterized by, for example, vegetables, cooking oil, fruit, berry, olive oil, rice, water as beverage, whole grain cereals, yogurt, poultry, lean fish, and boiled potatoes, as well as low intake of processed meat products, white bread, and pizza/tacos [30].

Omega-3 fatty acids. Low-risk women without a prior PTB have a similar incidence of PTB <37 weeks when given 12 eggs with either 133 mg docosahexaenoic acid (DHA) or with 33 mg DHA per week starting at 24–28 weeks [31]. Supplementation has been shown to prolong pregnancy by 3–8 days in different populations. When all RCTs are examined, women allocated a marine oil supplement had a mean gestation that was 2.6 days longer than women allocated to placebo or no treatment. This was not reflected in a clear difference between the two groups in PTB <37 weeks, although women allocated marine oil did have a lower risk of PTB <37 weeks (RR 0.69, 95% CI 0.49–0.99). Birth weight was slightly greater (47 g) in infants born to women in the marine oil group compared with controls. However, there were no overall differences between the groups in the proportion of LBW or small-for-GA babies. There was no clear difference in the relative risk of preeclampsia between the two groups [32]. A large RCT found that DHA 800 mg supplementation before 21 weeks was associated with a decrease in PTB <34 weeks (RR 0.49, 95% CI 0.25–0.94) [33]. In contrast, in a recent meta-analysis of nine randomized trials including over 3800 asymptomatic women with singleton pregnancies and no prior PTB, women who received omega-3 had a similar rate of PTB <37 weeks as women who received either a placebo or no supplementation (77% vs. 91%; RR 0.90, 95% CI 0.72–1.11) [34]. The evidence suggests minimal or no benefit of omega-3 supplementation in reduction in PTB <37 weeks in asymptomatic low-risk singleton pregnancies.

PrimaCare vitamin supplement contains 150 mg of omega-3 fatty acids and has not been evaluated in a trial. The possible beneficial effects of omega-3 fatty acids to later fetal/neonatal/infant cognition remain not fully proven. For omega-3 supplementation in high-risk women, see section “Risk: Prior PTB.”

Probiotics. Women who report habitual intake of probiotic dairy foods (e.g., yogurt) have been associated with lower incidence of PTB in non-RCT [35]. Currently, there are insufficient data from RCTs to demonstrate any impact of probiotic foods supplementation in low-risk women on PTB and its complications [36].

Other nutritional changes. There are no trials with specific aim of prevention of PTB to evaluate other nutritional changes, such as vitamin supplementation (see Chapter 2), HCG, and anticytokine supplements. Prepregnancy weight <120 lb (<50 kg) is a very significant risk factor for PTB and should be avoided if possible. Suggested pregnancy weight gain in pregnancy is 25–35 lb for women with normal BMI, but there are no trials on proper prepregnancy weight or pregnancy weight gain.

Vitamin C. Vitamin C supplementation alone is not associated with a reduced risk of PTB [37].

Vitamin E. Vitamin E supplementation alone or in combination with other supplements is not associated with prevention of PTB [38].
Vitamins C and E. Maternal supplementation with vitamin C 1000 mg and vitamin E 400 mg daily from 9 to 16 weeks until delivery in nulliparous low-risk women is not associated with a reduction in PTB [39].

Vitamin D. Level II evidence suggests protective effect of vitamin D supplementation in women with low 25-hydroxyvitamin D concentration (at or prior to 20 weeks) on rate of PTB in low-risk women [40]. Routine vitamin D level assessment and/or vitamin D replacement or supplementation is not indicated in women at risk for PTB, until a well-designed RCT confirms above findings.

Calcium supplementation. Maternal supplementation of calcium versus placebo is not associated with a reduction in PTB [41].

Magnesium supplementation. There is insufficient high quality evidence to show that dietary magnesium supplementation during pregnancy is not beneficial. In the analysis of all trials, oral magnesium treatment before 25 weeks showed no significant effect of magnesium on perinatal mortality, SGA, preeclampsia and PTB, compared with placebo [42]. Of the ten trials included in the review, only two were judged to be of high quality, and showed no association of magnesium and pregnancy outcomes.

Progesterone. The mechanism of action of progesterone for prevention of PTB is poorly understood but probably involves an anti-inflammatory action. Safety for the fetus/neonate has not been yet proven with 100% certainty, but progesterone is known not to be a teratogen, and long-term detrimental effects up to 18 years of age have not been shown. The effect of progesterone supplementation should be evaluated according to different patient populations, and according to type of progesterone. Here we review progesterone for low-risk women (see below for other risk groups).

In a small RCT including women in active military duty with only a 3% rate of prior PTB and unknown CL, 17P 1000 mg IM weekly starting at 16–20 weeks was not associated with any effect on incidence of PTB or perinatal outcomes compared with placebo [43]. No RCT has evaluated the effect of vaginal progesterone in this population.

In summary, there is insufficient evidence to determine the impact on PTB of progesterone in singleton gestations with no prior PTB, with unknown or normal CL.

Secondary Prevention

Secondary prevention strategies involve screening for a predictive risk factor (prediction) in asymptomatic women, and avoiding it or treating it (preventive intervention) (Table 17.3).

Risk: Smoking

**Intervention: Smoking cessation programs.** It is estimated that 10%–15% of PTBs may be due to smoking. In the United States, about 23% of women start pregnancy as a smoker, and 11% continue to smoke throughout pregnancy [44]. Psychosocial interventions (e.g., counseling, health education, feedback, incentives, social support) to support women to stop smoking in pregnancy has been associated with an increase the proportion of women who stop smoking in late pregnancy, and reduce LBW (RR 0.82, 95% CI 0.71–0.94) and PTB (18% reduction, RR 0.82, 95% CI 0.70–0.96) [45]. The American College of Obstetricians and Gynecologists (ACOG) has recommended use of the five As—ask, advice, assess, assist, arrange—approach [46]. The most effective intervention for smoking cessation in pregnancy is social support and a reward component (23% decrease) [47,48]. If above approach is not successful, consider nicotine replacement therapy (NRT) (see Chapter 22 in Maternal-Fetal Evidence Based Guidelines).

**Intervention: Nicotine replacement therapy.** NRT is associated with a trend for benefit in reduction in the incidence of smoking [49–51] (see Chapter 22 in Maternal-Fetal Evidence Based Guidelines). A recent RCT showed that NRT was associated with a higher validated smoking cessation rate 1-month postrandomization; however, there was no difference between group's smoking cessation rate at the time of delivery [52]. One concern about NRT use in pregnancy is the possibility of adverse effects of nicotine on the fetus, through alterations in uterine, placental, or blood flow, or directly on the brain. As there are still too few trials to assure it's safe use in pregnancy, and animal studies suggest nicotine may be toxic to the developing central nervous system, registries of women using NRT should be established to gather more outcome data. A 2-year follow up study showed that infants born to women who used NRT for smoking cessation in pregnancy were more likely to have unimpaired development [53]. There is insufficient evidence to assess the safety of NRT, nicotine gum, and also the effect on incidence of PTB. No trial has been done using bupropion (see Chapter 22 in Maternal-Fetal Evidence Based Guidelines).

**Other interventions.** Interventions to increase smoking cessation among the partners of pregnant women, with the additional aim of facilitating cessation by the women themselves, have been insufficiently studied (only one trial) [45]. Stages of change, or feedback, do not show benefit [45]. A recent RCT showed no benefit of combining supervised exercise and physical activity counseling, to improve the effectiveness of behavioral support for smoking cessation in pregnancy [54]. There has been increased awareness and use of e-cigarette or electronic nicotine delivery system (ENDS) in adolescents (especially in middle and high school students), health impact of which is not clear [55,56]. E-cigarettes are marketed as smoking cessation devices and as better alternatives to regular cigarettes. This perception of harm reduction in comparison to regular cigarettes may be prevalent due to the lack of Food and Drug Administration (FDA) regulation on e-cigarette advertising and this potentially can further increase use of e-cigarette in adolescents/reproductive age women and in pregnant women as well [55,57]. ENDS device do not burn tobacco leaves, instead delivers liquid nicotine (with other ingredients and by-products from metals, plastics, rubbers, ceramics, fibers and foams) as an aerosol by heating and vaporizing the liquid components through a battery charged atomizer [58] (see Chapter 22 in Maternal-Fetal Evidence Based Guidelines).

Risk: Short Cervix on Ultrasound

**Intervention: Activity restriction.** Activity restriction is not associated with prevention of PTB in asymptomatic singleton gestations with a TVU CL <30 mm, and in fact is associated with a 237% increase in PTB incidence [59].

**Intervention: Progesterone.** Regarding 17P, a multicenter RCT evaluating the effect of 17P in women with singleton gestations, no prior PTB, and short CL <30 mm compared with placebo showed no difference in rate of PTB <37 weeks (25.1% vs. 24.2%, RR 1.03, 95% CI 0.79–1.35) [60]. In another smaller RCT including women with singleton pregnancies (66% of
which had no prior PTB) with CL <25 mm between 16 and 24 weeks, 17P was associated with similar incidences of PTB and neonatal morbidity and mortality compared with cerclage [61]. Cerclage was significantly more effective than 17P at reducing the incidences of PTB <35 and <37 weeks in the subgroup with CL ≤15 mm [61]. In summary, 17P cannot be recommended for prevention of PTB in singletons gestations (no prior PTB) with a short TVU CL.

Regarding vaginal progesterone, there are several RCTs available. In 250 women with mostly (90%) singleton gestations and CL ≤15 mm at 20–25 weeks, of whom about 85% had no prior PTB, vaginal progesterone 200 mg nightly started at 24 weeks until 34 weeks was associated with a 44% significant decrease in SPTB <34 weeks (19% vs. 34%, RR 0.56, 95% CI 0.36–0.86), but no significant effects on neonatal morbidities (composite neonatal adverse outcomes: RR 0.57, 95% CI 0.23–1.31) [62]. A subgroup analysis of only women without prior PTB confirmed significant benefit of progesterone in preventing PTB <34 weeks (RR 0.54, 95% CI 0.34–0.88) [62]. The incidence of CL ≤15 mm was 1.7%. Based on the frequency of short CL and effectiveness for prevention of SPTB <34 weeks from the work of Fonseca et al., the number of women needed to be screened with CL in order to prevent one SPTB <34 weeks is approximately 387, if all women with a CL ≤15 mm receive vaginal progesterone. Once the short CL ≤15 mm is identified, the number needed to treat to prevent one PTB <34 weeks is 7.

In 458 women with singleton gestations and CL 10–20 mm at 19–23 6/7 weeks, of whom about 84% had no prior PTB, vaginal progesterone 90 mg daily started at 20–23 6/7 weeks until 36 6/7 weeks was associated with a 45% significant decrease in PTB <33 weeks (8% vs. 16%, RR 0.55, 95% CI 0.33–0.92) and 43% significant decrease in composite neonatal morbidity and mortality (8% vs. 14%, RR 0.57, 95% CI 0.33–0.99) [63]. The incidences of PTB <28 and <35 weeks, and RDS, were also significantly decreased. Analysis of only women without prior PTB confirmed significant benefit of progesterone in preventing PTB <33 weeks (8% vs. 15%, RR 0.50, 95% CI 0.27–0.90) [63]. The incidence of CL 10–20 mm was 2.3%. Based on the frequency of short CL and effectiveness for prevention of PTB <33 weeks from this study [63], the number of women needed to be screened with CL in order to prevent one PTB <33 weeks is approximately 604, if all women with a CL 10–20 mm receive vaginal progesterone. Once the short CL 10–20 mm is identified, the number needed to treat to prevent one PTB <33 weeks is 14.

A individual patient data meta-analysis of five high quality RCTs, also showed benefit of vaginal progesterone in asymptomatic women with sonographic short cervix (≤25 mm) in reduction of PTB <33 weeks (RR 0.58, 95% CI 0.42–0.80) and composite neonatal morbidity and mortality (RR 0.57, 95% CI 0.40–0.81) [64].

In summary, in women with singleton gestations, no prior SPTB, and short CL, vaginal progesterone is associated with reduction in PTB and composite perinatal morbidity and mortality. Based on these results, if a TVU CL ≤25 mm is identified at ≤24 weeks, vaginal progesterone should be offered for prevention of PTB [62–64]. There is insufficient evidence that any of the vaginal preparations or doses are superior, as they have not been compared. Vaginal progesterone 200 mg suppository has been used in the trial for CL ≤15 mm [62], and 90 mg gel for the trial for CL 10–20 mm [63]. Therefore, CL, but also cost, availability, and other factors may influence preferred dosing [65,66].

These results also support universal screening with a single TVU assessment of CL at around 18–24 weeks in singleton gestations without prior SPTB (Figure 17.2). TVU CL screening of singleton gestations fulfills all criteria for an effective screening program. [67]. For example, multiple cost-effectiveness analyses evaluating universal CL screening in singleton gestations, to identify those with short CL eligible for vaginal progesterone, have been published so far [65,66,68]. All reported that such a strategy would be cost-effective, and in fact cost-saving.

In one study, compared with other managements, including no screening, “universal” sonographic screening of CL in singletons was associated with a reduction of 95,920 PTBs <37 weeks annually in the United States, and was actually cost-saving (almost $13 billion saved) [65]. Even varying the variables (e.g., the cost of vaginal progesterone or of TVU screening), universal screening was the preferred strategy 99% of the time [65].

The other cost-effectiveness analysis, when analyzing universal screening of singleton gestations without prior PTB with TVU CL at 18–24 weeks, calculated over $12 million saved, 424 quality-adjusted life-years (QALY) gained, and 22 neonatal deaths or long-term neurologic deficits prevented for every 100,000 women screened, compared with no screening. Even varying the variables (e.g., the cost of vaginal progesterone or of TVU screening), universal screening was cost-effective over 99% of the time [66].

It should be noted that only 1.7%–2.3% of women were identified to have short CL in the two large trials published [62,63], and that the incidence of CL ≤20 mm at 18–24 weeks is even lower (about 0.8%) in singletons without prior SPTB [67]. There are more limited data to evaluate the effectiveness of vaginal progesterone for CL 21–25 mm [64].

Guidelines from Society for Maternal-Fetal Medicine (SMFM) and ACOG state that implementation of universal TVU CL screening should be viewed as reasonable, and can

![Figure 17.2 Transvaginal ultrasound (TVU) cervical length (CL) screening for the woman with a singleton gestation and no prior spontaneous preterm birth (SPTB) at 16–36 6/7 weeks.

*For example, daily 200 mg suppository or 90 mg gel from time of diagnosis of short CL to 36 weeks. (Adapted from Romero R et al., Am J Obstet Gynecol, 206, 124.e1–e19, 2012; Society for Maternal-Fetal Medicine and Berghella V, Am J Obstet Gynecol, 206, 376–386, 2012.)
be considered by individual practitioners; third-party payers should not deny reimbursements for this screening [69,70]. International Federation of Gynecology and Obstetrics (FIGO) recommends TVU CL screening [71]. TVU CL examinations should be done following strict quality criteria, in the US via Cervical Length Education and Review (CLEAR) and the Perinatal Quality Foundation (https://clear.perinatalquality.org), and in Europe through the Fetal Medicine Foundation [72]. Only TVU, and not transabdominal (TA) ultrasound, should be used for CL screening [69,70].

Screening with TVU CL at 18–24 weeks should be offered to all singleton gestations without prior SPTB. If CL <25 mm, vaginal progesterone (e.g., 200 mg suppositories daily until 36 weeks) should be recommended [62–64].

**Intervention: Cerclage (ultrasound-indicated cerclage [UIC]).** A UIC involves first screening of pregnancies with TVU of the cervix to determine during pregnancy the risk of PTB, since the majority of even women at high risk by obstetrical risk factors for PTB do not develop a short CL and deliver at term even without intervention. A short CL (<25 mm) on TVU in the second trimester (between 14 and 23 6/7 weeks) significantly increases the risk of PTB in all populations studied [74]. UIC is defined as a cerclage performed because a short CL has been detected on TVU during pregnancy, usually in the second trimester. This cerclage has also been called in the past therapeutic, salvage, or rescue cerclage, but these terms are confusing and should be avoided. UIC has differing effects in different populations.

In women with singleton gestations, no prior PTB or other risk factors for PTB, and CL <25 mm before 24 weeks, cerclage is associated with no significant effect of PTB since the relatively small numbers of women (n = 235) included in RCTs done so far. Therefore, cerclage cannot be recommended in this population, but more research is needed.

For singleton gestations with prior PTB and a short TVU CL, see under the section “Prior PTB and Short Cervix.”

**Intervention: Pessary.** There is contradictory evidence regarding the efficacy of pessary to prevent PTB in women with singleton gestations and a short TVU CL <25 mm in the second trimester. All studies so far used the Arabin pessary. While the first RCT revealed a 82% decrease in PTB <35 weeks RR (0.76, 95% CI 0.52–1.15) [75]. This is probably due to the relatively small numbers of women (n = 235) included in RCTs done so far. Therefore, cerclage cannot be recommended in this population, but more research is needed.

For singleton gestations with prior PTB and a short TVU CL, see under the section “Prior PTB and Short Cervix.”

**Intervention: Indomethacin.** There is insufficient evidence to evaluate the effect of indomethacin on incidence of PTB in women with a short CL on TVU [79], as no RCTs have been performed.

**Intervention: Antibiotics.** There is insufficient evidence to evaluate the effect of antibiotics on incidence of PTB in women with a short CL on TVU, as no RCTs have been performed.

**Risk: “Pregnancy High-Risk for PTB”**

**Intervention: Activity restriction/bed rest.** Activity restriction and/or bed rest are probably the most commonly prescribed intervention for PTB prevention despite no proven benefit and potential for increased maternal morbidity. There is no evidence supporting bed rest or activity restriction to prevent PTB [80,81]. Per ACOG, bed rest should not be routinely recommended for prevention of PTB [82]. Bed rest (rest 1 hour tid) in (asymptomatic and symptomatic) “high-risk” singleton pregnancies is not associated with prevention of PTB over no bed rest [83]. Bed rest can be associated with an increased incidence of complications; in-hospital extended strict bed rest for PTB or PPROM is associated with an up to 1%-2% incidence of thromboembolic disease. Moreover, muscle wasting, cardiovascular deconditioning, bone demineralization, impaired glucose tolerance, heartburn, constipation, failure of volume expansion, headaches, dizziness, fatigue, depression, anxiety, stress, as well as lost wages, lost domestic productivity, and other costs may be other detrimental consequences of bed rest.

It is true that rest decreases uterine activity, and exercise increases it, but these are small effects that do not change rates of PTB. In nonrandomized studies, exercise in pregnancy has been associated with a decrease in PTB [84], while physically demanding work, prolonged standing, shift and night work, and high cumulative work fatigue score have been associated with PTB. Despite its use in about 20% of pregnancies, bed rest or any activity restriction for prevention of PTB cannot be recommended. These interventions should be studied in trials before clinical use. If prescribed bed rest, women should be allowed to ambulate to the bathroom a few times a day to limit complications of strict bed rest. It is possible that women at real risk of PTB from the above or other risk factors have not been studied adequately with this intervention of bed rest.

**Intervention: Support.** Programs of additional support during at-risk pregnancy (varying definitions) usually by a professional (social worker, midwife, or nurse) do not reduce PTB or LBW [85]. “Additional support” was defined as some form of emotional support (e.g., counseling, reassurance, and sympathetic listening) with or without additional information/advice, occurring during home visits, clinic appointments, and/or by telephone; most of the times these were intensive programs started in the first or second trimesters to the end of pregnancy. Other significant outcomes are as follows: antenatal hospital admission and cesarean delivery are decreased [85].

**Intervention: Weekly manual examinations, education.** A program of weekly manual cervical examinations in addition to education for women at high-risk for PTB (≥10 on Creasy’s score) does not reduce PTB [86–88].

**Intervention: Antibiotics.** See “Intervention: Antibiotics” in the section “Risk: Prior PTB.”

**Intervention: Cerclage.** Different specific clinical scenarios have been studied for possible benefit of cerclage. For efficacy of history-indicated cerclage, TA cerclage, UIC, PEIC, as well as cerclage in twins (see below).

**Risk: Prior PTB [89]**

**Intervention: Low-dose aspirin.** A recent secondary analysis of an RCT (underpowered) showed no association of preconception low-dose aspirin in women with one or more prior pregnancy loss or PTB on overall PTB rates (Figures 17.3 and 17.4) [90].

**Intervention: Fish intake.** Moderate fish intake, up to three meals per week, before 22 weeks is associated with a reduction in repeat PTB, compared with women eating fish less than once per month (odds ratio [OR] 0.60, CI 0.38–0.95) [91]. Moderate fish intake should be encouraged in women with prior PTB for prevention of recurrent PTB (see the section “Primary Prevention”).
PTB <37 weeks and a singleton gestation. The same omega-3 fatty acid regimen does not reduce PTB in women with twins [92]. In another larger RCT, in singleton gestations with prior SPTB and on 17P 250 mg IM, omega-3 supplementation (1200 mg EPA and 800 mg DHA) from 16 to 22 weeks until 36 weeks was associated with similar incidences of PTB <37 weeks (38% vs. 41%; RR 0.91, 95% CI 0.77–1.07) and PTB <35 weeks (RR 0.95, 95% CI 0.72–1.25) [93]. Meta-analysis of these two RCTs revealed that women who received omega-3 had similar rates of PTB at <37 weeks (34.5% vs. 39.8%; RR, 0.81; 95% CI, 0.59–1.12) and PTB at <34 weeks (12.0% vs. 15.4%; RR, 0.62; 95% CI, 0.26–1.46) compared with control. The omega-3 groups had a statistically significantly longer latency (mean difference, 2.10 days; 95% CI, 1.98–2.22) and higher birthweight (mean difference, 102 g; 95% CI, 20–185) compared with control subjects [94]. In summary, in singleton gestations with prior SPTB on 17P, omega-3 supplementation does not seem to be beneficial in preventing recurrent PTB. The benefits in longer latency and higher birth weight may deserve further study (see the section “Primary Prevention”).

Intervention: Antibiotics. Antibiotics to prevent PTB in women with prior PTB have been evaluated either as preconception (one RCT) or as prenatal intervention.

In women with prior PTB <34 weeks, preconception oral azithromycin 1 g twice (4 days apart) and metronidazole 750 mg daily for 7 days are not associated with an effect of subsequent PTB or miscarriage rates [95].

Clindamycin cream 2% for 7 days at 26–32 weeks does not reduce PTB <37 weeks in women with a prior PTB 24–36 weeks but may increase PTB <34 weeks, especially in women without BV, so that antibiotics in this setting may actually be detrimental [96].

Cefetamet Pivoxil (not available in United States) 2 g × 1 at 28–32 weeks in women in Nairobi with prior PTB, fetal death, or LBW did not affect GA at delivery (PTB was not reported) [97].

Metronidazole 250 mg tid × 7 days and erythromycin base 333 mg tid × 14 days in women with a prior PTB or pre-pregnancy weight <50 kg do not prevent PTB <37 weeks but may increase PTB <34 weeks [98]. Antibiotic prophylaxis did not reduce the risk of PPROM or PTB (except in a subgroup of women with a prior PTB who had bacterial vaginosis in current pregnancy) [99].

In summary, antibiotics given just because of a prior PTB do not prevent recurrent PTB.

Intervention: Progesterone. The effect of progesterone supplementation should be evaluated according to different patient populations, and according to type of progesterone. Here we review progesterone for prior PTB.

For 17P, there are at least two RCTs available in women with prior PTB. In 43 women with mostly (>90%) singleton gestation and either prior PTB or prior >1 spontaneous abortion, 17P 250 mg IM weekly started as soon as prenatal care began was associated with significant reduction in PTB <37 weeks and perinatal mortality compared with placebo [100]. In 463 women with singleton gestation and prior SPTB at 20–36 6/7 weeks of a singleton gestation, compounded 17P 250 mg IM weekly started at 16–20 6/7 weeks was associated with reduction in the incidences of PTB <37 weeks (RR 0.66, 95% CI 0.54–0.81), PTB <35 weeks, and <32 weeks, as well as of supplemental oxygen, NEC and IVH, compared with placebo [101]. The number needed to treat to prevent one recurrent PTB was 5.4. Based mostly on this clinical trial, 17P 250 mg IM weekly

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**Figure 17.3** TVU CL screening for the woman with a singleton gestation and prior SPTB at 16–36 6/7 weeks, ≥250 mg intramuscularly (IM) every week from 16 to 20 weeks to 36 weeks; vaginal progesterone can be used as well as an alternative. a Every 2 weeks; if 26–29 mm, repeat in 1 week. (Adapted from Society for Maternal-Fetal Medicine and Berghella V, Am J Obstet Gynecol, 206, 376–386, 2012.)

**Figure 17.4** Clinical algorithm for care of asymptomatic women with multiple prior PTB or second-trimester losses (STLs) (see text). (Adapted from Iams JD and Berghella V, Am J Obstet Gynecol, 203(2), 89–100, 2010.)

**Intervention: Omega-3 fatty acids.** In one RCT, omega-3 fatty acids (fish oil, Pikaosal: 52% eicosapentaenoic acid [EPA], 23% DHA, and 2 mg tocopherol/mL; 4 capsules/day: 1.3 g EPA and 0.9 g DHA, total 2.7 g/day; started ≥16 weeks at an average of 29–30 weeks) were associated with reduction in PTB <37 weeks by 46% and PTB <35 weeks by 68% in women with a prior

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<table>
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<th>Singleton with prior SPTB</th>
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<tr>
<td>17-OH progesterone at 16 weeks*</td>
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<tr>
<td>Serial TVU CL at 16–23 6/7 weeks</td>
</tr>
<tr>
<td>CL &lt; 25 mm</td>
</tr>
<tr>
<td>Cerclage; continue 17-OH progesterone</td>
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<tr>
<td>CL ≥ 25 mm</td>
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<td>Continue 17-OH progesterone</td>
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**Figure 17.3** TVU CL screening for the woman with a singleton gestation and prior SPTB at 16–36 6/7 weeks, ≥250 mg intramuscularly (IM) every week from 16 to 20 weeks to 36 weeks; vaginal progesterone can be used as well as an alternative. a Every 2 weeks; if 26–29 mm, repeat in 1 week. (Adapted from Society for Maternal-Fetal Medicine and Berghella V, Am J Obstet Gynecol, 206, 376–386, 2012.)

**Figure 17.4** Clinical algorithm for care of asymptomatic women with multiple prior PTB or second-trimester losses (STLs) (see text). (Adapted from Iams JD and Berghella V, Am J Obstet Gynecol, 203(2), 89–100, 2010.)
started at 16–20 weeks is recommended for all women with prior SPTB 20–36 6/7 weeks [102–104] (Figure 17.3). The estimated number of prevented PTBs <37 weeks in the United States by this policy is about 9870.

For **vaginal progesterone**, there are at least two RCTs available in women with prior PTB. In 142 women with singleton gestations and mostly (>90%) prior PTB, vaginal progesterone 100 mg nightly from 24 to 34 weeks was associated with significant reduction in the incidences of PTB <37 weeks (RR 0.48, 95% CI 0.25–0.96) and <34 weeks, as well as contraction frequency, compared with placebo [105]. In 659 women with singleton gestation and prior SPTB 20–35 0/7 weeks, vaginal progesterone 90 mg every morning starting at 18–22 6/7 weeks until 37 0/7 weeks was not associated with significantly different rates of PTB <37, 36, 33, 29 weeks and neonatal morbidity and mortality [106]. Several women screened for this trial were excluded from inclusion because of short CL [106]. Therefore, vaginal progesterone cannot be recommended for prevention of recurrent PTB when data against placebo is considered.

For **comparison of 17P versus vaginal progesterone**, there are two RCTs. In 518 women with singleton gestation and prior PTB between 20 and 34 weeks, 17P 250 mg IM weekly was associated with similar incidences of PTB and neonatal morbidities and mortality compared with vaginal progesterone 90 mg (Crinone) daily, except for significant lower incidences of PTB 28–31 6/7 weeks, neonatal intensive care unit (NICU) admission, and maternal side effects in the vaginal progesterone group [107]. Another RCT reported similar outcomes, including incidence of PTB, with either vaginal progesterone or 17P in women with singleton gestations and prior PTB [108] Another recent nonblinded randomized trial of 78 singleton pregnancies from Iran, compared effectiveness of vaginal progesterone with 17P in women with either prior PTB or short cervix TVU CL <25 mm. The authors concluded that vaginal progesterone and IM progesterone have same level of effectiveness and reported similar rate of PTB in two groups [109]. Overall this RCT does not appear to be high quality and the study results should be used cautiously. More data are necessary to evaluate how 17P and vaginal progesterone compare in efficacy in singletons with a prior PTB, as other populations, but vaginal progesterone seems to be at least equivalent, if not possibly more efficacious, than 17P in women with prior SPTB.

For **oral progesterone**, there are two small RCTs. In 150 women with singleton gestation and prior SPTB 20–36 6/7 weeks, oral micronized progesterone 100 mg twice a day was associated with significantly reduced incidences of PTB <37 weeks (39.2% in the oral progesterone vs. 59.5% in the placebo group, p = .002) and NICU admission compared with placebo [110]. In 33 women with singleton gestation and prior PTB 20–36 6/7 weeks, oral progesterone 400 mg daily was associated with nonsignificant reductions in the incidence of PTB <37 weeks (26% vs. 57%) and ventilator use (0% vs. 21%) compared with placebo [111]. In summary, there is insufficient evidence to assess the efficacy of oral progesterone for prevention of PTB in women with prior PTB.

In summary, in women with prior SPTB and singleton gestation, progesterone administration is beneficial in preventing PTB. Although we have limited data comparing the different preparations of progesterone, there is at present stronger evidence of effectiveness for 17P than for vaginal progesterone, based on the two largest trials [101,106], but comparison RCTs seems to show vaginal progesterone is an effective alternative [107,109]. 17P 250 mg IM weekly starting at 16–20 weeks until 36 weeks should be recommended to women with singleton gestations and prior SPTB 20–36 6/7 weeks [101]. In cases in which 17P is unavailable, other progesterone preparations may be considered [105].

**Intervention: Cerclage.** A history-induced cerclage is placed based solely on prior ob-gyn history (previously called a prophylactic or elective cerclage). Trials on women with only one or two prior PTBs have not shown benefit from history-induced cerclage [112,133]. A history-induced cerclage has been associated with prevention of PTB in women with three or more STLs or PTBs [114]. Cerclage decreases the incidence of PTB <37 weeks from 53% (with no cerclage) to 32%, and the incidence of PTB <32 weeks from 32% (with no cerclage) to 15% in women with three or more prior PTBs or STLs [114] (Figure 17.4).

The other clinical indication for history-induced cerclage might include cervical insufficiency, defined by some as prior painless cervical dilation leading to recurrent STLs. Unfortunately, no trial has been done to confirm the efficacy of history-induced cerclage in reducing PTB in women with a diagnosis of cervical insufficiency. Other indications such as prior cone biopsy, Mullerian anomaly, diethylstilbestrol (DES) exposure, prior PTB are not associated with cervical insufficiency, and Ehler-Danlos have occasionally been used clinically but have not been confirmed by any trial as indications that benefit from history-induced cerclage. History-induced cerclage is usually performed at 12–15 weeks’ gestation, and its techniques have been well described [74].

**TA cerclage** has been associated with less recurrent PTB compared with controls receiving transvaginal cerclage in women with a history of a failed SPTB <33 weeks despite cerclage) transvaginal history-induced cerclage in a case-control study [115]. It should be noted that in this study antibiotics and progesterone were uniformly given to the TA cerclage women. There is no RCT on TA cerclage. The efficacy of TA cerclage for other clinical scenarios such as a cervix with no intravaginal portion has not been adequately studied. TA cerclage can be performed prophylactically at around 10–12 weeks, and its technique has been well described [74,115]. TA cerclage has been successfully performed also laparoscopically and robotically, in particular prepregnancy, but also in the early first trimester [116].

In summary, the vast majority of women with prior PTB (e.g., those with only 1 or 2 prior PTBs) do not benefit from universal history-induced cerclage, and can instead be followed with TVU CL screening starting usually at 16 weeks. A policy of TVU CL screening with cerclage for short CL is associated with similar incidences of PTB <37 weeks (31% vs. 32%; RR 0.97, 95% CI 0.73–1.29), PTB <34 weeks (17% vs. 23%; RR 0.76, 95% CI 0.48–1.20), and perinatal mortality (5% vs. 3%; RR 1.77, 95% CI 0.58–5.35) compared with universal history-induced cerclage, and omits an obviously unnecessary cerclage in about 58% of these women [117]. Therefore, all singleton gestations with a prior spontaneous PTB should be screened with TVU CL starting at 16 weeks, every 2 weeks, until 23 6/7 weeks. If the CL is 26–29 mm, repeat TVU CL can be done in 1 week instead of 2 weeks. If the CL <25 mm is detected before 24 weeks, singleton gestations with prior PTB should undergo UIC [118] (Figure 17.3) (see also the section “Risk: Short Cervix on Ultrasound”).

If cerclage is performed, it should be performed according to best technique. McDonald cerclage, with suture (usually mersilene tape) placed as close to the internal os as possible (as high as possible), under spinal anesthesia, is recommended. There is usually no need for preoperative antibiotics or tocolytics [119]. A double suture might improve outcomes compared...
with one suture, but the evidence is still insufficient for a recommendation [120]. Cervical occlusion has no significant additional effect on cerclage [121].

**Intervention: Oral tocolytics.** There is insufficient evidence (only 1 small old RCT) to support the use of prophylactic oral betamimetics for preventing PTB in women with prior PTB with a singleton pregnancy [122].

For a summary of care of pregnant women with prior PTB, see also Ref. [89].

**Risk: Prior PTB and Short Cervix on Ultrasound**

**Intervention: Progesterone.** In a secondary analysis of an RCT evaluating just 46 singleton gestations with prior SPTB <35 weeks and short CL <28 mm at 18–22 6/7 weeks, vaginal progesterone 90 mg daily started at 18–23 6/7 weeks was associated with significant decreases in incidences of both PTB <32 weeks and NICU admission compared with placebo [123].

In a small RCT which did not recruit the planned sample size, 17P in singleton gestation are at high risk for PTB (56% with prior PTB) with TVU CL <25 mm 20–31 6/7 weeks was noted to have no benefit in prolonging pregnancy compared with no 17P. The sample size was too small, and the intervention probably too late, to show significance [124].

In a randomized trial that did not recruit the planned sample size, 17P and cerclage had similar effect in women for CL <25 mm for prevention of PTB, but cerclage was more effective in women with CL <15 mm [61]. So while cerclage seems to be more efficacious (lower RRs) as the CL is shorter [125,126], progesterone seems to be most efficacious in cases of “moderate” short CL [62,63].

Until further evidence is available, we suggest continuation of 17P in women with prior PTB and short cervix in the index pregnancy. Some instead have suggested switching from 17P to vaginal progesterone, but not based on RCT data [127]. There is no good evidence suggesting benefit of addition of vaginal progesterone in women with prior PTB on 17P who are noted to have short cervix [127]. In summary, women with prior PTB should be screened with TVU CL from 16 to 24 weeks. 17P should be recommended to women with prior SPTB starting at 16 weeks, as described above. If the cervix shortens, there is insufficient evidence to assess efficacy of a different progesterone therapy, and therefore it is reasonable to continue 17P until 36 weeks.

**Intervention: Cerclage.** In women with a singleton gestation, prior SPTB, and CL <25 mm before 24 weeks, cerclage is associated with a significant 30% reduction in PTB <35 weeks (28.4% vs. 41.3%; RR 0.70, 95% CI 0.55–0.89), significant reductions also in PTB <37, <32, <26, and <24 weeks, as well as a significant 36% decrease in composite perinatal mortality and morbidity (15.6% vs. 24.8%; RR 0.64, 95% CI 0.45–0.91), compared with no cerclage [118]. Therefore, cerclage is recommended in women with a singleton gestation, prior SPTB, and CL <25 mm before 24 weeks [69,70].

There are no randomized trials on management of subsequent pregnancy in women who required UIC in their prior pregnancy. Level II evidence suggests similar outcomes with TVU CL screening with UIC for short cervix of 25 mm or less, or planned HIC in the subsequent pregnancy, in singleton gestations with prior UIC. Authors reported less than 50% of the TVU CL screening group require a repeat UIC in the subsequent pregnancy [128]. Another retrospective study of women with prior UIC also reported that majority of women who underwent CL surveillance in the next pregnancy did not require intervention for short cervix. The authors reported higher rate of PTB in women who received HIC in the subsequent pregnancy, which was not justified based on their risk status [129]. Based on the available level II evidence, TVU CL screening with UIC for short cervix is acceptable and possibly more effective than routine history-indicated cerclage in women with UIC in prior pregnancy. A consideration can be given to HIC in the small subset of women who delivered prior to 32 weeks in their prior pregnancy with UIC [128].

If the TVU CL shortens again later after the UIC has been placed, a second “reinforcing” cerclage has not been associated with prevention of PTB [130,131].

For singleton gestations without a prior PTB and with a short TVU CL, see the sections “Risk: Short Cervix on Ultrasound” and “Intervention: Cerclage” (Ultrasound Indicated Cerclage).

**Intervention: Vaginal progesterone versus cerclage.** There is no RCT comparing directly vaginal progesterone to cerclage in this population, and therefore there is insufficient evidence for a recommendation. While an indirect meta-analysis reported that either vaginal progesterone or cerclage are equally efficacious in the prevention of PTB in women with a sonographic short cervix in the mid trimester, singleton gestation, and previous PTB, the populations compared were not similar, introducing significant bias [132]. We offer 17P based on the prior PTB, and cerclage for the short CL, based on data reviewed above.

**Intervention: 17P versus cerclage**

**Intervention: Progesterone in addition to cerclage.** There is insufficient evidence to assess the size of any cumulative effect of 17P for prior PTB in singletons already with cerclage for short CL. 17P was associated with reduction in PTB <35 weeks in women with UIC in the current pregnancy for TVU CL <25 mm, but sample size was too small to show significance [133,134].

**Intervention: Cerclage in addition to progesterone.** There is insufficient evidence to assess the size of any cumulative effect of cerclage for short CL in singletons already on 17P for prior PTB. UIC was associated with reduction in PTB <35 weeks in women already on 17P for prior PTB in the current pregnancy, but sample size was too small to show significance [135].

**Risk: Cervical Insufficiency**

**Intervention: Cerclage.** No intervention has been specifically studied in this population (see the section “Risk: Prior PTB”). There are no randomized trials evaluating the role of history-indicated cerclage in women with prior PTB or pregnancy loss due to cervical insufficiency. There is insufficient evidence to recommend a history-indicated cerclage in women with less than three prior PTBs or STLs. A policy of TVU CL screening with UIC if CL shortens to <25 mm at <24 weeks has shown to be equivalent to a policy of universal history-indicated cerclage in women with a prior loss or PTB [74,117,136].

**Intervention: 17P in addition to cerclage.** There is insufficient evidence to assess the possible cumulative effect of 17P on PTB <35 weeks in women with a history-indicated cerclage [137].

**Risk: IVF ART**

**Intervention: Progesterone or HCG.** 17P or HCG supplementation in the first trimester is associated with an increase in the incidence of fetal heart activity on ultrasound by 238%, and of pregnancy ≥24 weeks’ rate by 380% compared with placebo [138].
Risk: Amnioncensis

**Intervention: Progesterone.** Natural progesterone 200 mg IM q day for 3 days postamniocentesis followed by 17P 340 mg IM 2x/week until the second week after the amniocentesis did not reduce PTB <25 weeks in women undergoing amniocentesis [139].

Risk: Uterine Contractions Detected by Home Uterine Activity Monitoring (HUAM)

**Intervention: Varied, per obstetrican.** Uterine contractions have been associated with PTB, but their predictive value is poor. HUAM usually consists of 1 hour of tocomonitoring twice daily at 24–36 weeks. HUAM with or without nursing contact and education is not associated with prevention of PTB. Some studies show earlier (at lower cervical dilatation) detection of PTL. The lack of benefit in prevention of PTB might have been secondary to lack of effective intervention (usually tocolysis) once PTL was diagnosed. Three different populations of women at high risk for PTB have been studied: singleton gestations with risk factors for PTB (e.g., prior PTB), twin gestations, and women status-post an episode of PTL. Unfortunately, there is no published meta-analysis of all trials, and most trials do not report results for each population specifically, and also report differing outcomes. A meta-analysis of published data shows no decrease in PTB <37 weeks in any of these three subgroups: mostly singletons at high risk (nine trials; n = 3613) [140–149]; RR 1.01, 95% CI 0.91–1.11; twins (five trials; n = 998) [64,67,69–71]; RR 0.91, 95% CI 0.80–1.04; or women s/p PTL episode (four trials; n = 218) [67,72–74]; RR 1.21, 95% CI 0.92–1.60. The largest study [149] and a recent meta-analysis [17] showed more unscheduled antenatal visits and prophylactic tocolytic use in the HUAM group compared with controls. Therefore, HUAM should not be routinely provided for prevention of PTB.

Risk: Cervical Dilatation

**Intervention: Physical Exam-Indicated Cerclage (PEIC).** PEIC (aka emergency, or urgent) is the cerclage placed because of changes in the cervix (dilatation, effacement, etc) detected by physical (manual) examination. Since about 50% of women with asymptomatic cervical dilatation ≥2 cm in the second trimester have microbial invasion of the amniotic cavity, an amnioncensis should be considered before offering this cerclage. There are insufficient data to assess efficacy of PEIC in women with cervical dilatation in the second trimester, as only one small trial has been reported. In women with membranes at or beyond the external os at around 20–24 weeks, PEIC (and indomethacin) is associated with a delay of delivery of about 4 weeks compared with controls (30 vs. 26 weeks) [150]. The major limitations of this study are the small sample size and the inclusion of twins. Over 25 retrospective observational series, mostly with no controls, have claimed benefit of PEIC. The largest cohort study reported a significant decrease of 92% in prevention of PTB <28 weeks with PEIC, compared with no cerclage, in singletons with ≥1 cm of cervical dilatation before 26 weeks [151]. A recent meta-analysis reported that PEIC is associated with a significant increase in neonatal survival and prolongation of pregnancy of approximately 1 month when compared with no cerclage. The strength of this conclusion is limited by the potential for bias in the included studies [152]. For PEIC, perioperative indomethacin and antibiotics have been associated with a significant 28 days prolongation of pregnancy [153]. In summary, PEIC in singleton gestations with a cervix dilated to ≥1 cm in the second trimester is associated with prevention in PTB and neonatal benefits. Clearly a large, well-designed prospective randomized trial is needed to confirm these benefits.

**Intervention: Indomethacin.** There is insufficient evidence to evaluate the effect of indomethacin on incidence of PTB in women with a short CL on TVU [154], as no RCT has been performed. An RCT of perioperative indomethacin and antibiotics in women receiving a PEIC between 16–23 weeks, showed significant prolongation of gestation by 28 days compared with control [153].

Risk: Positive fFN

**Intervention: Antibiotics.** fFN is a basement membrane protein present between the decidua/uterus and fetal membranes/placenta and produced by the trophoblast. Its presence (>50 ng/mL) at ≥22 weeks in the cervicovaginal canal has been associated with an increased risk for PTB. In fact fFN is one of the best predictors of PTB in all populations, including asymptomatic low-and high-risk women, twins, and women in PTL. Even at 13–22 weeks, higher (using 90th percentile) fFN levels are associated with two-to-threefold increase risk in subsequent SPTB. In women found to be fFN positive at 21–25 weeks, treatment with metronidazole 250 mg tid and erythromycin 250 mg qid × 10 days is associated with similar incidences of PTB <37 weeks to placebo. Among women with a prior SPTB, this antibiotic regimen is associated with a significantly higher incidence of PTB <37 weeks than the placebo group [155]. In women with at least one other risk factor for PTB (e.g., prior PTB), and positive fFN at 24–27 weeks, metronidazole 400 mg orally tid for 7 days was associated with no effect on reduction of PTB and maternal and perinatal outcomes compared with placebo, rather authors reported that metronidazole use can be detrimental due to a trend of higher PTB rate, antenatal admission and LBW in metronidazole group compared with placebo [156]. In summary, screening low-risk or high-risk asymptomatic pregnant women for fFN is not effective, as there is no intervention (the one tried has been antibiotics) shown to alter outcomes.

Risk: Prior Intrauterine Growth Restriction (IUGR) or Pre-eclampsia

**Intervention: Low-dose aspirin.** Compared with no aspirin or placebo, low-dose aspirin (usually 50–150 mg) is associated with decreased incidence of PTB (RR 0.22, 95% CI 0.10–0.49) in women at high risk for pregnancy complications, such as those with prior preeclampsia or prior IUGR [157]. A large systematic review showed 8% reduction in rate of PTB <37 weeks in women (many with prior preeclampsia or IUGR) treated with antiplatelet agents (RR 0.92, 95% CI 0.88–0.97) [158]. Low dose aspirin started between 12 and 28 weeks is recommended in women at high risk of recurrent preeclampsia [159]. (See Chapters 1 and 45 in Maternal-Fetal Evidence Based Guidelines).

Risk: Periodontal Disease

**Intervention: Periodontal therapy.** Periodontal disease has been associated with increased risk of PTB in several observational studies. Periodontal treatment such as scaling, root planning, plaque control, and daily rinsing have been evaluated as an intervention to decrease PTB in women with periodontal disease [160,161].

Periodontal treatment has not been associated with decrease in PTB in women with periodontal disease (RR 0.63, 95% CI 0.32–1.22), even when controlling for probing depth and attachment loss for periodontitis criteria, multiparity, prior PTB, or genitourinary infections [162].
Infections

Risk: Asymptomatic Bacteriuria

Intervention: Antibiotics. Asymptomatic bacteriuria occurs in 2%–10% of pregnancies, can lead to pyelonephritis, and is associated with an increased risk of PTB. Screening for asymptomatic bacteriuria and treating for urine colony count of >100,000 bacteria/ml reduce the incidence of PTB by 73% (RR 0.27, 95% CI 0.11–0.62). Two studies, n = 242 [163]. The optimal time to perform the urine culture is unknown; it seems reasonable to perform the urine culture and treat, as done in most studies, at the first prenatal visit. Quantitative urine culture of a midstream or clean catch urine is the gold standard for detecting asymptomatic bacteriuria in pregnancy. The choices of a sulfonamide or sulfonamide-containing combination, a penicillin, or nitrofurantoin, based on the results of susceptibility testing, are appropriate regimens for the management of asymptomatic bacteriuria. A short (3–7 days) course of therapy of asymptomatic bacteriuria has become accepted practice, as is as effective as longer therapy. A single-dose regimen of antibiotics may be less effective than a short course (4–7 day) regimen [164]. Women with asymptomatic bacteriuria in pregnancy should be treated by the standard regimen of antibiotics until more data becomes available on the cure rate of shorter course 3–5 days regimen compared with standard regimen. Although it is recommended that a urine culture be done following treatment, with retreatment as necessary, the evidence is insufficient to specifically evaluate the effectiveness of this strategy. Treatment of asymptomatic pregnant women with lower colony counts is not currently recommended, but further study of appropriate strategies to manage these women is warranted. Asymptomatic women with even low (100+) colony-forming units (CFU) of GBS in the urine culture at 27–31 weeks have decreased PTB <37 weeks (5.4% in the penicillin group vs. 38% in the placebo group, p < .002) when treated with penicillin 1 million IU three times per day for 6 days compared with placebo [165].

Antibiotic treatment compared with placebo or no treatment is effective in clearing asymptomatic bacteriuria. Antibiotic treatment of asymptomatic bacteriuria is then clinically indicated to reduce the risk of pyelonephritis in pregnancy. If untreated, the overall incidence of pyelonephritis is about 21%. Overall, the number of women needed to treat to prevent one episode of pyelonephritis is seven, and treatment of asymptomatic bacteriuria will reduce the number of women needed to treat to prevent one episode of pyelonephritis to approximately 37 weeks, which in early studies prior to the availability of effective antimicrobial therapy, was 7 days) was associated with PTB <37 weeks, PTB <34 weeks, PTB <32 weeks, or PPROM [166]. However, treatment before 20 weeks may reduce the risk of PTB <37 weeks (OR 0.72, 95% CI 0.55–0.95).

In women with a previous PTB, treatment did not affect the risk of subsequent PTB <37 weeks, with a 17%–25% nonsignificant trend for benefit [166,167]. It may decrease the risk of PPROM and LBW. Subgroup analysis of treatment with metronidazole or clindamycin does not alter incidence of PTB <37 weeks [167].

Risk: Trichomonas vaginalis

Intervention: Antibiotics. Antibiotics (metronidazole only one tested) do not prevent PTB in women with asymptomatic T. vaginalis (TV) infection [167–171]. In fact, metronidazole as studied (two 2-g doses 48 hours apart; possibly excessive dose) is associated with 78% higher incidence of PTB <37 weeks [170], and similar incidences of PTB <32 weeks and perinatal mortality [100]. Even in women with a prior PTB, metronidazole is associated with an 84% higher risk of PTB [167]. Metronidazole does eradicate TV in >90% of pregnant women with TV. Therefore, at least for the purpose of decreasing PTB, asymptomatic women should not be screened for TV and treated with metronidazole at doses studied so far if positive for TV. Symptomatic women with TV should still be adequately treated with metronidazole as a single 2-g oral dose, or 500 mg twice a day for seven days (see Chapter 36 in Maternal-Fetal Evidence Based Guidelines).

Risk: BV, Candida, and Trichomonas

Intervention: Antibiotics. In one large RCT, screening for BV (treatment clindamycin 2% vaginal cream for 6 days), Candida (treatment local clotrimazole 100 mg for 6 days), and Trichomonas (treatment local metronidazole 500 mg for 7 days) was associated with lower risks of PTB <37 weeks (3% vs. 5%; RR 0.55, 95% CI 0.41–0.75). The incidence of PTB for BW <2500 g and <1500 g were significantly lower in the intervention group (RR 0.48, 95% CI 0.34–0.66 and RR 0.34; 95% CI 0.15–0.75, respectively) compared with screening with results unavailable to the managing physician [172]. Very few women were positive for Trichomonas, while about 21% of women had either BV or candida, or both.

Risk: GBS Vaginal–Cervical Colonization

Intervention: Antibiotics. GBS colonization of the cervicovaginal tract is common in pregnancy (10%–20%), and has been associated with a slight (OR 1.5–3, usually) increased risk of PTB. Antibiotic therapy (with erythromycin) does not prevent PTB in women with GBS colonization, or affect stillbirths. Subanalysis by heavy colonization did not change results [173].

Risk: Ureaplasma Vaginal–Cervical Colonization

Intervention: Antibiotics. Ureaplasma urealyticum and/or mycoplasma hominis colonization of the cervicovaginal
tract is common in pregnancy and has been associated with a possible increased risk of PTB. There is insufficient evidence to show whether giving antibiotics to women with ureaplasma in the vagina prevents PTB. The only trial did not report data on PTB [174]. Compared with placebo, erythromycin is associated with a nonsignificant 30% decrease in incidence of LBW <2500 g (RR 0.70, 95% CI 0.46–1.07). Although some studies appeared to meet the inclusion criteria for this review, in most studies ureaplasma was not an essential entry criterion or studies reported just a post hoc subgroup analysis of ureaplasma.

**Multiple Gestations (See Chapter 44 in Maternal-Fetal Evidence Based Guidelines)**

**Intervention: Bed Rest**

In uncomplicated twin pregnancies, prophylactic bed rest in the hospital does not reduce PTB, perinatal mortality, LBW, and other complications of pregnancy [175]. In fact, the incidence of PTB <34 weeks is significantly increased by 84% [176–180].

In twin pregnancies with cervical dilatation, bed rest in the hospital does not decrease PTB in women in Zimbabwe [181]. In the trial in which it was recorded, only 6% of women appreciated in-hospital bed rest. For complications, see the section “Intervention: Bed Rest” (singleton pregnancies).

**Intervention: Multifetal Reduction**

There is no trial to assess the effect of multifetal reduction to prevent PTB. Compared with triplets/higher order multiples, triplets/higher order multiples reduced to twins have to prevent PTB. Compared with triplets/higher order multiples reduced to twins have no trial to assess the effect of multifetal reduction to prevent PTB.

**Intervention: CL Screening**

Routine second-trimester TVU assessment of CL in twin gestation is not associated with improved outcomes when incorporated into the standard management of otherwise low-risk twin pregnancies in an RCT [182]. See below for progesterone, cerclage, and pessary as interventions for short CL in twins.

**Intervention: Progesterone**

17P. 17P is not associated with prevention of PTB or neonatal adverse outcomes in the over 6 RCTs including twins [183–188]. A meta-analysis revealed no benefit of 17P in unselected twin pregnancies [189].

In twins with short CL, there was no effect of 17P on PTB rates, but numbers are limited [186,190]. A meta-analysis showed that 17-alpha-hydroxy progesterone caproate is not beneficial in reducing PTB or adverse perinatal outcome in 175 twin pregnancies (with >90% of mothers with no prior spontaneous PTB) with a TVU CL ≤25 mm before 24 weeks.

In women with twins and prior PTB, 17P did not affect incidence of PTB [187].

In triplet gestations, 17P is not associated with an effect on the incidence of PTB [191,192].

**Intervention: Vaginal Progesterone**

Vaginal progesterone 90 mg daily starting at 24 weeks for 10 weeks, in 500 women with unselected twin gestation, was not associated with significant effects in incidences of PTB or perinatal morbidity and mortality [193]. Other RCTs showed also no effect of vaginal progesterone in unselected twins [194,195]. A recent double-blinded placebo controlled RCT on nonselected twin pregnancies with no prior PTB investigated use of daily vaginal progesterone or placebo from 18 to 21 weeks to 34 weeks for prevention of PTB <34 weeks [196]. There was no difference in mean GA at delivery, the rate of SPTB <34 weeks (18.5% in the progesterone group and 14.6% in the placebo group, OR 1.32, 95% CI 0.24–2.37), and no reduction of neonatal morbidity and mortality. Another RCT evaluated two different dose of vaginal progesterone (200 mg vs. 400 mg daily) to placebo in dichorionic multiple gestations, and showed no difference in rate of PTB <37 weeks in the three groups [197]. A meta-analysis revealed no benefit from vaginal progesterone in unselected twins [189]. In summary, vaginal progesterone should not be given to unselected twin pregnancies for prevention of PTB.

In women with twin pregnancies and a short TVU CL, a subgroup analysis of an individual patient data meta-analysis of multiple gestations with TVU CL ≤25 mm showed that vaginal progesterone is associated with a non-significant reduction in PTB <35 weeks (30.4% vs. 44.8%, RR 0.70, 95% CI 0.34–1.44), and significant reduction in composite neonatal morbidity and mortality (23.9% vs. 39.7%, RR 0.52, 95% CI 0.29–0.93) [64].

In summary, further trials are needed to evaluate the efficacy of vaginal progesterone in multiple gestations with short cervix. There is insufficient evidence to assess the effect of vaginal progesterone on unselected triplet gestations [194].

In women with prior SPTB, and a current multiple gestation, some experts have suggested the use of 17P starting at 16 weeks based on the historic risk factor [89], but there is insufficient level 1 evidence to make this a strong recommendation.

**Intervention: Cerclage**

History-indicated cerclage does not prevent PTB in unselected twin gestations in one small RCT [198].

In summary, there is insufficient evidence to assess the efficacy of UIC in twin pregnancies with a short TVU CL. UIC does not prevent PTB in a meta-analysis of the 49 twin gestations and TVU CL <25 mm included in the three RCTs published so far [75]. An individual patient level data meta-analysis of three RCTs showed no benefit of cerclage compared with no cerclage in preventing PTB <34 weeks [199]. For multiple gestations, there is no evidence that cerclage is an effective intervention for preventing PTBs and reducing perinatal deaths or neonatal morbidity [200].

A recent retrospective cohort study of asymptomatic twin gestation with short cervix (TVU CL ≤25 mm) between 16–24 weeks showed UC was not associated with perinatal outcomes compared with controls. However, in the planned subgroup analysis of asymptomatic twin pregnancies with TVU CL ≤15 mm before 24 weeks, UC was associated with a significant prolongation of pregnancy by almost 4 more weeks, significantly decreased SPTB <34 weeks by 49%, and admission to neonatal intensive care unit by 58% compared with controls [201]. Given these contradictory data, while UIC should not be recommended for twin gestations currently, further research is warranted.

There is insufficient evidence to assess the effectiveness of PEIC in twins, as no RCT has been done.
Intervention: Pessary

In unselected twin gestations, the largest RCT found no benefit of pessary use [202]. In summary, there sufficient evidence to recommend against pessary use for prevention of PTB in unselected twins.

There is contradictory evidence regarding the efficacy of pessary to prevent PTB in women with twin gestations and a short TVU CL in the second trimester. All studies so far used the Arabin pessary. One RCT on twins with TVU CL ≤25 mm revealed that SPTB <34 weeks was significantly less frequent in the pessary than in the expectant management group (11/68 [16.2%] vs. 26/66 [39.4%]; RR: 0.41; 95% CI 0.22–0.76) [203]. A secondary analysis of an RCT suggests pessary use is associated with decreased rate of PTB <28 weeks and <32 weeks (but not <37 weeks) in twin pregnancies with TVU CL less than 38 mm before 23 weeks GA [204]. In the largest RCT so far, pessary on twins with TVU CL ≤25 mm was not associated with prevention of PTB ≤34 weeks (33/106 [31%], vs. 28/108 [26%]; RR 1.20; 95% CI 0.78–1.83) or other PTB or neonatal outcomes [202]. In summary, there insufficient evidence to recommend pessary use for prevention of PTB in twins with short CL. In summary, pessary cannot be recommended for prevention of PTB in singleton gestations with a short TVU CL.

For preconception counseling, prenatal care, antepartum testing, mode of delivery, anesthesia, and postpartum/breastfeeding in multiple gestations, see Chapter 44 in Obstetrics: Normal and Problem Pregnancies. Chapter 28, 627–658.e12.

REFERENCES


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For preconception counseling, prenatal care, antepartum testing, mode of delivery, anesthesia, and postpartum/breastfeeding in multiple gestations, see Chapter 44 in Obstetrics: Normal and Problem Pregnancies. Chapter 28, 627–658.e12.

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Preterm labor

B. Anthony Armson

KEY POINTS
- See Chapter 17 for primary and secondary prevention of preterm birth (PTB) in asymptomatic women, and for risk factors and complications.
- The diagnosis of preterm labor (PTL) is based on the clinical criteria of regular uterine contractions (≥4/20 minutes or ≥8/hour) at 20–36 6/7 weeks with either:
  - Manually detected cervical dilatation of ≥3 cm, or
  - Transvaginal ultrasound (TVU) cervical length (CL) <20 mm, or
  - TVU CL 20–29 mm and positive fetal fibronectin (FFN)
- Threatened PTL (regular uterine contractions (≥4/20 minutes or ≥8/hour) at 20–36 6/7 weeks) should be assessed and managed with knowledge of TVU CL, and FFN results (Figure 18.1).
- Women with threatened PTL but TVU CL ≥30 mm have a ≤2% chance of delivering within 1 week, and a >95% chance of delivering ≥35 weeks without therapy, and should therefore not receive any treatment.
- Corticosteroids (e.g., betamethasone 12 mg intramuscular (IM) every 24 hours × 2 doses) given to pregnant women between 24 and 33 6/7 weeks prior to PTB (either spontaneous or indicated) are effective in preventing respiratory distress syndrome (RDS), intraventricular hemorrhage (IVH), necrotizing enterocolitis (NEC), and neonatal mortality. Steroids may be also considered for decreasing respiratory morbidity and other neonatal outcomes in some women ≥34 weeks.
- Single rescue course of antenatal corticosteroids should only be considered if more than 2 weeks have elapsed since the initial course of corticosteroids and a new episode of PTL or preterm premature rupture of membranes (PPROM) or impending risk of PTB presents again at <33 weeks. A single rescue course of betamethasone, two 12-mg doses 24 hours apart, received before 33 weeks at least 14 or more days after the first course, which was administered before 30 weeks, is associated with decreased RDS, ventilatory support, surfactant use, and composite neonatal morbidity. More than two courses of corticosteroids for fetal maturity should be avoided.
- Intravenous magnesium sulfate (loading dose of 4–6 g infused for 20–30 minutes, followed by a maintenance infusion of 1–2 g/hour) given at 24–31 6/7 weeks immediately (within 12 hours) before PTB is associated with a significant decrease in the incidence of cerebral palsy.
- Other interventions studied to prevent PTB in women with PTL and intact membranes, including bed rest, hydration, sedation, and antibiotics, have not been shown to be beneficial in the management of PTL.
- No tocolytic agent has been shown to reduce perinatal mortality.
- Tocolytics should not be used without concomitant use of corticosteroids for fetal maturity.
- There is no tocolytic agent that has been shown to be superior in safety and effectiveness. Cyclooxygenase (COX) inhibitors are the only class of primary tocolytics shown to decrease PTB <37 weeks compared with placebo. COX inhibitors and oxytocin receptor antagonists (ORAs) have been shown to significantly prolong pregnancy by 48 hours and by 7 days compared with placebo. COX inhibitors, calcium channel blockers (CCBs), and ORA have significantly less side effects than betamimetics. CCB and COX inhibitors are the tocolytic agents best supported by evidence for safety and effectiveness.
- In general, maintenance tocolysis has not been proven to prevent PTB or reduce perinatal morbidity/mortality, and therefore maintenance tocolysis should not be used. An exception is vaginal progesterone, which has been shown to be beneficial for maintenance tocolysis in small randomized controlled trials (RCTs) and a meta-analysis of these, and deserves more study.
- There is insufficient evidence to evaluate multiple tocolytic agents for primary tocolysis, refractory (primary agent is failing, so another is started) tocolysis, or repeated (after successful primary tocolysis) tocolysis.
- In preterm neonates, delayed cord clamping for 30–60 (maximum 120) seconds is associated with fewer transfusions for anemia, better circulatory stability, less IVH and lower risk of NEC compared with early clamping at <30 seconds.

DIAGNOSIS
PTL was often defined in the past as regular uterine contractions (≥4/20 minutes or ≥8/hour) with either manually-detected cervical change, or cervical effacement ≥80 percent, or cervical dilatation ≥3 cm at 20–36 6/7 weeks [1]. A more evidence-based definition using current diagnostic technology is regular uterine contractions (≥4/20 minutes or ≥8/hour) with TVU CL <20 mm, or positive FFN and TVU CL 20–29 mm, at 20–36 6/7 weeks. Threatened PTL can be defined as regular uterine contractions (≥4/20 minute or ≥8/hour) between 20 and 36 6/7 weeks, before cervical dilatation and TVU CL has been assessed.

SYMPTOMS
Symptoms of PTL are often nonspecific and include menstrual-like cramps, abdominal “tightenings,” mild irregular contractions, low backache, pelvic pressure, increased vaginal discharge, spotting or bleeding.
Pregnant patient with SIUP between 23/0/7 and 33 6/7 weeks c/o PTL (contractions >6/hour, abdominal pain/pressure, vaginal bleeding)

- Consider admission
- Corticosteroids for fetal maturity
- GBS prophylaxis
- Consult NICU and MFM
- Establish fetal presentation, EFW, AFI, placental location
- Order laboratories as appropriate, start IVF
- Consider amniocentesis to rule out infection

≥ 3 cm dilated or cervical change

- Indocin 50–100 mg load, then 25–50 mg q6 hrs to achieve 48 hours of steroids.
- Consider magnesium sulfate for fetal neuroprotection if delivery is considered imminent

< 3 cm dilated or no cervical change

Perform TVU CL and obtain FFN

- o SSE to r/o SROM → if +, then Rx per PPROM protocol, including latency Abx
- o Consider obtaining FFN, GBS, GC/Chl and Urine Cxs
- o Continuous EFM/tocometer

CL < 20 mm

- FFN pos
- Persistent ctx w/o cervical change → manage at physician’s discretion
- Discharge home with dx: preterm ctx

CL 20–29 mm

- FFN neg
- Persistent ctx w/o cervical change → manage at physician’s discretion

CL ≥ 30 mm

Figure 18.1 Suggested algorithm for the management of preterm labor (PTL). AFV, amniotic fluid volume; Amnio, amniocentesis; BP, blood pressure; Chl, chlamydia; c/o, complains of; CTX, contractions; dx, diagnosis; EFM, external fetal monitoring; EFW, estimated fetal weight; FFN, fetal fibronectin; GA, gestational age; GBS, group B streptococcus; GC, gonorrhea; IVF, intravenous fluids; MFM, maternal-fetal medicine specialist; NICU, neonatal intensive care unit; PPROM, preterm premature rupture of membranes; R/O, rule out; Rx, treat; SLIUP, single live intrauterine pregnancy; SROM, spontaneous rupture of membranes; SSE, sterile speculum examination; TVU CL, transvaginal ultrasound cervical length; w/, with.

WORKUP

- History: ensure correctness of gestational age (GA) estimation; evaluate signs and symptoms of PTL, and ALL risk factors for PTB (Table 17.1; Figure 17.1).
- Physical examination: maternal vital signs; frequency, intensity and duration of uterine contractions; fetal heart rate pattern; assess uterine tenderness, firmness, fetal position.
- Perform speculum examination:
  - Estimate cervical dilatation [2]
  - Assess presence/amount of uterine bleeding
  - Assess for PPROM (nitrazine, pooling, ferning)
  - Obtain swab for FFN testing
  - If no PPROM, perform digital cervical examination
  - If cervix <3 cm dilated, send FFN swab and perform TVU CL
  - If cervix ≥3 cm, discard FFN swab and manage PTL

- Laboratory tests: rectovaginal group B streptococcus (GBS) culture; gonorrhea and chlamydia; urinalysis and urine culture (Figure 18.1); FFN in women <34 weeks gestation with cervical dilation <3 cm and TVU CL 20–29 mm.

Transvaginal Ultrasound Cervical Length

Symptomatic women with CL ≥30 mm have a low risk (<5%) of delivering preterm, regardless of FFN result. For women with CL 20–29 mm, the PTB rate is somewhat increased but still <5% within 7 days if the FFN test is negative. A CL <20 mm is associated with a high risk (>25%) of PTB within 7 days. With this degree of cervical shortening, FFN testing does not improve the predictive accuracy of CL measurement alone [3,4].

Compared with no knowledge, knowledge of TVU CL results is associated with a nonsignificant decrease in PTB <37 weeks (22.3% vs. 34.7%, respectively; relative risk [RR] 0.59, 95% confidence interval [CI] 0.26–1.32). Delivery occurred at a later GA (by about 4–5 days) in the knowledge versus no knowledge groups [5]. Given the clinical and cost-effectiveness of TVU CL in the assessment of suspected PTL,
TVU CL should be used in the management of women with threatened PTL (Figure 18.1).

**Fetal Fibronectin**

FFN testing has been shown to predict PTB with moderate accuracy in symptomatic women resulting in health care cost benefits by identifying women who do not require intervention [6]. In a meta-analysis of six RCTs, compared with no knowledge, knowledge of FFN results in women with threatened PTL had no effect on the incidence of PTB <37 weeks or any other outcome, maternal or neonatal, including time in triage, PTB <34, 32, or 28 weeks; GA at delivery; birth weight less than 2500 g; perinatal death; maternal hospitalization; tocolysis; steroids for fetal lung maturity [7]. The benefit in knowledge of FFN was seen only in one RCT, in which TVU CL was the main screening test, with FFN used only for “indeterminate” results [8] (Figure 18.1).

**MANAGEMENT**

**Principles of Management**

Before treatment is considered, the diagnosis and risk of PTB must be established. Women with preterm uterine contractions but negative FFN and TVU CL ≥30 mm have a <2% chance of delivering within 1 week, and a >95% chance of delivering ≥35 weeks without therapy, and should therefore not receive any treatment [4] (Figure 18.1).

Women with preterm uterine contractions and cervical dilation ≥3 cm, TVU CL <20 mm or TVU CL 20–29 mm and positive FFN have a moderate to high risk of PTB within 7 days, are those with true PTL, and should be managed with strong consideration for admission, steroids, magnesium for neuroprophylaxis, and possibly tocolysis, depending on GA.

If PTL is diagnosed without using TVU CL, about 70%–95% at 26–28 weeks and 75%–95% at 29–30 weeks, whereas intact survival at 18 months is about 50% after 25 weeks [12]. Disabilities in mental and psychomotor development, neuromotor function (including cerebral palsy), or sensory and communication function are present in at least 50% of fetuses born ≤25 weeks’ gestation [13].

**Table 18.1** Estimated Incidences of Intra-Amniotic Infection in Women in Different Clinical Scenarios

<table>
<thead>
<tr>
<th>Condition</th>
<th>Percent</th>
</tr>
</thead>
<tbody>
<tr>
<td>GA &lt;37 weeks</td>
<td></td>
</tr>
<tr>
<td>Asymptomatic second trimester</td>
<td>0.5</td>
</tr>
<tr>
<td>PTL (intact membranes)</td>
<td>13</td>
</tr>
<tr>
<td>PPROM, no labor</td>
<td>25</td>
</tr>
<tr>
<td>PPROM, labor</td>
<td>39</td>
</tr>
<tr>
<td>Cervix ≥2 cm/80% in second trimester</td>
<td>50</td>
</tr>
<tr>
<td>GA ≥37 weeks</td>
<td></td>
</tr>
<tr>
<td>Labor</td>
<td>19</td>
</tr>
<tr>
<td>PROM</td>
<td>34</td>
</tr>
</tbody>
</table>

**Table 18.2** General Guidance Regarding Obstetric Interventions for Perivable Birth

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonatal assessment for resuscitation</td>
<td>Not recommended</td>
<td>Consider</td>
<td>Consider</td>
</tr>
<tr>
<td>Antenatal steroids</td>
<td>Not recommended</td>
<td>Not recommended</td>
<td>Not recommended</td>
</tr>
<tr>
<td>Tocolytics to allow steroids</td>
<td>Not recommended</td>
<td>Not recommended</td>
<td>Not recommended</td>
</tr>
<tr>
<td>Magnesium sulfate for neuroprotection</td>
<td>Not recommended</td>
<td>Not recommended</td>
<td>Not recommended</td>
</tr>
<tr>
<td>Antibiotics for PROM to prolong latency</td>
<td>Consider</td>
<td>Consider</td>
<td>Consider</td>
</tr>
<tr>
<td>Antibiotics for GBS prophylaxis</td>
<td>Not recommended</td>
<td>Not recommended</td>
<td>Not recommended</td>
</tr>
<tr>
<td>Continuous intrapartum electronic fetal monitoring</td>
<td>Not recommended</td>
<td>Not recommended</td>
<td>Not recommended</td>
</tr>
<tr>
<td>Cesarean delivery for fetal indication</td>
<td>Not recommended</td>
<td>Not recommended</td>
<td>Consider</td>
</tr>
</tbody>
</table>

**Source:** Adapted from Rogue TN, et al., Obstet Gynecol, 123:1083–1096, 2014.

**Abbreviation:** GBS, group B streptococcus.

*Survival of infants born in the perivable period is dependent on resuscitation and support. Several factors (e.g., intrauterine growth restriction, small fetal size, the presence of fetal malformations or aneuploidy, and pulmonary hypoplasia due to prolonged membrane rupture) can impact the potential for survival and the determination of viability. Personal and family values should be extensively discussed, allowing individual decisions.

*Persistently abnormal fetal heart rate patterns or biophysical testing (category II–III), malpresentation.
Pregnancies at risk for periviable PTB are particularly challenging to counsel and manage. The periviability period varies according to several factors, including by level of care provided in the hospital and the country where the care is being provided. In many developed countries, this period, in 2015, is between 22 and 24 weeks. Table 18.2 provides some guidance for care, but should be adjusted according to local capabilities [13]. Patients with an intrauterine demise or undergoing pregnancy termination are not to be included in this group. For patients in whom the GA has not been clearly established, plans for intervention and information provided to the patient should be based on the best estimate of GA. Several algorithms have been developed to provide patient-specific mortality and morbidity information [14], as well as an online calculator [15]. Status can be reevaluated with each additional gestational week.

**Prophylaxis to Prevent Neonatal Morbidity/ Mortality from PTB (Fetal Maturation)**

**Corticosteroids**

Betamethasone, dexamethasone (only two corticosteroids that cross the placenta reliably).

**Dose:** One course: Betamethasone 12 mg SQ every 24 hours x 2 doses or dexamethasone 6 mg SQ q12h x 4 doses.

**Mechanism of action:** Enhanced maturational changes in lung architecture and induction of lung enzymes resulting in biochemical maturation.

**Evidence for effectiveness:** Corticosteroids given prior to PTB (either spontaneous or indicated) are effective in preventing RDS, IVH, NEC, and neonatal mortality [16]. Antenatal administration of 24 mg of betamethasone (12 mg intramuscularly q24h), or of 24 mg of dexamethasone (6 mg intramuscularly q12h), to women expected to give birth preterm is associated with significant 31% reduction in neonatal mortality, 34% reduction in RDS and respiratory support, 46% reduction in IVH, 54% reduction in NEC, 20% reduction in intensive care admissions, and 44% reduction in systemic infections in the first 48 hours of life in preterm infants. There are also decreased needs for surfactant, oxygen, and mechanical ventilation in the neonatal period. Treatment with antenatal corticosteroids does not increase risk to the mother of death, chorioamnionitis, or puerperal sepsis.

These benefits apply to GAs of at least 24–33 6/7 weeks, and are not limited by gender or race. There is insufficient data to assess effectiveness before or after these GAs. At 22–25 weeks, cohort studies have shown benefit in decreasing neonatal death [17]. The effects are optimal at 48 hours to 7 days from first dose [17], but treatment should not be withheld even if delivery appears imminent. Antenatal corticosteroids should be administered when PTB is considered imminent within seven days, the GA estimate is accurate, adequate neonatal care is available and there is no clinical evidence of maternal infection.

There is new evidence that steroids are associated with neonatal benefits also ≥34 weeks, as three RCTs have been done on steroids ≥34 6/7 weeks in women at risk of PTB, and two RCTs in women at ≥37 weeks [18]. Women who received antenatal corticosteroids ≥34 weeks had a significantly lower incidence of RDS (RR 0.76, 95% CI 0.62–0.93), mild RDS (RR 0.40, 95% CI 0.23–0.69), moderate RDS (RR 0.39, 95% CI 0.18–0.89), transient tachypnea of the newborn (RR 0.62, 95% CI 0.50 to 0.77), severe RDS (RR 0.66, 95% CI 0.53–0.82), surfactant use (RR 0.61, 95% CI 0.38–0.99), mechanical ventilation (RR 0.62, 95% CI 0.41–0.94), significantly lower time on oxygen (mean difference [MD] –2.06 hours, 95% CI –2.17 to –1.95), lower maximum inspired oxygen concentration (MD –0.66%, 95% CI –0.69 to –0.63), lower LOS in NICU (MD –764 days, 95% CI –765 to –764), higher APGAR score at 1 and at 5 minutes (MD 0.06, 95% CI 0.05–0.07) compared with those who did not [18]. **Steroids should be considered for decreasing respiratory morbidities and other neonatal outcomes in women ≥34 weeks [19].**

**Type of steroid:** There is no clear evidence of superiority between betamethasone and dexamethasone. Based on a recent Cochrane review, dexamethasone was associated with a reduced risk of IVH compared with betamethasone (RR 0.44, 95% CI 0.21–0.92) [20]. There were no statistically significant differences in other perinatal outcomes including perinatal death, RDS, NICU admissions, however. In one study, infants exposed to dexamethasone had a significantly shorter stay in the NICU. Results for biophysical parameters were inconsistent, but no important differences were seen for these or other secondary outcomes. Indirect comparisons of betamethasone and dexamethasone suggested that betamethasone is more effective in reducing RDS risk than dexamethasone, while chorioamnionitis and puerperal sepsis was more likely with dexamethasone [20].

Oral dexamethasone significantly increased the incidence of neonatal sepsis (RR 8.48, 95% CI 1.11–64.93) compared with intramuscular dexamethasone in one trial of 183 newborns [20].

Betamethasone administered at 12-hourly compared with 24-hourly intervals has been associated with reduced maternal length of stay, but no other differences in maternal or neonatal outcomes [20].

In a recent international cluster randomized trial designed to increase the use of antenatal corticosteroid therapy in low-resource settings, involving almost 100,000 pregnant women, identification of women at risk for PTB and increased use of antenatal corticosteroids increased overall newborn mortality by 12% (RR 1.12, 95% CI 1.01–1.22), perinatal mortality by 11% (RR 1.11, 95% CI 1.04–1.19) and suspected maternal infection rate by 45% (OR 1.45, 95% CI 1.33–1.58) [21]. Possible explanations for these unexpected findings in low-resource settings include unreliable dating criteria to establish GA, the reliance on birth weight rather than estimated GA to define PTB and evaluate other important outcomes, and the coding of suspected maternal infection by treatment rather than diagnosis. Inconsistent access and limited resources to provide optimal neonatal intensive care and potential targeting of term, low birth weight infants for antepartum corticosteroid treatment may have also contributed to these results. As recently reaffirmed by the World Health Organization, the study’s findings should not alter current recommendations regarding the use of antenatal corticosteroids to improve newborn outcomes [22].

**Weekly repeat courses of antenatal corticosteroids:** Evidence for repeating antenatal corticosteroids weekly is provided by a recent Cochrane systematic review of 10 trials (4738 women/5700 babies) [23]. Administration of repeat dose(s) to women who remained at risk for PTB seven or more days after the initial dose was associated with a reduction in the risk of RDS (RR 0.83, 95% CI 0.75–0.91) and the risk of serious neonatal adverse outcome (RR 0.84, 95% CI 0.75–0.94). In the largest international multicenter RCT, multiple repeat doses were associated with decreased birth weight, length and head circumference at birth [24]. At early childhood follow-up, no statistically significant differences were seen for infants exposed to repeat prenatal corticosteroids compared with unexposed infants for the primary outcomes (total deaths; survival free of any disability or major disability; disability; or serious outcome) or in the secondary outcome growth assessments [23].
Single repeat (rescue) course of antenatal corticosteroids: Given the potential short and long term adverse effects of multiple doses of corticosteroids, combined with the optimal benefit 7 days following administration, timing the first course is extremely important. A single rescue course should only be considered if PTB has not occurred within 14 days of the initial dose, and subsequent clinical assessment demonstrates a new episode of PTL or PPROM and PTB appears highly likely within 7 days [25]. A single rescue course of betamethasone, two 12-mg dose 24 hours apart, received before 33 weeks at least 14 or more days after the first course, that was administered before 30 weeks, is associated with decreased RDS, ventilatory support, surfactant use, and composite neonatal morbidity (RR 0.65, 95% CI 0.44–0.97) [25,26]. Given the concerns regarding multiple courses of corticosteroids, more than two courses should be avoided.

Contraindications: None.

Side effects: When used for only 1 course, no significant side effects are seen, except for transient maternal hyperglycemia from 12 hours to about 5–7 days after the dose. This effect results in false-positive glucose screening tests or difficulty in managing diabetes. There is no significant increase in maternal or fetal/neonatal infection. If ≥4 courses are used, there is an association with birth weight <10th percentile and probably with small (<10th percentile) neonatal head circumference, with evidence of later “catch-up” [23,24]. No adverse consequences of prophylactic corticosteroids for PTB in either mothers or infants, even at 10+ years follow-up, have been identified, but long-term follow-up data is limited.

Thyrotropin-Releasing Hormone
Prenatal thyrotropin-releasing hormone (TRH), in addition to corticosteroids, given to women at risk of very PTB, does not improve infant outcomes and can cause maternal side effects [27]. Overall, prenatal TRH, in addition to corticosteroids, does not reduce the risk of neonatal respiratory disease or chronic oxygen dependence, and does not improve any of the fetal, neonatal, or childhood outcomes. Prenatal TRH may actually have adverse effects for women and their infants. Side effects (nausea, vomiting, lightheadedness, urgency, flushing) are more likely to occur in women receiving TRH. Among infants, prenatal TRH increases 16% the risk of needling ventilation, 48% having a low Apgar score at 5 minutes, and was associated with poorer childhood outcomes at follow-up [24].

Phenobarbital
The use of prophylactic maternal phenobarbital administration prior to preterm delivery does not prevent periventricular hemorrhage (PVH) or protect from neurological disability in preterm infants [28]. Prenatal maternal phenobarbital is associated with a significant 35% reduction in the rates of all grades of PVH and 59% reduction in severe grades PVH (3 and 4) in the infants. The results were influenced by poor quality trials that excessively influence the analysis due to their higher rates of severe PVH. When only higher quality trials were included, phenobarbital is not associated with any beneficial effects, including similar incidences of all grades of PVH and severe grades of PVH to placebo. No difference in the incidence of neurodevelopmental abnormalities at childhood follow-up assessed between 18 and 36 months of age was observed. Maternal sedation is a common side effect in women receiving phenobarbital [28].

Vitamin K
Vitamin K administered to women prior to very PTB has not been shown to significantly prevent PVHs or other neurodevelopmental abnormalities in preterm infants. Antenatal vitamin K is associated with a nonsignificant reduction in all grades of PVH (RR 0.76, 95% CI 0.54–1.06) and a significant reduction in severe PVH (grades 3 and 4) (RR 0.58, 95% CI 0.30–0.91) [29]. When the two quasi-randomized trials are excluded, antenatal vitamin K is not associated with a reduction in all grades of PVH (RR 0.87, 95% CI 0.60–1.26) or severe PVH (RR 0.82, 95% CI 0.49–1.36). Treatment with vitamin K results in a significant reduction in the Bayley Mental Development Index at 2 years of age; but, these results are derived from one trial with a high lost to follow-up rate. No difference is found in the incidence of other neurodevelopmental abnormalities at childhood follow-up at 18–24 months or 7 years of age between children exposed to vitamin K and children not exposed [29].

Ambroxol
Giving ambroxol to women at risk of PTB to prevent neonatal RDS is not supported by available evidence. Prenatal administration of ambroxol to reduce the risk of RDS in preterm infants has not been shown to reduce the incidence of RDS or perinatal mortality when compared with betamethasone or placebo. Maternal adverse effects were similar in ambroxol and betamethasone-treated women. Since the trials included in the Cochrane systematic review were of very low to moderate quality, there is insufficient evidence to support or refute this intervention [30].

Magnesium Sulfate for Neuroprotection
Antenatal magnesium sulfate therapy given to women at risk of PTB significantly reduces the risk of cerebral palsy in their child (RR 0.68, 95% CI 0.54–0.87). There was also a significant reduction in the rate of substantial gross motor dysfunction (RR 0.61, 95% CI 0.44–0.85). No statistically significant effect of antenatal magnesium sulfate therapy is detected on pediatric mortality (RR 1.04, 95% CI 0.92–1.17), or on other neurological impairments or disabilities in the first few years of life. There are higher rates of minor maternal side effects in the magnesium groups, but no significant effects on major maternal complications [31].

Magnesium sulfate should be administered to women with singleton or twin pregnancies 23 0/7–31 6/7 weeks for neuroprotection when PTB is imminent. Imminent PTB is defined as a high likelihood of birth due to active labor with ≥4 cm cervical dilatation with or without PPROM or planned PTB for maternal or fetal indications. Women with an indicated PTB anticipated within 2–24 hours (e.g., for severe preeclampsia) are candidates for magnesium sulfate.

Intravenous magnesium sulfate is administered with a loading dose of 4–6 g infused for 20–30 minutes, followed by a maintenance infusion of 1–2 g/hour [22]. If delivery has not occurred after 12 hours and is no longer considered imminent (e.g., if the patient is not having regular uterine contractions), the infusion should be discontinued and resumed when delivery is deemed imminent again (e.g., when contractions develop). If at least 6 hours have passed since the discontinuation of the magnesium sulfate, another loading dose should be given. Administration of magnesium sulfate for prevention of cerebral palsy should not delay the delivery.

Nontocolytic Interventions for PTL
Bed Rest
Bed rest in hospital or at home in singleton gestations complicated by PTL or PPROM has been evaluated in one multiintervention RCT [32]. Bed rest offered no benefit over no treatment or placebo in preventing PTB. In multiple pregnancies, bed rest in the hospital did not reduce the risk of PTB or
perinatal mortality whether uncomplicated twin pregnancy, or twin or triplet pregnancy with cervical dilatation [33].

Hydration
There is no benefit of intravenous hydration in preventing PTB. Intravenous hydration does not seem to be beneficial, even during the period of evaluation soon after admission, in women with PTL. Women with evidence of dehydration may, however, benefit from the intervention. Compared with bed rest alone, hydration is associated with similar incidences of PTB <37 weeks, <34 weeks, or <32 weeks, and of admission to neonatal intensive care unit (NICU) [34]. Cost of treatment was slightly higher (US$39) in the hydration group for hospital costs during a visit of less than 24 hours. Women studied were at low risk, as only 30% of women required tocolysis, and <30% had PTB. No studies evaluated oral hydration [34].

Screen for Infections
Several infections are associated with a higher risk of PTL and PTB. These include chlamydia, gonorrhea, syphilis, trichomoniasis, and others, as well as bacterial vaginosis (see Chapter 17). There is insufficient evidence to assess the effect of screening and treating for genitourinary infections in women with PTL, as no RCTs have been performed.

Antibiotics
There is no evidence of benefit with the use of prophylactic antibiotic treatment for PTL with intact membranes on important neonatal outcomes [35]. PTB <36 or 37 weeks was similar in antibiotics and placebo groups. There is a trend for a 52% increase in neonatal mortality for those who received antibiotics (RR 1.52, 95% CI 0.99–2.34), with similar overall perinatal mortality (RR 1.22, 95% CI 0.88–1.70) [30]. The only benefit is a 26% reduction in maternal infection with the use of prophylactic antibiotics. Follow-up at 7 years of the largest RCT showed increased incidence of functional impairment associated with erythromycin, and increased incidence of cerebral palsy associated with either antibiotic studied (erythromycin or co-amoxiclav) [36]. Given these data, antibiotics should not be used for prevention of PTB in women with PTL and intact membranes.

Tocolysis
Principles
Tocolytic therapy may provide short-term prolongation of pregnancy, allowing antenatal corticosteroid administration, magnesium sulfate for neuroprotection and/or maternal transport to a tertiary care facility. There is no evidence that treatment of PTL with tocolytic agents improves perinatal outcomes, however [22]. CCB and COX inhibitors are the agents best supported by evidence of safety and effectiveness (see Table 18.3).

Contraindications
See Table 18.4.

Primary Tocolysis—Single Agent
Betamimetics
Types: Ritodrine, terbutaline.

Dose: Ritodrine: 50–100 mg/min intravenous (IV) initial dose, increase 50 mg/min q10minute (max 350 mg/min) (orally [po]: 1–20 mg po q2–4h). Terbutaline: 0.25 mg subcutaneously (SQ) q20minute at first, then 2–3 hours; or 5–10 mg/min IV, max 80 mg/min; or 2.5–5 mg po q2–4h (hold for maternal hazards ratio [HR] >120/min).

Mechanism of action: Stimulate B2 receptor through cyclic adenosine monophosphate (AMP), so no free calcium for myometrial contraction.

Table 18.3 Principles of Tocolytic Therapy

- At 24–33 6/7 week, steroids for fetal maturation should always be given if tocolysis is initiated. Tocolytics should not be used without concomitant use of steroids for fetal maturation.
- Tocolysis is typically used for 48 hours to allow steroid effect. Given side effects, consider stopping tocolytic therapy at 48 hours after steroids given if PTL under control.
- No tocolytic agent has been shown to improve perinatal mortality.
- There is no tocolytic agent that is more safe and efficacious. COX inhibitors are the only class of primary tocolytics shown to decrease PTB <37 weeks compared with placebo. COX inhibitors and ORA have been shown to significantly prolong pregnancy at 48 hours and 7 days compared with placebo. COX inhibitors, CCB, and ORA, properly used, have significantly less side effects than betamimetics. Currently, CCB and COX inhibitors are the tocolytic agents best supported by evidence for safety and effectiveness.
- There is no maintenance tocolytic that prevents PTB or perinatal morbidity/mortality, and therefore maintenance tocolysis should in general not be used. One possible exception is vaginal progesterone, with more data needed. There is insufficient evidence to evaluate multiple tocolytic agents for primary tocolysis, refractory (primary agent is failing, so another is started) tocolysis, or repeated (after successful primary tocolysis) tocolysis.

Abbreviations: < CCB, calcium channel blockers; COX, cyclooxygenase; ORA, oxytocin receptor antagonists; PTB preterm birth.

Table 18.4 Contraindications to Tocolytic Therapy

Maternal
Chorioamnionitis
Severe vaginal bleeding/abruption
Preeclampsia
Medical contraindications to specific tocolytic agent
Other maternal medical condition that makes continuing the pregnancy inadvisable

Fetal
Death
Major (especially if lethal) fetal anomaly or chromosome abnormality
Other fetal conditions in which prolongation of pregnancy is inadvisable

Documented fetal maturity

Evidence for effectiveness (Table 18.5): Compared with placebo, betamimetics are associated with a decrease in the number of women in PTL giving birth within 48 hours (RR 0.68, 95% CI 0.53–0.88), and a decrease in the number of births within 7 days (RR 0.80, 95% CI 0.65–0.98) but there was no reduction in PTB <37 weeks [37]. No benefit is demonstrated for betamimetics on perinatal or neonatal death, and on RDS. A few trials reported the following outcomes, with no difference detected: cerebral palsy, infant death, and NEC. Betamimetics are significantly associated with the following side effects (see below): withdrawal from treatment due to adverse effects, chest pain, dyspnea, tachycardia, palpitation, tremor, headaches, hypokalemia, hyperglycemia, nausea/vomiting, nasal stuffiness, and fetal tachycardia [34]. There is insufficient evidence to assess which of the studied betamimetics is most effective and/or associated with fewer side effects, with most data reported for ritodrine. For comparison with
other tocolytics, RCTs are too small and varied to make meaningful comparisons [37].

**Specific contraindications:** Cardiac arrhythmia or other significant cardiac disease, DM, poorly controlled thyroid disease (for ritodrine).

### Table 18.5 Summary of the Evidence for Tocolytic Therapy

<table>
<thead>
<tr>
<th>Tocolytics</th>
<th>&lt;48 Hours</th>
<th>&lt;7 Days</th>
<th>&lt;34 Weeks</th>
<th>&lt;37 Weeks</th>
<th>Perinatal Mortality</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Betamimetics</strong></td>
<td>0.68 (0.53–0.88)</td>
<td>0.84 (0.46–1.55)</td>
<td>NC</td>
<td>0.95 (0.88–1.03)</td>
<td>0.90 (0.27–3.00)</td>
</tr>
<tr>
<td>CCB</td>
<td>0.30 (0.21–0.43)</td>
<td>NC</td>
<td>NC</td>
<td>NC</td>
<td>NC</td>
</tr>
<tr>
<td>COX</td>
<td>0.20 (0.03–1.28)</td>
<td>0.41 (0.10–1.66)</td>
<td>NC</td>
<td>0.21 (0.07–0.62)</td>
<td>0.80 (0.25–2.58)</td>
</tr>
<tr>
<td>Mg</td>
<td>0.56 (0.27–1.14)</td>
<td>NC</td>
<td>NC</td>
<td>0.62 (0.46–0.83)</td>
<td>4.56 (1.00–20.86)</td>
</tr>
<tr>
<td>ORA</td>
<td>1.05 (1.15–7.43)</td>
<td>0.74 (0.61–0.91)</td>
<td>1.33 (0.84–2.14)</td>
<td>1.17 (0.99–1.37)</td>
<td>2.25 (0.79–6.30)</td>
</tr>
<tr>
<td>NOD</td>
<td>1.19 (0.74–1.90)</td>
<td>0.93 (0.61–1.41)</td>
<td>NC</td>
<td>0.43 (0.06–2.89)</td>
<td>(n = 186) (NND)</td>
</tr>
<tr>
<td>Progesterone</td>
<td>NC</td>
<td>NC</td>
<td>0.62 (0.30–1.27)</td>
<td>NC</td>
<td>0.62 (0.39–0.98)</td>
</tr>
</tbody>
</table>

#### Comparisons

<table>
<thead>
<tr>
<th>Tocolytics</th>
<th>Primary—single agent vs. placebo</th>
<th>Significant results are shown in bold.</th>
</tr>
</thead>
<tbody>
<tr>
<td>CCB vs. betamimetics</td>
<td>0.86 (0.67–1.10)</td>
<td>0.76 (0.59–0.99)</td>
</tr>
<tr>
<td>COX vs. betamimetics</td>
<td>0.27 (0.08–0.96)</td>
<td>0.88 (0.52–1.46)</td>
</tr>
<tr>
<td>COX vs. CCB</td>
<td>1.08 (0.59–2.01)</td>
<td>0.26 (0.52–1.46)</td>
</tr>
<tr>
<td>COX vs. Mg</td>
<td>0.87 (0.59–1.29)</td>
<td>NC</td>
</tr>
<tr>
<td>Mg vs. betamimetics</td>
<td>0.90 (0.72–1.65)</td>
<td>NC</td>
</tr>
<tr>
<td>Mg vs. CCB</td>
<td>1.19 (0.86–1.65)</td>
<td>NC</td>
</tr>
<tr>
<td>Tocolytics</td>
<td>0.97 (0.77–1.21)</td>
<td>NC</td>
</tr>
<tr>
<td>Mg vs. COX</td>
<td>1.08 (0.91–1.27)</td>
<td>NC</td>
</tr>
<tr>
<td>ORA vs. betamimetics</td>
<td>0.89 (0.66–1.22)</td>
<td>0.91 (0.69–1.20)</td>
</tr>
<tr>
<td>ORA vs. CCB</td>
<td>1.09 (0.44–2.73)</td>
<td>NC</td>
</tr>
<tr>
<td>NOD vs. betamimetics</td>
<td>0.96 (0.87–1.05)</td>
<td>1.03 (0.92–1.15)</td>
</tr>
<tr>
<td>NOD vs. CCB</td>
<td>0.97 (0.77–1.21)</td>
<td>NC</td>
</tr>
</tbody>
</table>

#### Maintenance therapy after treatment of PTL (vs. placebo or no treatment)

<table>
<thead>
<tr>
<th>Tocolytics</th>
<th>Perinatal Mortality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Betamimetics</td>
<td>NC</td>
</tr>
<tr>
<td>CCB</td>
<td>2.41 (0.86–6.74)</td>
</tr>
<tr>
<td>COX</td>
<td>0.46 (0.07–3.00)</td>
</tr>
<tr>
<td>Mg</td>
<td>1.07 (0.88–1.30)</td>
</tr>
<tr>
<td>ORA</td>
<td>NC</td>
</tr>
<tr>
<td>Progesterone</td>
<td>NC</td>
</tr>
<tr>
<td>17P</td>
<td>0.85 (0.47–1.55)</td>
</tr>
<tr>
<td>Vaginal</td>
<td>NC</td>
</tr>
</tbody>
</table>

**Abbreviations:** NC, not calculable from the available reports; NOD, nitric oxide donors; NND, neonatal death; Mg, magnesium sulfate; Rx, therapy; excl. cong. anom, excluding congenital anomalies.

**Note:** Data are shown as relative risk (95% confidence intervals). Significant results are shown in bold.

*Data courtesy of Vincenzo Berghella.*

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**Side effects:**

Material: Hyperglycemia (glucose 140–200 mg/dL in 20%–50%; mechanism: decreased peripheral insulin sensitivity and increased endogenous glucose production); hyperinsulinemia; hypokalemia (K < 3 mEq/L in 50%); tremors,
nervousness, shortness of breath (10%), chest pain (5%–10%), tachycardia/palpitations, arrhythmia (3%); electrocardiogram (EGK) changes (2%–3%); hypotension (2%–3%); pulmonary edema (<1%–5%); mechanism: reduced sodium excretion—sodium and therefore fluid retention). Ritodrine: altered thyroid function, antiarrhythmia.

**Fetal/Neonatal:** Ritodrine: Neonatal tachycardia, hypoglycemia, hypocalcemia, hyperbilirubinemia, hypotension, IVH. Terbutaline: tachycardia, hyperinsulinemia, hyperglycemia, myocardial and septal hypertrophy, myocardial ischemia.

### Calcium Channel Blockers

**Types:** Nifedipine, nicardipine.

**Dose:** Nifedipine 20–30 mg × 1, then 10–20 mg q4–8h (max 90 mg/day) (nicardipine similar dosing).

**Mechanism of action:** Impair calcium channels, so inhibit influx of calcium into cell, and therefore myometrial contraction.

**Evidence for effectiveness** (Table 18.5): Two small RCTs comparing CCB with placebo showed a reduction in PTB <48 hours (RR 0.30, 95% CI 0.21–0.43) and an increase in maternal adverse effects, with insufficient evidence to assess effect on PTB within 7 days, and <37 weeks [38].

When compared with other tocolytic agents [betamimetics, nonsteroidal anti-inflammatory drugs (NSAIDs), glyceryl trinitrate (GTN), magnesium sulfate, and ORAs] CCB increased the interval from initiation of treatment to delivery, increased GA at birth and decreased preterm and very PTB. RDS, NEC, IVH, hyperbilirubinemia and admission to the NICU were also decreased [38]. CCB also reduced the requirement for women to have treatment ceased for adverse drug reaction. There are insufficient data regarding the effects of different dosage regimens and formulations of CCB on maternal and neonatal outcomes; the most studied is nifedipine, at dosage shown above. CCB should be preferred to beta-mimetics for tocolysis.

**Specific contraindications:** Cardiac disease; hypertension (<90/50); concomitant use of magnesium; caution in renal disease.

**Side effects:**

**Maternal:** Flushing, headache, dizziness, nausea, transient hypotension. Caution in women with hypertension and renal disease, as well as women on magnesium (cardiovascular collapse).

**Fetal/Neonatal:** None.

### COX Inhibitors

**Types:** Nonselective COX inhibitors: indomethacin (indocin). Selective COX inhibitors (preferential COX-2 inhibitor): sulindac, rofecoxib (Vioxx), celecoxib, nimesulide.

**Dose:** Indomethacin: 50–100 mg loading dose (rectal or vaginal route preferred, oral otherwise), then 25–50 mg q6h for 48 hours max, and always <32 weeks. Sulindac 200 mg po q12h × 48 hours. Ketorolac: 60 mg IM, then 30 mg IM q6h × 48 hours.

**Mechanism of action:** Inhibit prostaglandin synthesis, therefore inhibit myometrial contraction.

**Evidence for effectiveness** (Table 18.5): The nonselective COX inhibitor, indomethacin, was used in most trials. When compared with placebo, COX inhibition (indomethacin only) results in a 79% reduction in PTB <37 weeks in a small trial, an increase in GA of 3.5 weeks and a >700 g increase in birth weight [39]. No difference was shown in birth within 48 hours of initiation of treatment (RR 0.20, 95% CI 0.03–1.28). No differences were detected in neonatal morbidity or mortality.

**Compared with other betamimetics, COX inhibition resulted in a 47% reduction in PTB <37 weeks’ gestation and a 73% reduction in PTB within 48h [39]. No differences were detected in the fetal or neonatal outcomes such as perinatal mortality, RDS, IVH, NEC, premature closure of the ductus, persistent pulmonary hypertension of the newborn (PPHN). No differences were found when COX inhibitors were compared with magnesium sulfate or CCBS [39].

A comparison of nonselective COX inhibitors versus selective COX-2 inhibitors did not demonstrate any differences in maternal or neonatal outcomes [39]. Due to small numbers, all estimates of effect are imprecise and need to be interpreted with caution.

**Specific contraindications:** Renal or hepatic disease, active peptic ulcer disease, poorly controlled hypertension, NSAID-sensitive asthma, coagulation disorders/thrombocytopenia.

**Side effects:** When used for only 48 hours, no serious maternal and fetal/neonatal side effects occur, and fetal surveillance is not indicated. Usually COX inhibitors are better tolerated by mother than other tocolytics such as magnesium and betamimetics.

**Maternal:** As with any NSAIDs, mild gastrointestinal (GI) upset—nausea, heartburn (take with some food/milk) (COX-1). GI bleeding (COX-1), coagulation, and platelet abnormalities (COX-1), asthma if ASA-sensitive. May obscure elevation in temperature. Long-term rofecoxib (Vioxx) use in adults has been associated with stroke, so this drug is now not available in many countries.

**Fetal/neonatal:** In trials, 403 women received short-term tocolysis (up to 48 hours) with COX inhibitors (mainly indomethacin) and there was only one case of antenatal closure of the ductus arteriosus. There was no increase in the incidence of patent ductus arteriosus (PDA) postnatally (eight treated with COX inhibitors versus eight treated with placebo or other tocolytics) [40]. No difference in incidences of IVH, BPD, PDA, NEC, or perinatal mortality was noted in a review of trials aimed at evaluating safety [41]. Use for >48 hours, especially ≥32 weeks, is associated with significant fetal effects such as constriction of the ductus arteriosus, which can lead to hydrops, pulmonary hypertension and death, and renal insufficiency, manifested in utero by oligohydramnios. Other effects with prolonged use such as hyperbilirubinemia, NEC, IVH have not been shown with <72 use. Selective COX-2 inhibitors have not been shown consistently to be any safer for the fetus/neonate than nonselective COX inhibitors such as indomethacin. Therefore, continuous use of COX inhibitors for >48 hours and ≥32 weeks is contraindicated.

### Magnesium Sulfate (MgSO₄)

**Dose:** 40 g MgSO₄ in 1 L 1/2 normal saline. Initial: 4–6 g/30 minute, then 2–4 g/hour. A dose of 5 g/hour has not been shown beneficial in perinatal outcome compared with a dose of 2 g/hour, and is associated with significant side effects [42]. Weaning MgSO₄ tocolysis has no benefits and a few harmful side effects compared with stopping MgSO₄ abruptly [43].

**Mechanism of action:** Intracellular calcium antagonist.

**Evidence for effectiveness** (Table 18.5): Compared with placebo, there is insufficient evidence to show if MgSO₄ reduces the incidence of PTB or perinatal morbidity and mortality [44].
Compared with all controls (including other tocolytics), MgSO₄ did not prevent PTB at 48 hours, PTB <37 weeks or <32 weeks. Perinatal death was higher (only two perinatal deaths), while perinatal morbidities were similar. Dose of magnesium did not affect efficacy. Given these results, there is no convincing evidence for recommending magnesium for tocolysis [45].

Management: Aim for 4–7 MgSO₄ level. Monitor urinary output. Follow deep tendon reflexes: ↓ at level ≥8, absent ≥10. At ≥10, respiratory depression; at ≥15, risk of cardiac arrest.

Specific contraindications: Myasthenia gravis.

Side effects:
- Maternal: Flushing, lethargy, headache, muscle weakness, diplopia, dry mouth, pulmonary edema (1%; mechanism: intravenous overhydration), cardiac arrest.
- Fetal/neonatal: Lethargy, hypotonia, hypocalcemia, respiratory depression. Prolonged use: demineralization.

Oxytocin Receptor Antagonists

Types: Atosiban (Tractocile); barusiban.

Dose: Atosiban 6.75 mg bolus, then 300 mg/min IV × 3 hours, then 100 mg/min (max 45 hours).

Mechanism of action: Competitive inhibitor of oxytocin via blockade of oxytocin receptor.

Evidence for effectiveness (Table 18.5): Compared with placebo, atosiban does not reduce incidence of PTB or improve neonatal outcome [46]. In one trial, atosiban was associated with an increase in extreme PTB <28 weeks and infant deaths at 12 months of age compared with placebo [46]. However, this trial randomized significantly more women to atosiban before 26 weeks' gestation. Compared with placebo, use of atosiban results in about 140 g lower infant birth weight and milder maternal adverse drug reactions [46]. Compared with betamimetics, atosiban is associated with similar incidences of PTB and perinatal morbidity/mortality, and with fewer maternal drug reactions requiring treatment cessation [46]. There is insufficient evidence to assess the effectiveness of a different ORA, barusiban, as only required treatment cessation [46]. There is insufficient evidence to support the use of oral betamimetics as primary tocolysis as they have less side effects than the (effective) betamimetics.

Side effects: Minimal to none.

Nitric Oxide Donors

Type: Nitroglycerine.

Dose: Nitroglycerine transdermal patch 0.4 mg/hour.

Mechanism of action: Direct relaxation of uterine muscle.

Evidence for effectiveness (Table 18.5): There is currently insufficient evidence to support the routine administration of nitric oxide donors (NOD) for prevention of PTB in women with PTL [48]. Compared with placebo, there was no evidence that NOD prolonged pregnancy beyond 48 hours or improved neonatal outcomes. When compared with other tocolytic agents (betamimetics, magnesium sulfate, CCBS or combination of tocolytics), there was no evidence that NOD perform better than other tocolytics. Nitroglycerine has been the only NOD used in trials.

Specific contraindications: NOD should not be used in women with hypotension or with preload dependent cardiac lesions, such as aortic insufficiency.

Side effects:
- Maternal: NOD cause dilatation of arterial smooth muscle and commonly cause headache and may result in hypotension. Other side effects include dizziness, flushing and palpitations.

Fetal/neonatal: Although maternal hypotension could compromise utero-placental blood flow, no adverse fetal or neonatal effects have been reported.

Progesterone

There is insufficient data to assess the efficacy for progesterone as primary tocolysis. There are some data suggesting that progesterone may reduce PTB and increase birth weight and that progesterone may reduce uterine contractility, prolong pregnancy and attenuate cervical shortening. However, current evidence does not support a role for progesterone as a tocolytic agent [49].

Primary Tocolysis—Multiple Agents Simultaneously

Indomethacin, Ampicillin-Sulbactam, and Magnesium versus Magnesium Alone

Compared with placebos, indomethacin and ampicillin-sulbactam did not prevent PTB in women in PTL already receiving magnesium sulfate tocolysis [50].

Primary Tocolysis—Additional Agents versus One Agent Only

Progesterone versus Placebo, in Addition to Ritodrine

Compared with placebo, in women receiving ritodrine tocolysis, the addition of progesterone was not associated with a significant reduction in PTB <37 weeks (16% vs. 33% in placebo) in a very small trial [51]. In this RCT, 44 women with mostly (>90%) singleton gestations and threatened PTL at less than 35 weeks treated with ritodrine, natural progesterone 400 mg orally q6h × 24 hours (then 400 mg q8h for next 24 hours, and then 300 mg q8h onward) was associated with similar incidence of PTB, with less quantity of ritodrine administered and shorter maternal hospital stay compared with placebo [51].

Refractory Tocolysis—Primary Agent Is Failing

Indomethacin was similar to sulindac in prevention of PTB in women failing primary magnesium sulfate tocolysis in a small trial [52].

Maintenance Tocolysis—After Successful Primary Tocolysis

Betamimetics (Oral)

Dose: Ritodrine: 1–20 mg po q2–4h. Terbutaline: 2.5–5 mg po q2–4h.

Evidence for effectiveness: Compared with placebo, oral betamimetic therapy for maintenance tocolysis does not prevent PTB, recurrent PTL, recurrent hospitalizations, or perinatal morbidity and mortality [53]. Some adverse effects such as tachycardia are more frequent in the betamimetics group. Given this ample evidence from 13 trials, there is absolutely no evidence to support the use of oral betamimetics after PTL has resolved.

Terbutaline Pump

Dose: 0.05 mg/hour.

Evidence for effectiveness: Compared with placebo, terbutaline pump does not prevent PTB or improve perinatal morbidity and mortality. Side effects and costs associated with this therapy further advice against its use [54].
Calcium Channel Blockers
There is insufficient evidence to assess the efficacy of CCB maintenance therapy after successful tocolysis. In three small RCTs, the incidence of PTB <37 weeks was similar to placebo, or no treatment [55].

COX Inhibitors
Compared with placebo, after successful tocolysis, oral sulindac either 200 mg q12h x 7 days or 100 mg q12h until 34 weeks does not reduce PTB compared with placebo [56]. Given the association with fetal/neonatal complications with COX inhibitor use for >48 hours, COX inhibitors should not be used for maintenance tocolysis.

Compared with oral terbutaline, oral indomethacin is associated with a similar incidence of PTB when used for maintenance tocolysis after successful IV tocolysis, but indomethacin is associated with significant constriction of ductus arteriosus and oligohydramnios when used for >48 hours [57]. Therefore, indomethacin should not be used for maintenance tocolysis, especially after 26 weeks.

Magnesium Sulfate
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Compared with placebo, after successful tocolysis, oral sulindac either 200 mg q12h x 7 days or 100 mg q12h until 34 weeks does not reduce PTB compared with placebo [56]. Given the association with fetal/neonatal complications with COX inhibitor use for >48 hours, COX inhibitors should not be used for maintenance tocolysis.

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Magnesium Sulfate
Compared with placebo, or no treatment, magnesium maintenance therapy does not prevent PTB (RR 1.05, 95% CI 0.80–1.40) or affect perinatal morbidity and mortality (mortality, RR 5.00, 95% CI 0.25–99.16) [58]. It has also similar effect on PTB as alternative tocolytic drugs (RR 0.99, 95% CI 0.57–1.72) (i.e., it is equally ineffective). Therefore, magnesium sulfate should not be used as a maintenance tocolytic.

Oxytocin Receptor Antagonists
Compared with placebo, ORA (atosiban) maintenance therapy (30 µg/min) via pump up to 36 weeks does not prevent PTB or affect perinatal morbidity and mortality, with a 5 days (32.6 vs. 27.6, p = .02) longer interval to delivery in one trial [59]. There are no side effects compared with placebo except for injection-site reactions. ORA are not available in oral form for maintenance.

Progesterone
Women with a singleton gestation who received 17P maintenance tocolysis for arrested PTL had a similar rate of PTB <37 weeks (42% vs. 51%; RR 0.78; 95% CI 0.50–1.22) and PTB <34 weeks (25% vs. 34%; RR 0.60; 95% CI 0.28–1.12) compared with controls [62,63]. Women who received 17P had significantly later GA at delivery (MD 2.28 weeks; 95% CI 1.46–13.51), longer latency (MD 8.36 days; 95% CI, 3.20–13.51), and higher birth weight (MD 224 g; 95% CI, 71–378) as compared with controls. As 17P for maintenance tocolysis is associated with a significant prolongation of pregnancy, and significantly higher birth weight [60,61], further research is suggested.

Maintenance tocolysis with vaginal progesterone for arrested PTL is associated with a significantly lower rate of PTB <37 weeks (42% vs. 58%; RR 0.71; 95% CI 0.57–0.90), significantly longer latency (MD 14 days), later GA at delivery (MD 1.3 weeks), lower rate of recurrent PTL (24% vs. 46%; RR 0.51; 95% CI 0.31–0.84), and lower rate of neonatal sepsis (2% vs. 7%; RR 0.34; 95% CI 0.12–0.98) compared with placebo or no treatment [62]. However, due to the poor quality of the trials, maintenance therapy with vaginal progesterone should be studied further before a strong recommendation can be made.

In summary, there is insufficient evidence to recommend progesterone for prevention of PTB in women who remain pregnant after an episode of PTL, but recent evidence about vaginal progesterone for maintenance tocolysis is encouraging.

PTL RESOLVED: HOME VERSUS IN-HOSPITAL CARE
After PTL has resolved (and cervical dilatation has not progressed ≥4 cm), home management is associated with similar incidences of reaching ≥36 weeks compared with hospital management [63,64]. Hospitalization may increase maternal stress, vaginal examinations, time in recumbent position (and its consequences), and decreased plasma volume. For the many women with arrested PTL, continued hospitalization after steroids administration is unnecessary. Women sent home with a diagnosis of false PTL are not at increased risk for early (<34 weeks) PTB or neonatal mortality, but they are at risk for later (≥34 weeks) PTB [1].

PRECONCEPTION COUNSELING
Given its major impact of perinatal morbidity and mortality, it is important to review risk factors for PTB in every pregnant woman. In the woman with a risk factor (e.g., prior PTB), it is important to review prognosis, possible complications, and management of a future pregnancy (see above).

PRENATAL CARE
Preconception counseling as above, if not already done. Management should follow recommendations as above (Table 18.3) (see also Chapter 2).

ANTEPARTUM TESTING
No specific fetal testing is indicated. Home uterine activity monitoring (HUAM), discussed in Chapter 17, is not effective in preventing any complication.

MODE OF DELIVERY
There is insufficient evidence to evaluate the use of a policy for uniform planned cesarean delivery (CD) compared with expectant management and selective CD for preterm (24–36 weeks) infants [65]. Mothers in the planned CD group had more morbidity. There was no significant difference between planned CD and expectant management in perinatal morbidity or mortality or in abnormal childhood follow-up. Differentiation of data between breech and vertex presentations is difficult, with numbers too small for definite conclusions.

Delayed Cord Clamping
In preterm neonates, delayed cord clamping by about 30–60 seconds (120 maximum) is associated with fewer transfusions for anemia, less hypotension, and less IVH than early clamping at less than 30 seconds [66].

Milking of cord has been evaluated in two small RCTs in preterm neonates, so there is insufficient evidence for recommendation, even if it appears as beneficial as delayed cord clamping. Compared with no milking, milking of cord has been associated with less need for blood transfusions and less need for circulatory and respiratory support [67]. Compared with 30-second delayed cord clamping, milking of cord four times achieved a similar amount of placento-fetal blood transfusion [68].

ANALGESIA/ANESTHESIA
No specific changes from term intrapartum management (see Chapter 11).
POSTPARTUM/BREAST-FEEDING/ COUNSELING

As in other pregnancies, breast-feeding is encouraged as tolerated for the preterm infant. Milk expression using breast pump is also encouraged. Extensive counseling should be provided regarding rate of recurrence of PTB, and future management in pregnancy. Treatment with antibiotics before pregnancy does not prevent recurrent PTB. In women with a prior spontaneous PTB <34 weeks, oral azithromycin and metronidazole every 4 months after the PTB and before the next conception does not significantly reduce subsequent PTB [69]. Vaginal and IM progesterone has been shown to reduce the risk of recurrent PTB and adverse perinatal outcomes in women with a history of spontaneous singleton PTB. Following spontaneous singleton PTB, women should be counseled about the benefits of progesterone prophylaxis in subsequent pregnancies [70] (see also Chapter 17).

REFERENCES


29. Gonzalez Garay AG, Reveiz L, Velasco Hidalgo L, Solis Galicia C. Ambroxol for women at risk of preterm birth for preventing


KEY POINTS

• Definite diagnosis is by direct visualization of fluid (pooling), with nitrazine cervicovaginal swab and ferning as usual confirmatory tests. In dubious cases, additional biochemical tests may aid in the diagnosis.

• Complications of preterm premature rupture of membranes (PPROM) include premature labor/delivery with related complications of prematurity such as respiratory distress syndrome (RDS), intraventricular hemorrhage (IVH) and periventricular leukomalacia (PVL), infection and necrotizing enterocolitis (NEC), maternal or neonatal infections (chorioamnionitis, endometritis, and sepsis); abruptio placentae, cord prolapse, and, especially for PPROM <24 weeks, perinatal death, pulmonary hypoplasia (PH), compression deformities, long-term infant morbidities, increased need for cesarean delivery (CD), and retained placenta.

• Corticosteroids should be administered in women with PPROM at 23 0/7–33 6/7 weeks, as this intervention is associated with lower incidences of RDS, IVH, NEC, and a trend for a lower neonatal death rate.

• Antibiotics (in particular ampicillin and erythromycin or erythromycin alone) are associated with less chorioamnionitis, preterm birth (PTB) within 48 hours, PTB within 7 days, neonatal infection, surfactant use, oxygen therapy, and abnormal cerebral ultrasound scan (including IVH). RDS and NEC are also decreased with ampicillin and erythromycin treatment.

• Tocolytic therapy in women with PPROM is not associated with maternal or perinatal benefits and should, in general, be avoided.

• Expectant management of PPROM in women with cerclage in place is associated with increased maternal and fetal/neonatal infection risks. Therefore, the cerclage should, in general, be removed when the diagnosis of PPROM is made. At maximum, cerclage might be left in place for about 48 hours to allow steroid therapy.

• Before 34 weeks, conservative management of PPROM is usually indicated, if possible. Delivery is indicated for NRFHT, preterm labor (PTL), chorioamnionitis, or ≥34 weeks (Figure 19.1). In women with PPROM between 34 and 36 6/7 weeks, in the absence of overt signs of infection or fetal compromise, a policy of cautious expectant management can be considered.

• There are no strong published data to assess the benefit of interventions for PPROM <23 weeks. Series have reported live birth rates from 20% to 90% associated with expectant management, with high rates of perinatal morbidity.

• The management of PPROM in twin pregnancies should not differ from that of singletons.

DEFINITION

PPROM refers to chorioamnionic membrane rupture before the onset of labor in pregnancies at <37 weeks of gestation.

DIAGNOSIS

Definite diagnosis is by direct visualization of fluid (pooling) in the posterior vaginal fornix at sterile speculum examination. History of persistent leakage of fluid and ultrasonographic diagnosis of oligohydramnios are two other confirmatory but not diagnostic findings. Statistical measures (sensitivity, specificity, positive predictive value, negative predictive value) of the main PROM diagnostic tests are reported in Table 19.1 [1–5].

Traditional confirmatory tests:

• Nitrazine test: Evaluation of vaginal pH by a cervical–vaginal swab collected in a sterile way from the posterior vaginal fornix. Vaginal pH generally varies from 4.5 to 6. In presence of amniotic fluid it becomes >7. A nitrazine paper veers from yellow to blue if pH is >7. False positive results in presence of blood, seminal fluid, alkaline antiseptics, cervical–vaginal infections or alkaline urines. False negative results in case of prolonged rupture of membranes.

• Ferning test: The presence of arborization on a slide of amniotic fluid collected in a sterile way from posterior vaginal fornix. Arborization represents the crystallization of amniotic fluid due to its high contents of salts and proteins. False positive results for contamination by cervical–vaginal mucus, seminal fluid, fingerprints, and urine crystals. False negative results for blood or meconium contamination or inadequacy in slide preparation.

Main biochemical tests:

• Placental alpha-microglobulin test (Amnisure™): Placental alpha-microglobulin-1 (PAMG-1) is a glycoprotein of amniotic fluid (2,000–25,000 ng/mL); it can be detected also in lower concentrations in maternal blood (5–25 ng/mL) and in cervical–vaginal secretions (0.05–0.2 ng/mL) if fetal membranes are intact. PAMG-1 is a good marker for PROM diagnosis due to this different concentration between amniotic fluid and cervical–vaginal secretions. PAMG-1 is dosed by an immune-chromatographic test able to detect also little concentrations of this glycoprotein in cervical–vaginal secretions (Amnisure™ PROM test). It is an easy, rapid (5–10 minutes), noninvasive test (speculum examination is not necessary). This test can be performed from 11 to 42 weeks of pregnancy. The result is not influenced by the presence of seminal fluid, urine, blood or vaginal infections.

• Insulin-like growth factor binding protein (IGFBP)-1 (PROM test): Immune-chromatographic test that detects amniotic fluid in the vaginal secretions. Monoclonal antibodies identify IGFBP-1, whose concentration is elevated.
in amniotic fluid (IGFBP-1 ≥10 μg/L: positive test). Actim™ PROM test is easy and rapid. The result is not influenced by the presence of urine or seminal fluid, while it can be altered by contamination with blood.

- **Diagnostic panty liner with polymer-embedded strip:** A pad with a reactive strip put in contact with external genitalia detects the presence of amniotic fluid.

  - **Fetal fibronectin:** This test is sensitive but not specific for PROM: a negative result suggests the absence of PROM with high accuracy, while a positive result is not diagnostic for PROM. Fetal fibronectin test is not recommended in case of PROM. **Transabdominal amniinfusion of dye** (indigo carmine, Evans blue, fluorescein) can be used as a confirmatory test in

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**Figure 19.1** Management of preterm premature rupture of membrane (PPROM). (Adapted from Blumenfeld YJ et al., *Obstet Gynecol*, 116, 1381–1386, 2010.)
doubtful cases at low gestational ages (GAs). Methylene blue must be avoided as it can cause fetal meta-hemoglobinemia [6,7].

**SYMPTOMS**

Over 90% of women with PPROM report a history of “gush of fluid.”

**INCIDENCE**

PPROM occurs in <1% at <24 weeks, and about 1%–3% at 24–33 weeks, 3%–5% at 34–36 weeks compared with about 8%–10% for PROM at term. PPROM is associated with about 25%–30% of all PTBs.

**ETIOLOGY/BASIC PATHOPHYSIOLOGY**

Etiology is complex and multifactorial (see Chapter 17). Possible mechanisms leading to PPROM are choriodecidual infection, collagen degradation, decreased membrane collagen content, localized membrane defects, uterine overdistention, and programmed amniotic cell death [8]. Evidence that supports a causal association between PPROM and infection is vast and includes the fact that microorganisms in the amniotic fluid are more frequently present and the rate of histological chorioamnionitis is higher in PPROM than in intact membranes preterm delivery and the frequency of PPROM is significantly higher in women with lower genital tract infections (e.g., group B streptococcus [GBS] and bacterial vaginosis). Microorganisms that colonize the lower genital tract produce phospholipases, which can stimulate the production of prostaglandins and lead to uterine contractions; the immune response in endocervix and/or fetal membranes leads to the production of multiple inflammatory mediators (particularly matrix metalloproteinases) that can weaken membranes and result in PPROM [9]. Invasive uterine procedures performed during pregnancy (such as amniocentesis, chorionic villus sampling, fetoscopy, and cervical cerclage) can damage the membranes, causing them to leak.

**CLASSIFICATION**

PPROM can be classified into PPROM <23 weeks (usually 16–22 6/7 weeks, and called also preivable or midtrimester or very early PPROM—see also the section “PPROM <23 WEEKS”) and PPROM at 23 0/7–36 6/7 weeks. PPROM at 23–36 weeks can be further subdivided into PPROM at 23–33 6/7 weeks (early PPROM) and PPROM at 34–36 6/7 weeks (late preterm PPROM—for management, see also Chapter 20).

**RISK FACTORS**

Main risk factors for spontaneous PPROM are listed in Table 19.2 [10]. However, most cases of preterm PROM occur in otherwise healthy women without identifiable risk factors. See also Chapter 17.

**COMPLICATIONS**

Complications are inversely correlated with GA at PPROM and at delivery.

- **PTL/delivery:** In 50% of PPROM, labor occurs within 24 hours, and in 80%–90% within 7 days. Median latency to delivery after PPROM is similar from 24 to 28 weeks’ gestation (about 9 days), and shortens with PPROM ≥29 weeks [11]. The latency with PPROM >30 weeks is usually only 2–4 days.

**Preterm delivery** and complications of prematurity are the most important causes of perinatal mortality and morbidity; complications decrease with advancing GA. Infective/inflammatory factors involved in PPROM etiopathogenesis may enhance the risk of fetal white matter damage [12].

- **Most common neonatal morbidities are RDS, IVH, PVL, and NEC.**

- **Infections:** Mother is at risk of chorioamnionitis, endometritis, and sepsis. Serious maternal consequences are uncommon. Mean incidence of chorioamnionitis is about 3%–15%. Major neonatal infections occur in 5% of PPROM and 15%–20% of cases developing chorioamnionitis. Fetal infection can precede clinically evident chorioamnionitis, resulting in neonatal pulmonary and cerebral morbidities.

- **Other complications such as abruptio placentae, cord prolapse, perinatal death, PH, compression syndrome, long-term infant morbidities, increased need for CD, and retained placenta are most common at very early PPROM and are discussed more in detail for PPROM <23 weeks.**

**MANAGEMENT**

**Prevention**

See Chapter 17 (Figure 19.1) [13].

**Preconception Counseling**

Women with prior PPROM have a 20%–30% chance of PTB, including a 15%–20% chance of recurrent PPROM in the next pregnancy. Recurrence is higher in black race. These incidences are inversely related with GA at PPROM (the earlier the GA at PPROM, the higher the recurrence rate) and interpregnancy interval (shorter than 6 months particularly associated with higher recurrence) (see Chapters 1 and 17).

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**Table 19.1 Predictive Accuracy of PROM Diagnostic Tests**

<table>
<thead>
<tr>
<th>Test</th>
<th>Sensitivity (%)</th>
<th>Specificity (%)</th>
<th>Positive Predictive Value (%)</th>
<th>Negative Predictive Value (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nitrazine test</td>
<td>85</td>
<td>39.7</td>
<td>49.3</td>
<td>79.3</td>
</tr>
<tr>
<td>Ferning test</td>
<td>84</td>
<td>78.7</td>
<td>79.7</td>
<td>83.1</td>
</tr>
<tr>
<td>Placental alpha microglobulin test (AmniSure)</td>
<td>92.7</td>
<td>100</td>
<td>100</td>
<td>95.2</td>
</tr>
<tr>
<td>Insulin-like growth factor binding protein-1 (PROM test)</td>
<td>87.5</td>
<td>94.4</td>
<td>92.1</td>
<td>91.1</td>
</tr>
<tr>
<td>Diagnostic panty liner</td>
<td>95.4</td>
<td>80</td>
<td>82.4</td>
<td>94.7</td>
</tr>
<tr>
<td>Fetal fibronectin</td>
<td>94.5</td>
<td>89.1</td>
<td>89.7</td>
<td>94.2</td>
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</tbody>
</table>


Abbreviation: PROM, premature rupture of membrane.
Counseling should also review all risk factors for PTB (Table 19.2). Several risk factors, such as smoking, low body mass index (BMI <20 kg/m²), inadequate maternal weight gain, exposure to infections, illicit drug use, nutritional deficiencies of copper and ascorbic acid, young or advanced maternal age, anemia, occupational factors such as prolonged walking or standing, strenuous working conditions, and long weekly work hours, psychological factors such as depression, anxiety, and chronic stress can be modified and best addressed preconception see also Chapter 17). The woman with a history of PPROM should also be made aware that interventions in pregnancy, such as for example progesterone and cerclage, can further decrease her chance of recurrence (see Chapter 17).

### Prenatal Counseling

Counseling regarding prognosis, possible complications, management, and expectations for neonatal outcome should be provided. Prognosis depends on GA at PPROM and GA at delivery. Latency is inversely related with GA at PPROM (Table 19.3) [14] and can be prolonged by some interventions. Fetal maturity depends on GA and is probably not enhanced or delayed by PPROM.

### Workup

The current gold standard for the diagnosis of PROM is based on three traditional tests: visual pooling of clear fluid in the posterior fornix of the vagina or leakage of fluid from the cervical os at speculum examination; ferning test and nitrazine test tests (Figure 19.1). Such tests become progressively less accurate when more than 1 hour has elapsed after the membranes have ruptured. We still perform these three tests initially, with no need for additional tests in diagnosis in about 99% of women with possible PPROM. Alternative tests have been introduced, which are particularly helpful when traditional tests are indeterminate. See Table 19.1 for accuracy of diagnosis of PPROM with different.

Screening for gonorrhea and chlamydia is indicated, especially in high-risk groups. GBS culture should be sent from anorectal and vaginal areas. Avoid manual/digital examination of the cervix in any woman with suspected PPROM, and also after PPROM is diagnosed by speculum examination. Digital examination is associated with shorter latency and higher incidences of infection [15,16].

Ultrasound should evaluate at least presentation, biometry for GA, anatomy, placenta and cord location, and amniotic fluid. The lower is the amniotic fluid volume (usually measured by amniotic fluid index [AFI] in most studies, but maximum vertical pocket (MVP) is preferred currently), the higher is the incidence of perinatal infection and the shorter is the latency period [17,18].

Fifty to seventy percent of PPROM have low amniotic fluid volume on initial sonography. Low amniotic fluid volume is associated with an increased risk of umbilical cord compression and shorter latency but the predictive value is low [19]. Patients with nonvertex presentations have higher risk for prolapsed umbilical cord and of an unintended vaginal delivery when compared with vertex presentations [20].

See also Chapter 17.

### Infection Precautions

Women with active herpes simplex virus (HSV) infection or human immunodeficiency virus (HIV) with viral loads >1000

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### Table 19.2 Selected Risk Factors for Spontaneous PPROM

<table>
<thead>
<tr>
<th>Maternal Factors</th>
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<tbody>
<tr>
<td>• PPROM in a prior pregnancy (recurrence risk is 16%–32% as compared with 4% in women with a prior uncomplicated term delivery)</td>
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<tr>
<td>• Antepartum vaginal bleeding</td>
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<td>• Chronic steroid therapy</td>
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<td>• Collagen vascular disorders (such as Ehlers–Danlos syndrome, systemic lupus erythematosus)</td>
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<td>• Direct abdominal trauma</td>
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<td>• Preterm labor</td>
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<td>• Exposure to infections</td>
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<tr>
<td>• Cigarette smoking</td>
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<tr>
<td>• Illicit drugs (cocaïne)</td>
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<tr>
<td>• Anemia</td>
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<tr>
<td>• Low body mass index (&lt;20 kg/m²)</td>
<td></td>
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<tr>
<td>• Inadequate maternal weight gain</td>
<td></td>
</tr>
<tr>
<td>• Nutritional deficiencies of copper and ascorbic acid</td>
<td></td>
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<tr>
<td>• Low socioeconomic status</td>
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</tr>
<tr>
<td>• Unmarried status</td>
<td></td>
</tr>
<tr>
<td>• Young or advanced maternal age</td>
<td></td>
</tr>
<tr>
<td>• Occupational factors (strenuous working conditions, long weekly work hours)</td>
<td></td>
</tr>
<tr>
<td>• Psychological factors</td>
<td></td>
</tr>
</tbody>
</table>

### Uteroplacental Factors

- Uterine anomalies (such as uterine septum)
- Placental abruption (may account for 10%–15% of preterm PROM)
- Advanced cervical dilatation (cervical insufficiency)
- Prior cervical conization
- Cervical shortening in the second trimester (<2.5 cm)
- Uterine overdistension (polyhydramnios, multiple pregnancy)
- Intra-amniotic infection (chorioamnionitis)
- Multiple bimanual vaginal examinations (but not sterile speculum or transvaginal ultrasound examinations)

### Fetal Factors

- Multiple pregnancy (preterm PROM complicates 7%–10% of twin pregnancies)
- Fetal anomalies (malformations, aneuploidies)

**Source:** Adapted from Caughey AB et al., *Rev Obstet Gynecol*, 1, 11–22, 2008.

### Table 19.3 Latency Depending on GA at PPROM

<table>
<thead>
<tr>
<th>GA at PPROM (weeks)</th>
<th>Mean Latency</th>
<th>Delivery</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;24</td>
<td>7 days</td>
<td>&lt;48 Hours</td>
</tr>
<tr>
<td>24–33 6/7</td>
<td>3–6 days</td>
<td>20%</td>
</tr>
<tr>
<td>34–36 6/7</td>
<td>24 hours</td>
<td>50%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>70%–80%</td>
</tr>
</tbody>
</table>

**Source:** Adapted from Schucker JL and Mercer BM, *Semin Perinatol*, 20, 389–400, 1996.

**Abbreviation:** GA, gestational age.
are not, in general, expectantly managed, especially if PPROM has occurred after 32 weeks (see Chapters 32 and 50 in Maternal-Fetal Evidence Based Guidelines). Management needs to be individualized with PPROM 23–31 6/7 weeks in the presence of these infections.

**Amniocentesis**

Amniocentesis can be of use for the evaluation of the following:

- **The diagnosis.** If diagnosis is in doubt, 1 mL of indigo carmine (not currently available in the United States; alternatives are Evans blue or fluorescein) in 9 mL of normal saline can be injected into the amniotic cavity under continuous ultrasound guidance. Presence of blue on a pad worn on the perineum for 2–4 hours confirms the diagnosis.
- **The infectious state of the amniotic cavity.** Send amniotic fluid glucose (<15 mg/dL associated with positive culture), Gram stain, and culture.
- **The fetal maturity.** Results of similar accuracy to amniocentesis can be obtained noninvasively by collecting vaginal fluid using a bedpan [21]. Despite oligohydramnios, there is a 90% rate of success for transabdominal amniocentesis in PROM. There are insufficient data to assess the effect of amniocentesis on outcomes in PPROM. A recent Cochrane review [22] reported no benefits for amniocentesis on perinatal outcomes in a single randomized study with 47 patients.

**Fetal Maturity Assessment**

Assessment of fetal maturity can be obtained from amniotic fluid by amniocentesis or from the vaginal pool (see Chapter 58 in Maternal-Fetal Medicine Evidence Based Guidelines). The predictive value of lung maturity tests is not modified by PPROM. Phosphatidylglycerol (PG), surfactant/albumin ratio (TDx/FLM), and lamellar body counts (LBC) are accurate when tested in the vaginal pool. PG is not accurate in the presence of meconium or blood, LBC is not accurate in the presence of meconium, while the TDx/FLM is accurate with blood and/or meconium in the vaginal pool, yielding results similar to those observed with samples obtained with amniocentesis.

**Meconium**

Meconium-stained amniotic fluid is associated with clinical chorioamnionitis and positive amniotic fluid cultures. In the absence of symptoms and signs of chorioamnionitis, meconium alone is not an indication for intervention.

**Hospitalization**

There is insufficient evidence to compare hospital versus home management for PPROM. A Cochrane review [23] including only two small trials shows no significant differences between home management and hospitalization in pPROM. Home management can be offered only to consenting, reliable patients with the following: absence of infection, dependable transportation, living near hospital, evaluation in hospital before discharge, vertex presentation, vertical pocket of amniotic fluid >2 cm, recording of temperature and pulse every 6 hours, fetal movements count, twice-weekly non-stress tests (NST), and weekly ultrasound. Only 18% of patients with PPROM meet these criteria, so that most are managed in the hospital. Eleven percent of women with PPROM managed at home delivered unexpectedly at outside hospitals [24]. Women were monitored for 48–72 hours before randomization to hospitalization versus no hospitalization in the two randomized controlled trials (RCTs) on this subject [23]. Compared with hospitalization, no hospitalization was associated with similar incidences of perinatal mortality (relative risk [RR] 1.93, 95% confidence interval [CI] 0.19–20.05), serious neonatal morbidity, chorioamnionitis, GA at delivery, birth weight and admission to neonatal intensive care unit, as well as CD (RR 0.28, 95% CI 0.07–1.15). There was no information on serious maternal morbidity or mortality. Mothers randomized to care at home spent approximately 10 fewer days as inpatients and were more satisfied with their care. Furthermore, home care was associated with reduced costs [23].

**Maternal Surveillance**

All women with PPROM should be monitored for signs of infection by assessment of clinical parameters (e.g., fever, maternal/fetal tachycardia, uterine tenderness, and purulent vaginal discharge). A diagnosis of chorioamnionitis is usually made by the presence of two or more of these criteria. The presence of a fever of unknown origin in the presence of PPROM is highly suspicious for chorioamnionitis, so that an amniocentesis should be considered if expectant management is still being considered (Chapter 22).

There is no clear evidence to support the use of C-reactive protein (CRP) for the early diagnosis of chorioamnionitis (specificity 38%–55%). CRP has also a low sensitivity to identify intrauterine infection [25,26]. Maternal leukocytosis at admission is associated with higher adverse infant neurodevelopmental outcomes at 2 years of age in a study [27].

**Fetal Surveillance (Antepartum Testing)**

The two most common types of fetal surveillance are NST and biophysical profile score (BPS) [28]. Abnormalities of these tests can be somewhat predictive of fetal infection and umbilical cord compression related to oligohydramnios. There is insufficient evidence to assess the optimal type or frequency of testing. The NST or BPS performed daily have poor sensitivity (39% and 25%, respectively) and similar predictive values for predicting infection [29]. No improvement in perinatal outcome has been reported in one trial [30]. Given lower cost, the NST is usually suggested for daily to twice-a-week fetal surveillance. Monitoring may be more frequent with oligohydramnios, because it is associated with an increased risk of umbilical cord compression and shorter latency, with a preference for BPS as a backup if the NST is nonreassuring [19]. One non-RCT study supports continuous fetal monitoring (CFM) in the management of PPROM [31], but there is insufficient evidence for a recommendation, and most practitioners and patients adopt intermittent (e.g., 1 hour every 8 hours) fetal monitoring for inpatient management of PPROM. Moreover, continuous prolonged external fetal monitoring may not be practically feasible [32].

**Amniinfusion for Prolonging Latency**

Transabdominal amniinfusion is associated with a reduction in neonatal death (RR 0.30), neonatal sepsis (RR 0.26), PH (RR 0.22), puerperal sepsis (RR 0.20) and less likelihood to deliver within seven days of membrane rupture (RR 0.18) [33]. These results are encouraging but limited by the sparse data and low methodological robustness. Transcervical amniinfusion to prevent NRFHT is discussed below, under section “Delivery.”

**Corticosteroids for Fetal/Neonatal Maturation and Benefit**

A single course of corticosteroids is recommended for pregnant women between 24 0/7 weeks and 33 6/7 weeks of gestation.
who are at risk of PTB [34]. Antenatal corticosteroid therapy in case of PPROM is associated with lower combined fetal and neonatal death (RR 0.62, 95% CI 0.46–0.82), RDS (RR 0.67, 95% CI 0.55–0.82), cerebroventricular hemorrhage (RR 0.47, 95% CI 0.28–0.79), and NEC (RR 0.39, 95% CI 0.18–0.86). There are no statistically significant differences for maternal death, chorioamnionitis, puerperal sepsis or neonatal infection with or without antenatal corticosteroids administration in the setting of PPROM [35]. A course of antenatal steroid therapy (24 mg of betamethasone—12 mg intramuscular (IM) every 24 hours—or 24 mg of dexamethasone—6 mg IM every 12 hours) should not be repeated in patients with PPROM, since weekly courses do improve severe RDS, resulting in less composite neonatal morbidity among neonates delivered at 24–27 weeks, but are associated with shorter latency, higher risks of chorioamnionitis and neonatal sepsis, and no improvement in overall composite neonatal morbidity [36,37].

A single rescue course of antenatal corticosteroids may be considered if the antecedent treatment was given more than 2 weeks prior, the GA is less than 32/6/7 weeks, and the women are judged to be likely to give birth within the next week [38,39]. Prior to 26 weeks’ gestation there is paucity of data on the efficacy of the current dosing regimens of corticosteroids in women with PPROM; moreover at low GAs there are only a few primitive alveoli on which the drug can exert an effect to improve lung function. A recent review does not support or refute the administration of antenatal corticosteroids to women at risk of PTB <26 weeks’ gestation [40]. If the initial course was given at less than 26 weeks of gestation a single repeated course may be considered if the delivery is expected within 1 week [41]. A single rescue corticosteroid course in PPROM is not associated with an increased rate of neonatal sepsis [42], maternal chorioamnionitis or neonatal morbidity [43] (see also Chapter 17).

**Antibiotics for Prolongation of Latency and Fetal/Neonatal Benefit**

Compared with placebo, antibiotics for women with PPROM are associated with short-term benefits for both women and neonates and should be routinely given [44,45]. Benefits of maternal antibiotic therapy when there is PPROM are as follows:

- **Maternal:** 44% less chorioamnionitis
- **Fetal/Neonatal:**
  - 29% reduction in PTB within 48 hours
  - 21% reduction in PTB within 7 days
  - 33% reduction in neonatal infection

- 21% reduction in positive neonatal blood culture
- 17% reduction in use of surfactant
- 12% reduction in oxygen therapy
- 19% reduction in abnormal cerebral ultrasound scan (including IVH) prior to discharge from hospital
- A 10% prolongation of pregnancy
- Decreasing trend in perinatal mortality (RR 0.93, 95% CI 0.76–1.14)
- Decrease in RDS and NEC with ampicillin and erythromycin treatment [46]

A meta-analysis limited to PPROM before 34 weeks shows similar results [47]. The ORACLE study evaluated the children’s health at 7 years of age and found no difference in any functional impairment after prescription of erythromycin, with or without co-amoxiclav, compared with those born to mothers who received no erythromycin, or after prescription of coamoxiclav, with or without erythromycin, compared with those born to mothers who received no co-amoxiclav. Long-term adverse effects of antepartum prophylactic antibiotics for PPROM have not been observed in children followed to age 7 years [48]. This finding is in contrast to the observation from the same authors that in patients with spontaneous PTB and intact membranes, the rate of cerebral palsy was increased in children exposed to antibiotics in utero [49]. These results would suggest further caution should be used when considering the routine treatment of women with antibiotics if there is uncertainty about the diagnosis of PROM.

Benefits in short-term outcomes (prolongation of pregnancy, infection, need for respiratory therapy, less abnormal cerebral ultrasound before discharge from hospital, etc.) should be balanced against a lack of evidence for benefit for other outcomes, including perinatal mortality, and for long-term outcomes.

**Type**

There is insufficient evidence on the optimal antibiotic type (and regimen) in women with PPROM. Ampicillin and erythromycin [46] or erythromycin alone [44] are associated with significant benefits in neonatal outcomes and should be used routinely in women with PPROM 24–34 weeks [34,44,46,47,50–52] (Table 19.4). A combination of ampicillin and erythromycin—for example, ampicillin 2 g and erythromycin 250 mg both intravenously (IV) every 6 hours for 48 hours, followed by amoxicillin 250 mg and erythromycin base 333 mg both orally (PO) every 8 hours for 5 days, for a total of 7 days—in women with PPROM with no concomitant steroids use showed an improvement in

### Table 19.4 Possible Antibiotic Regimens

<table>
<thead>
<tr>
<th>Reference</th>
<th>Antibiotic</th>
<th>Dose</th>
</tr>
</thead>
<tbody>
<tr>
<td>34, 46</td>
<td>Ampicillin</td>
<td>2 g IV every 6 hours and 250 mg IV every 6 hours for 48 hours followed by 250 mg PO every 8 hours and 333 mg PO every 8 hours for 5 days.</td>
</tr>
<tr>
<td>44, 52</td>
<td>Erythromycin</td>
<td>250 mg PO every 6 hours for a maximum of 10 days</td>
</tr>
<tr>
<td>50</td>
<td>Cefazolin</td>
<td>1 g IV every 6 hours and 250 mg PO every 6 hours for 7 days</td>
</tr>
<tr>
<td>51</td>
<td>Ampicillin</td>
<td>500 mg PO single dose every 24 hours for 3 days followed by 500 mg PO every 6 hours for 4 days</td>
</tr>
<tr>
<td>51</td>
<td>Azithromycin</td>
<td>500 mg PO every 6 hours for 4 days</td>
</tr>
</tbody>
</table>


**Abbreviations:** IV, intravenous; PO, by mouth.
neonatal health by significantly reducing the rates of infants with one or more major infant morbidity (composite morbidity: death, RDS, early sepsis, severe IVH, and severe NEC) from 53% to 44% [45]. Compared with placebo, erythromycin 250 mg PO four times a day for 10 days is associated with decreases in neonatal death, chronic lung disease, major cerebral abnormalities, and prolongation of pregnancy in women with PPROM <37 weeks who received steroids in >75% of cases [44].

Azithromycin can be used instead of erythromycin, but there are less data on its efficacy in women with PPROM. A retrospective comparison between azithromycin and erythromycin in addition to ampicillin in preterm PROM shows no differences in latency or maternal or fetal outcomes [51]. Cephalosporins are also commonly used in pregnancy, and have few side effects and a broad-spectrum antibacterial effect [53]. A randomized trial comparing cefazolin plus macrolide (erythromycin or clarithromycin) versus cefazolin alone in PPROM showed no difference among the three antibiotic regimens in term of newborn outcome [50]. In a large trial, amoxicillin/clavulanate was associated with an increased risk of neonatal NEC, although there is no consistent trend toward a positive or negative effect of broad-spectrum antibiotics for NEC in the literature [28,44]. According to Cochrane review, co-amoxiclav should be avoided in women at risk of PTB due to increased risk of neonatal NEC (RR 4.72, 95% CI 1.57–14.23). Possible antibiotic regimens are listed in Table 19.4.

The mechanism of action of single antibiotic drugs should also be considered when selecting the best antibiotic regimens. Macrolides may be considered a good choice as they diffuse slowly in the tissues and act mainly on gram + and chlamydia; moreover, the microorganism disruption in the phagocyte prevents prostaglandins release and the consequent inflammatory cascade activation.

Detection of specific cervicovaginal pathogens should be appropriately treated (see Chapters 32 through 37 of Maternal-Fetal Evidence Based Guidelines) but the routine cervicovaginal swab analysis is not indicated.

Clinical chorioamnionitis requires therapeutic antibiotics. For example, ampicillin (2 g IV every 6 hours) or cefazolin (1 g IV every 6 hours), plus gentamicin (3–5 mg/kg single dose IV every 24 hours). If CD is performed, prophylactic antibiotics are indicated (see Chapter 13). Intrapartum GBS prophylaxis should be given until culture results are available, and to carriers. There are insufficient data to assess the need for this intervention in women with PPROM in whom GBS is sensitive to antibiotics—like ampicillin—already given for PPROM. The suggested regimen is penicillin (5 million units IV, and then 2.5 million units every 4 hours), or (if unavailable) ampicillin (2 g IV, then 1 g every 4 hours).

**Tocolysis for Prolongation of Latency and Fetal/Neonatal Benefit**

In a Cochrane review tocolysis was associated with longer latency (mean difference 73.12 hours; 95% CI 20.21–126.03) and fewer births within 48 hours (average RR 0.55; 95% CI 0.32–0.95). However, tocolysis was associated with increased incidence of 5-minute Apgar of less than seven (RR 6.05; 95% CI 1.65–22.23) and increased need for ventilation of the neonate (RR 2.46; 95% CI 1.14–5.34), while it was not associated with a significant effect on perinatal mortality. For women with PPROM before 34 weeks, there was a significantly increased risk of chorioamnionitis; neonatal outcomes were not significantly different. There were no significant differences in maternal/neonatal outcomes in subgroup analyses of different classes of tocolytics, antibiotic, corticosteroid or combined antibiotic/corticosteroid.

The review suggests there is insufficient evidence to support tocolytic therapy for women with PPROM, as there was an increase in maternal chorioamnionitis without significant benefits to the infant. However, studies did not consistently administer antibiotics and corticosteroids, both of which are now considered standard of care [54]. Compared with short-term tocolysis to allow steroid effect, long-term tocolysis >48 hours is associated with a nonsignificant prolongation of pregnancy, no differences in neonatal complications, but increases in chorioamnionitis and postpartum endometritis. It should therefore be avoided [55].

**Magnesium Sulfate for Fetal Neuroprotection**

A protective role of magnesium sulfate against the risk of cerebral palsy in preterm infants born before 32 weeks has been demonstrated by randomized trials (cumulative absolute risk reduction of 1.7) [56–60]. Results in PPROM cases were not reported separately in any study but constituted almost the 90% of the population in the largest study [60]. In the absence of evidence for an optimal dose of magnesium sulfate, scientific societies suggest adopting the minimum dosage used in the published trials and when delivery appears imminent at <32 weeks [61–63]. According to our Institution’s protocol, we administer magnesium sulfate 4 g load over 20 minutes, followed by 1 g/hour for a maximum duration of treatment of 12 hours (see Chapter 18).

**Progesterone**

17 alpha-hydroxyprogesterone caproate (17P) is not associated with increased latency versus placebo in women with PPROM [64]. In women with singleton gestations and PPROM at 24–30 weeks, 17P 250 mg IM is associated with no effect on interval to delivery, GA at delivery, or neonatal mortality and morbidity in a small RCT [65]. In a recent RCT, weekly 17OHP-C injections did not prolong pregnancy or reduce perinatal morbidity in patients with PPROM [66]. In summary, there is insufficient evidence to recommend the use of progesterone in women with PPROM. In a woman who has been receiving 17P for prior spontaneous PTB, in the absence of evidence to the contrary, it is reasonable to continue 17P once membranes have ruptured [67].

**Vitamin C and E**

There is insufficient evidence to assess the effect of vitamin supplementation in women with PPROM. In one small trial, compared with placebo, vitamin C 500 mg and vitamin E 400 IU daily in women with PPROM at 26–34 weeks were associated with 7-day prolongation in latency, but no other effects on maternal or neonatal morbidity and mortality [68]. In a recent RCT, the use of vitamins C 1000 mg and E 400 IU in women with PPROM was associated with a longer latency period before delivery. Adverse neonatal and maternal outcomes, which are often associated with prolonged latency periods, were similar between the groups [69].

**Cerclage Removal**

PPROM occurs in about 38% of women with cerclage in place [70]. The benefit of a retained cerclage to prolong latency and decrease complications related to prematurity has to be weighed against the risk of adverse maternal and neonatal outcomes. Compared with immediate cerclage removal, leaving the cerclage in place in women with PPROM is associated with longer latency between PPROM and delivery, but higher incidence of maternal (chorioamnionitis, endometritis) and neonatal (sepsis) infections [71,72]. Leaving the cerclage in place >24 hours is associated with a longer latency >48 hours (94% vs. 51%; OR
16.1, 95% CI 3.7–71.3), but also significantly increased maternal and fetal/neonatal infection risks, such as chorioamnionitis (43% vs. 20%; OR 2.9, 95% CI 1.7–5.0) and neonatal mortality from sepsis (12% vs. 1%; OR 13.2, 95% CI 1.6–108.3), and a trend for more neonatal sepsis (13% vs. 6%; OR 2.4, 95% CI 0.9–6.0) and neonatal mortality (17% vs. 10%; OR 2.0, 95% CI 0.9–4.3) [39,42–44,70]. A recent small randomized trial, terminated after an interim analysis, showed no differences between the two management groups in terms of latency, infection and composite neonatal outcomes. However, there was a numerical trend in the direction of less infectious morbidity with immediate removal of cerclage [73]. In a recent meta-analysis, cerclage retention after PPROM did not significantly prolong the gestational latency period, but it increased the rates of delivery after the first 48 hours. It did not significantly increase the rates of neonatal sepsis or neonatal death. Maternal chorioamnionitis was more prevalent among women with cerclage retention (OR 1.78) [74]. Therefore, in most cases, cerclage should be removed in women upon diagnosis of PPROM.

Delivery

Timing

Once PPROM occurs, and steroids for fetal maturity have been administered, the options for management include early delivery versus expectant management until later in pregnancy. In the absence of clear indications for expedient delivery, such as nonreassuring fetal status, clinical chorioamnionitis, and abruptio placenta, the optimal GA for delivery is less clear. A longer latency is associated with higher rate of chorioamnionitis, and a shorter latency with higher risk of severe prematurity. Delivery before 34 weeks is associated with risk of neonatal complications, including severe morbidity and death.

A meta-analysis of seven RCTs, including 690 women, concluded there was insufficient evidence to guide clinical practice regarding the risks and benefits of expectant management versus delivery in PPROM. In particular early delivery is associated with similar incidence of intratravine deaths (RR 0.26, 95% CI 0.04–1.52), neonatal sepsis (RR 1.33, 95% CI 0.72–2.47), RDS (RR 0.98, 95% CI 0.74–1.29), cerebroventricular hemorrhage (RR 1.90 95% CI 0.52–6.92), NEC (RR 0.58, 95% CI 0.08–4.08), duration of neonatal hospitalization, neonatal deaths (RR 1.59, 95% CI 0.61–4.16), and perinatal mortality (RR 0.98, 95% CI 0.41–2.36), compared with expectant management. Regarding maternal outcomes, early delivery is associated with an increase in the incidence of CD (RR 1.51, 95% CI 1.08–2.10), endometritis (RR 2.32, 95% CI 1.33–4.07), and about 1 day less in duration of maternal hospital stay, but no effect on chorioamnionitis (RR 0.44, 95% CI 0.17–1.14) [75]. Unfortunately, most trials of this meta-analysis did not use steroids for fetal maturity, or antibiotics for infection prophylaxis [76–79]. Because steroids and antibiotics are associated with major perinatal benefits, these RCTs therefore have limited clinical validity. According to most authorities based on the data reviewed above, women with PROM before 34/07 weeks should be managed expectantly if no maternal or fetal contraindications exist [34].

Recently PPROMEXIL 1–2 trials have compared immediate delivery after PPROM in near term (34–37 weeks) gestation versus expectant management. The studies conclude that the risk of neonatal sepsis after PPROM near term is low and induction of labor does not reduce this risk [80,81]. A recent secondary analysis of PPROMEXIL trials, with early onset neonatal sepsis as main outcome measure, suggests that women with PROM between 34 and 37 weeks might benefit from immediate delivery if they have GBS vaginal colonization, while in GBS-negative women labor induction could be delayed until 37 weeks [82]. Another recent RCT (PPROMT trial) has compared immediate delivery with expectant management after PPROM close to term (34–37 weeks) in the absence of overt signs of infection. The primary outcome, neonatal sepsis, did not differ between the study groups, just like the composite secondary outcome of neonatal morbidity and mortality. However, neonates born to mothers in the immediate delivery group had increased rates of respiratory distress (RR 1.6) and any mechanical ventilation (RR 1.4) and spent more time in intensive care (median 4.0 days). The study concludes that, in the absence of overt signs of infection or fetal compromise, a policy of expectant management with appropriate surveillance of maternal and fetal wellbeing should be followed in pregnant women who present with ruptured membranes close to term [83].

At >34/07 weeks, American College of Obstetricians and Gynecologists (ACOG) recommendations still suggest delivery in women with PPROM [34], but recent evidence suggest a cautious expectant management policy even >34/07 weeks as a viable alternative, especially if GBS negative.

Ammioinfusion for Preventing NRFHT

There is insufficient evidence to assess the effect of routine amnioinfusion in all women with PPROM, for prophylaxis either before labor or during labor. Compared with no amnioinfusion, transcervical amnioinfusion (warmed saline at 10 mL/minute for 1 hour, then 3 mL/minute—total volume infused mean 1160 mL) at the time of labor for women with PPROM at 26–35 weeks is associated with statistically similar but more favorable incidences of CD, low Apgar scores, and neonatal death in one small trial [84]. In the amnioinfusion group, the number of severe fetal heart rate decelerations per hour during the first stage of labor was reduced by just about one in this small trial. Fetal umbilical artery pH at delivery was higher. These outcomes are consistent with the benefits found for amnioinfusion for cord compression. Transcervical amnioinfusion reduced persistent variable decelerations during labor (RR 0.52) also in a trial with 86 participants [85].

In conclusion, to date, routine intrapartum amnioinfusion cannot be recommended for all women with PPROM given the limited data (see Chapter 10).

Mode and Site of Delivery

Mode of delivery should not be altered solely for the presence of PPROM. Malpresentation is more common with PTB, with the incidence of malpresentation being indirectly related to weeks of GA (see Chapter 22). Early preterm delivery should occur in a facility and by personnel capable of providing all the necessary support to the mother and fetus/neonate born preterm. Transferring women at risk of very PTB to perinatal centers for delivery decreases the neonatal mortality and morbidity rate considerably [86,87]. Predicting the time of delivery in PPROM is often impossible.

Anesthesia

No specific precautions.

Postpartum

See Chapter 17. Women with PPROM are at high risk of recurrence in subsequent pregnancies, with an association between GA at the time of PPROM, latency period, interval between pregnancies, and PPROM recurrence. Patients with a history of cervical insufficiency and fetal loss/miscarriage should be
considered at increased risk of midtrimester PPROM in the following pregnancy. See the section “Prevention.”

Special considerations must be given when PPROM occurs before fetal viability, after invasive procedures, and in twin gestation.

**PPROM <23 WEEKS**

**Definition**

PROM <23 weeks or “previable PPROM” (arbitrary definition that varies among investigators and from year to year because of advances in neonatal intensive care) is a relevant and unique nosological entity because of its significant association with fetal and neonatal morbidity and mortality.

**Incidence**

The incidence is about 0.6% of pregnancies.

**Etiology/Basic Pathophysiology**

There are two different categories: spontaneous and iatrogenic. Risk factors for spontaneous PPROM <23 weeks are similar to those for PTL and for PPROM later in pregnancy (see Table 19.2 and 17.1 in Chapter 17). Fluid leakage or PPROM occurs in about 1% of genetic amniocenteses [88], 3%–5% of diagnostic fetoscopies, and 10% of invasive fetoscopies [89], see below.

**Complications**

Incidences of most complications are inversely proportional to GA at PPROM, latency, residual amniotic fluid volume, and GA at delivery.

**Fetal/Neonatal**

*Neonatal death.* Published data on neonatal survival after early PPROM varies widely (20%–68%) [90–94]. The mortality rates may be underestimated by selection bias due to the high rate of elective termination prior to viability and consequent inclusion of only patients continuing pregnancy or who have experienced initial latency. A study was performed to investigate if, in settings where elective termination of pregnancy (TOP) is frequent, perinatal outcomes could be better (assuming that patients with a poorer prognosis opted more frequently for TOP). Perinatal outcomes (mean GA at delivery, latency, birthweight and survival) were better in the centers with lower TOP rate; in particular survival among live births was 95.8% in the center with lower TOP and 65% in center with higher TOP [93]. A retrospective study on 58 women with prolonged PPROM at less than 24 weeks reported a survival rate in newborns of 90%. Pulmonary morbidities were common in the neonates [95]. The most recent reports on perinatal survival in PROM <24 weeks involve patients routinely managed with antibiotics, antenatal steroids, postnatal surfactant, and high-frequency ventilation. A comparison between PPROM less than 25.0 weeks and 25.0–31.9 weeks showed higher rate of severe composite neonatal morbidity and composite severe childhood morbidity in case of early PPROM. Neonatal death occurred in nearly 17% of early PPROM [94]. Neonatal mortality is comparable to that in preterm deliveries matched for GA without PPROM.

*Chorioamnionitis.* Antenatal infection is the major complication limiting the latency interval. If clinical infection occurs at any time during the latency period, delivery is indicated. Chorioamnionitis complicates about 40% of midtrimester PPROM [96,97]. The occurrence of chorioamnionitis is higher early in the latency period; more than 50% of cases occur within the first 7 days after rupture, with the maximum clinical occurrence on 2–5 days [98,99]. After the first week of latency, the incidence falls, suggesting that subclinical uterine or chorioamniotic infection that weakens membranes and causes rupture infection was probably present prior to membrane rupture, whereas bacteria migration is a less important component. The risk of chorioamnionitis is inversely proportional to residual amniotic fluid volume [100].

*Placental abruption.* Abruptio placentae is more frequent in pregnancies with midtrimester PPROM, occurring up to 44% of cases compared with 0.8% of the general obstetric population [101]. The risk is highest with lower GA at PPROM and with vaginal bleeding occurring prior to or after membrane rupture [101,102].

*Cord prolapse.* The incidence of cord prolapsed is about 2%. The risk is higher (11% in one study) in the setting of nonvertex fetal presentations [20].

<table>
<thead>
<tr>
<th>Gestational age (weeks)</th>
<th>Stage of lung development</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>4 to 6</td>
<td>EMBRIONIC</td>
<td>Lung bud arises</td>
</tr>
<tr>
<td>7 to 16</td>
<td>PSEUDOGLANDULAR</td>
<td>Non respiratory bronchi and bronchioles develop</td>
</tr>
<tr>
<td>17 to 24</td>
<td>CANALICULAR</td>
<td>First gas exchanging acini and pulmonary capillaries are forming</td>
</tr>
<tr>
<td>25 to 37</td>
<td>TERMINAL SAC</td>
<td>Subsaccules and alveoli develop with extensive capillary invasion and expansion of the alveolar blood barrier surface area</td>
</tr>
<tr>
<td>38 to age 3 years</td>
<td>ALVEOLAR</td>
<td>Subsaccules become alveoli</td>
</tr>
</tbody>
</table>

Figure 19.2  Phases of lung development.
Fetal death. Fetal demise is primarily related to abruption, cord prolapse and compression, or infection. The average risk of fetal death after midtrimester PPROM is about 10%, and is inversely related to GA at PPROM and residual amniotic fluid volume [102].

Pulmonary hypoplasia. PH is a decrease in the number of lung cells, airways, and alveoli, mainly due in PPROM to alterations of normal amniotic fluid pressure and egress of lung fluid during the canalicular stage of lung development (ending at nearly 24 weeks) (Figure 19.2). The gold standard for the diagnosis of PH is lung weight by autopsy. The incidence of PH resulting from midtrimester PPROM varies from 13% to 28% [102–106]. The mortality rate in neonates with this condition is 50% to 95% [84, 88, 89]. The main independent reported risk factors for development of PH are as follows: (a) early GA at membrane rupture [103, 105, 106–109]; a risk of 50%–60% was reported when PPROM occurs at 20 weeks or less; and (b) low residual amniotic fluid volume [89,110,104,106]. PH is more common among pregnancies with GA at rupture [106,108]. Incidences of PH with severe, moderate, and absent-to-mild oligohydramnios are 43%, 21%, and 7%, respectively [103].

Tests proposed to identify PH prenatally are ultrasound measurement of amniotic fluid, fetal breathing movements, fetal chest circumference, lung length, lung volume, Doppler studies of pulmonary vessels, and O2 tests. In general, the predictive accuracy is poor and may be improved by combining tests [105,109,111].

Fetal compression syndrome. In early PPROM, asymmetric intrauterine pressure and restriction in fetal movement can lead to limb position deformities and craniofacial defects of variable severity, originally described in the context of renal agenesis. The mean frequency of skeletal deformities is 7%. Duration of latency and severity of oligohydramnios independently increase the risk of skeletal abnormalities and act synergistically [107]. The GA at PPROM is not a significant determinant due to the progressive and continuous development of the axial skeleton.

Increased likelihood of skeletal deformities observed among infants diagnosed with PH suggests that the two disorders share common risk factors. Surgical correction is not generally required as they resolve with postnatal growth and physiotherapy.

Other morbidities. Other neonatal morbidities are similar to that in PTB and are related to GA at PPROM. These include RDS, bronchopulmonary dysplasia (BPD), IVH, NEC, sepsis, and retinopathy of prematurity. The incidence of neonatal IVH and cystic PVL increases in cases complicated by clinical chorioamnionitis [112]. Since midtrimester PPROM is associated with early delivery and infections, it constitutes a potential risk factor for long-term neurological morbidities. There may be a higher risk of PVL in infants born after prolonged PPROM compared with other types of prematurity [113]. Prolonged latency in midtrimester PROM patients is not associated with an increasing frequency of abnormal neonatal cranial ultrasound examination [99].

Long-term morbidities. About 63%–84% of survivors after midtrimester PPROM will be neurologically intact [104]. Cerebral palsy rate in a group of 275 pregnancies complicated by early PPROM was 9.8% [94].

Maternal

Cesarean delivery. The CD rate increases secondary to more common fetal heart rate abnormalities (related to oligohydramnios and chorioamnionitis), malpresentation, and abruptio. A classical uterine incision may be required to reduce fetal trauma at early GA (oligohydramnios, fetal malpresentation, and lower uterine segment characteristics constitute risk factors).

Retained placenta. The risk of undergoing either uterine exploration or curettage is 9%–18%, and more likely if rupture occurs prior to 20 weeks of gestation.

Postpartum endometritis. This condition occurs in about 13% of cases. Postpartum maternal sepsis (about 0.8%) and death (about 1/1000) are uncommon. The risk of maternal infection is inversely proportional to the latency period.

Management

Counseling

Parents should be counseled regarding the prognosis, complications, and management of PPROM <23 weeks. The options are expectant management or delivery. The impact of immediate delivery on neonatal outcome, and potential benefits and risks of conservative management, should be reviewed. PPROM related to genetic amniocentesis is associated with favorable outcomes even with expectant management (91%–99% perinatal survival), which is very different than the prognosis with spontaneous PPROM <23 weeks.

Incidences of most complications are inversely proportional to GA at PPROM, latency, residual amniotic fluid volume, and GA at delivery.

The latency period between membrane rupture and delivery is critical in determining perinatal outcome.

Latency is indirectly correlated with GA at PPROM (Table 19.3) [14]. While the mean latency is about 7 days, the median latency may be up to about 10–21 days because of the few pregnancies that gain a lot more than 14 days. Up to 14% of women with midtrimester PPROM stop amniotic fluid loss, presumably due to resealing of membranes [98,114]. This subgroup of cases has outcomes similar to pregnancies uncomplicated by membrane rupture. In 10%–20% of patients, amniotic fluid loss continues, but a partial reaccumulation during expectant management is observed [17,105].

Oligohydramnios (AFI <5 or MVP <2 cm) on admission, during latency period, or at the last ultrasonographic examination is associated with shorter latency, and higher occurrences of PH, chorioamnionitis, and perinatal mortality [108,115,116]. Conversely, adequate residual amniotic fluid volume identifies cases with elevated odds of perinatal survival (85%–93%) and better long-term neurological outcomes [102,115].

In a retrospective cohort of 92 cases of PPROM under 24 weeks, the overall neonatal survival rate at discharge was 85%; the survival rate was lower and the developmental delay more frequent in patients with persistent oligohydramnios compared with those with normal amniotic fluid volume [117].

Workup

Similar to PROM >23 weeks.
**Therapy**

See Figure 19.1.

*Delivery* in previable PPROM is indicated in the presence of:

- Intrauterine death
  - Spontaneous onset of labor
  - Evidence of maternal and/or fetal infections
  - Any other obstetric medical condition necessitating delivery (e.g., abruptio placentae and preeclampsia)
  - Pregnancy termination request

Delivery (termination) can be carried out usually by induction or by dilatation and extraction (D&X) by experienced operators, with no trials comparing these two modalities.

For management of women who elect expectant management, unfortunately there are no trials on any interventions (e.g., steroids, antibiotics, and magnesium sulfate) for previable PPROM. Data for any benefit of most interventions are available only for PPROM diagnosed at ≥23 weeks, so these interventions should be used with caution before this GA, with the patients understanding these limitations in medical knowledge. If pregnancy continues, patients need to be counseled on clinical variables and complications that may affect outcome and treatment serially at diagnosis, during the latency period, and at delivery. Given the high incidence of infection, cerclage should be removed in women with cerclage in place at the time of PPROM.

**Transabdominal Amnioinfusion**

Serial transabdominal amnioinfusions in the second trimester aim to enhance amniotic fluid volume, reduce lung hypoplasia, prolong latency and improve pregnancy outcomes. Data come mainly from nonrandomized prospective or retrospective trials. A meta-analysis on this topic, not including PPROM <23 weeks, reports a perinatal mortality reduction and a longer latency period after amnioinfusion [118], while the Cochrane review doesn’t identify any randomized study about transabdominal amnioinfusion before 26 weeks [119]. A recent randomized study on 56 women (serial weekly transabdominal or expectant management) shows no major differences in maternal, perinatal or pregnancy outcomes. It is considered a pilot study, but it demonstrates that such a study is feasible [120]. Because amnioinfusion in previable PPROM has not been validated in large randomized studies, it can be proposed only in research programs after informed consent of the patient.

**Resealing Techniques**

Resealing techniques as amniopatch (see section PPROM Following Invasive Procedures (Iatrogenic PPROM) and Strategies to Repair) has been proposed, but their efficacy in spontaneous PROM is low.

**Proposed Management**

In women desiring expectant management with PPROM <23 weeks, there is no data to support benefit of hospitalization, optimal GA for corticosteroids, antibiotics, or other interventions (Figure 19.1). Some consider hospital bed rest during the first few days, but this is not supported by level 1 or other data. Some administer an initial course of broad-spectrum antibiotic prophylaxis (7 days or until culture results), but again this is not supported by any RCT or other data. Avoid vaginal examination until labor or delivery. During expectant management, monitor for the onset of infectious complications, observing temperature, maternal or fetal tachycardia, uterine contractions or tenderness, or purulent vaginal discharge. As outpatient, instruct to avoid intercourse, check temperature, and refer to hospital for vaginal bleeding and contractions. At 23/0/7 weeks, suggest (readmission to the hospital and active expectant management, including administration of at least one course of steroids and antibiotics, assuming not already given.

**PPROM FOLLOWING INVASIVE PROCEDURES (IATROGENIC PPROM) AND STRATEGIES TO REPAIR**

Iatrogenic PPROM (iPPROM) and spontaneous PPROM have different pathophysiologicals. In iPPROM the medical instrument creates the hole in the membranes, which subsequently fails to close. In studies of women undergoing second-trimester amniocentesis for prenatal diagnosis, the risk of PROM is approximately 1% [88]. When leakage of amniotic fluid occurs after amniocentesis, the outcome is better than after spontaneous preterm PROM and reaccumulation of normal amniotic fluid volume is more probable [110,121]. In one series of 11 patients with PROM after genetic amniocentesis, there was one previable pregnancy loss, reaccumulation of normal amniotic fluid occurred within 1 month in 72% of patients, and the perinatal survival rate was 91% [88]. After appropriate counseling, patients with PROM after genetic amniocentesis typically are managed expectantly as outpatients.

PPROM may also occur following fetoscopy for fetal surgical procedures (e.g., laser therapy for twin–twin transfusion syndrome [TTTS] or for twin-reversed arterial perfusion [TRAP sequence], or tracheal occlusion for congenital diaphragmatic hernia) or other minimally invasive fetal surgery (e.g., intrauterine blood sampling and transfusion or fetal shunting for lower urinary tract obstructions and pleural effusions). The incidence of iPPROM after these procedures varies significantly in the literature. iPPROM occurred in about 30% of cases treated by minimally invasive fetal surgery in one report [122]. Parameters that have been suggested to impact iPPROM rates are the diameter of the surgical instrument and the number of entries to the uterine cavity [123]. Other possible risk factors are duration and difficulty of the procedure, operator experience, membrane friction due to instrument manipulation, type of anesthesia, GA at intervention, number of interventions and placental location. In one review the maximum diameter of the instrument predicted iPPROM rate, GA at birth and fetal survival [122].

Several techniques have been developed in an attempt to artificially reseal the fetal membranes and prevent leakage of amniotic fluid. These include intra-amniotic injection of platelets and cryoprecipitate (amniopatch), sealing the cervical canal, and fetoscopic laser coagulation [89,124–126]. Compared with spontaneous PPROM, iPPROM is not only a different entity etiologically, but also in its better response to therapeutic measures. One report demonstrated successful membrane sealing for persistent oligohydramnios after amniocentesis and fetoscopy in seven women, using intraamniotic injection
of platelets and cryoprecipitate through a 22-gauge needle [89]. Other case reports include successful membrane sealing with intrauterine injection of gelatin sponge, fibrinogen and thrombin combinations, and with fibrin glue with powdered collagen [126–129]. Another study reported cessation of leakage and restoration of normal amniotic fluid volume in twelve women with intracervical instillation of fibrin sealants [124].

According to a review of cases of iPPROM, amniopatch effectively seals the fetal membranes in over two-thirds of cases [130]. A retrospective analysis of 24 amniopatch procedures performed for PPROM after a needle-based procedure or after fetoscopic intervention reported a success rate of 58% [131]. Unfortunately, the safety of these approaches has not been evaluated in large studies, and there are several anecdotal reports of maternal and perinatal morbidity and mortality. Even if such practices have not yet been incorporated into clinical practice, current experience suggests that, in cases of fluid leakage following an invasive fetal procedure, the use of sealing techniques is a therapeutic option that should be researched further.

PPROM IN TWIN GESTATIONS

Overall, PPROM complicates 7%–8% of twin pregnancies at a mean GA of 30–32 weeks (see also Chapter 44, in Maternal-Fetal Evidence Based Guidelines). PPROM occurs at an earlier GA among multiple gestations with 36% of twin PPROM cases occurring at less than 28 weeks. PPROM in singleton pregnancies is related mainly to infection/inflammation, while PPROM in twin gestation may probably be more related to uterine overdistension. The latency period seems to be shorter in twins compared with singleton pregnancies. Most studies report a median latency of less than 24 hours with only 16%–50% of twin pregnancies remaining undelivered at 48 hours, decreasing to 7%–22% at 7 days. Latency tends to be longer when PPROM occurs before 30 weeks of gestation [132]. The membrane rupture usually occurs in the lower sac (90% of cases) [133].

Studies comparing obstetric outcome between singleton and twin pregnancies with PPROM reported no difference in neonatal outcome [134,135]. A retrospective cohort of 23 multifetal pregnancies complicated by PPROM before 26 weeks showed a median latency of 11 days with expectant management. Of the 46 newborns, 20 (43%) survived to hospital discharge. Of those, 12 (60%) experienced severe neonatal morbidity. The multiple with ruptured membranes was more likely to experience intrauterine fetal demise, but all other outcomes did not differ by membrane status [136]. In 48 multiple pregnancies complicated by PPROM and delivering at a median GA of 31 weeks, neonatal morbidity and mortality were not different between the presenting and nonpresenting twin and no difference was found between fetuses with or without ruptured sac. The outcomes were not affected by duration of the latency period [137]. These data suggest that rupture of membranes per se probably do not worsen the outcomes, and support a conservative management of twin pregnancies with PPROM. The incidence of both clinical and subclinical chorioamnionitis has been reported as significantly higher in singleton pregnancies compared with twin pregnancies complicated by PPROM [133,138]. A study evaluating neonatal and infant outcomes in 151 twin gestations with PPROM at 24–31 weeks reported a neonatal mortality rate of 9.0% and an overall cerebral palsy rate of 7.3% [139].

The management of PPROM in twin pregnancies does not differ significantly from that of singletons. Antenatal steroids, antibiotics to prolong latency, magnesium sulfate for neuroprotection with delivery no later than 34 weeks of gestation are all recommended in twins with PPROM.

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Naef RW III, Albright JR, Ross EL, et al. Premature rupture of membranes at 34 to 37 weeks’ gestation: Aggressive versus conservative management. *Am J Obstet Gynecol*. 1998;178:126. [RCT, \( n = 120 \) women with PPROM at 34–36 weeks of gestation: chorioamnionitis 2% in induction vs. 16% in expectantly managed group. Tocolytics, steroids were not used; penicillin to all women for GBS prophylaxis]


Premature rupture of membranes at or near term

Kimberly Ma and Sally Segel

KEY POINTS

• The diagnosis of premature rupture of membranes (PROMs) at term is based on pooling, ferning, and nitrazine tests on speculum examination. Rapid bedside immunoassays such as placental alpha-microglobulin-1 or others may be helpful in cases in which the diagnosis of rupture of membrane (ROM) is suspected, but cannot be confirmed with pooling, ferning, and/or nitrazine tests. Initial digital examinations should be avoided, as it is associated with a higher risk of infection.

• The main complication is intrauterine infection; this incidence increases with duration of PROM, and, with longer latency, the risk of neonatal infection also increases.

• Women with PROM at term should be hospitalized and induced with oxytocin within 6 to 12 hours of PROM, or earlier as feasible. Most women with PROM at term, if given a choice, prefer induction. Oxytocin induction is safe, effective, and cost-effective. Misoprostol induction is an alternative that is just as effective when 25 mg every 4 hour dosing is used, but there is limited data regarding its safety. Foley catheter ripening is another safe and efficacious alternative, especially in cases of unfavorable cervical examination.

• Antibiotics are recommended if the patient is group B streptococcus (GBS) positive, or complications (choioamnionitis, PROM >18 hours) develop during labor. In women expected to have a latency from PROM to delivery of over 12 hours, antibiotics are associated with significantly decreased chorioamnionitis and endometritis.

DEFINITION

Premature rupture of membranes (PROM) is rupture of fetal membranes prior to the onset of labor [1]. Preterm premature rupture of membranes (PPROM) is premature rupture of fetal membranes ≤37 weeks [1]. This chapter includes guidance for near-term (34–36 6/7 weeks) as well as term (≥37 weeks) PROM. For PPROM <34 weeks, see Chapter 19.

DIAGNOSIS

The diagnosis of PROM and PPROM is based on history and physical examination findings (see also Chapter 19). A maternal history suggestive of PROM includes a gush of fluid soaking clothes or a constant trickle of fluid that does not stop. Physical examination for PROM starts with a sterile speculum examination [1–3]. Visualization of amniotic fluid passing from the cervical canal is diagnostic of this condition [1–3].

Another helpful test is the nitrazine test. The pH of the vagina is usually 4.5 to 6.0, whereas amniotic fluid has a pH 7.0 to 7.7; nitrazine paper turns blue with pH >6.5 [1,2]. A false positive nitrazine test can be caused from blood, semen, alkaline antiseptic, or bacterial vaginosis. A false-negative nitrazine test can be caused by prolonged leaking of amniotic fluid or minimal residual fluid. The sensitivity of nitrazine is 90.2% (81.3%–100%) and the specificity is 79.3% (16%–100%) [2].

A third helpful test is called “ferning.” A swab of vaginal fluid from the posterior fornix is placed on a slide and allowed to air dry. Arborization (ferning) under microscopic visualization suggests rupture of membranes. Cervical mucus can cause a false positive result. Ferning has a sensitivity of 90.8% (62.0%–98.5%) and a specificity of 95.3% (88.2%–100%) [2].

The combination of vaginal pool of fluid, nitrazine, and ferning has a sensitivity of 90.8% and a specificity of 95.6% [2]. These tests are more accurate when patients present in labor and less accurate when nonlaboring [3]. If these tests are equivocal, an ultrasound can be performed to evaluate for amniotic fluid, but oligohydramnios is not diagnostic of PROM, since it can be associated with other etiologies, such as placental insufficiency.

Placental alpha-microglobulin-1 is a 34-kDa glycoprotein abundant in amniotic fluid (2,000–25,000 ng/mL); there is a negligible amount of this glycoprotein in vaginal fluid with intact fetal membranes (0.05–2.0 ng/mL) [2,4,5]. AmniSure has a bedside immunoassay that uses the delta in concentration of placental alpha-microglobulin-1 for diagnostic accuracy. AmniSure has a sensitivity of 98.7% to 98.9% and a specificity of 87.5% to 100% [4,5].

Insulin-like growth factor–binding protein 1 (IGFBP-1) has a high concentration in the amniotic fluid compared with other body fluids such as vaginal secretions, urine, or semen [6]. An immunochromatography dipstick method (Actim PROM) is available to diagnose rupture of membranes. The test is more popular in Europe compared with the United States and is most accurate when the timing of the test is close to the timing of rupture of membranes. The sensitivity of the test is 95%–100% with a specificity of 93%–98%. [7–10].

A negative fetal fibronectin can be helpful in ruling out rupture of membranes, however a positive test cannot distinguish between intact and rupture of membranes [11]. Alpha-fetoprotein levels are higher in amniotic fluid compared with urine, semen, or normal vaginal discharge. A recent study showed a 96.2% sensitivity and 100% specificity, however may be falsely positive if a vaginal infection is present and requires further study in a larger cohort [12]. Further studies are needed to assess reliability with special attention paid to time of presumed membrane rupture [2,3]. Placental alpha-microglobulin-1, IGFBP-1, and these other tests have limited clinical applicability currently given the high accuracy of pooling, ferning, and nitrazine tests. Nonetheless, they can be helpful in cases in which the diagnosis of ROM is suspected, but cannot be confirmed with pooling, ferning, and/or nitrazine tests.

INCIDENCE

PROM occurs in approximately 8% of all term deliveries [1].
ETIOLOGY
The etiology of PROM without signs of infection or bleeding is often unknown, and this should be considered a physiologic, not pathologic, event.

RISK FACTORS
The risk factors associated with rupture of fetal membranes include low socioeconomic status, low body mass index (BMI) <19.8 kg/m², nutritional deficiencies of ascorbic acid and copper, connective tissue disorders, smoking, cold knife cone biopsy, cervical cerclage, pulmonary disease, uterine overdistension, and amniocentesis [1]. In addition, pregnant women with a previous preterm birth, midtrimester short cervix, and preterm labor are at increased risk for PROM; however, the majority of cases have no identifiable cause [1].

COMPLICATIONS
The most common complications of PROM are chorioamnionitis, endomyometritis, and postpartum hemorrhage. With increasing time interval from rupture of membranes, there is a significant increase in these complications: about 12 hours for chorioamnionitis, 16 hours for endomyometritis, and 8 hours for postpartum hemorrhage [13]. As the latency from ROM to delivery increases, so does the risk for neonatal infection. Maternal GBS colonization also increases the risk for neonatal infection. Incidence of cesarean section is not affected by management with either induction or expectant management, but depends on other risk factors (e.g., nulliparity) [14].

MANAGEMENT
Initial Evaluation
Initial evaluation of PROM involves confirmation of the diagnosis. Digital examination can increase the risk of infection and as a result should be avoided [14]. The cervix can be visually assessed for dilation and effacement [15]. Fetal presentation should be confirmed by ultrasound. Fetal well-being and the presence of uterine contractions should be assessed by external monitoring.

Counseling
Risk factors, complications, and management of PROM should be reviewed with the patient. About 50% of women with PROM at term deliver within 6–12 hours and 95% of women will deliver within 28 hours of membrane rupture [1,14]. Principles of management can be reviewed (Table 20.1).

Hospitalization
Compared with management in the hospital, expectant management of PROM at term at home is associated with a 52% increase in need for maternal antibiotics for nulliparas and 97% more neonatal infections [16]. PROM at term should be managed in hospital.

Table 20.1 Management of PPROM at ≥34 Weeks

<table>
<thead>
<tr>
<th>Management of PPROM at ≥34 Weeks</th>
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<tbody>
<tr>
<td>• Deliver</td>
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<tr>
<td>• Manage in hospital</td>
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<tr>
<td>• There is insufficient evidence to recommend one induction agent over another, but oxytocin has been the best studied for safety and effectiveness. The balloon is also safe and effective, and can be considered especially when the cervix is &lt;3 cm dilated</td>
</tr>
<tr>
<td>• Antibiotics are recommended if the patient is GBS positive, or complications (chorioamnionitis, PROM &gt; 18 hours) develop during labor. In women expected to have a latency from PROM to delivery of over 12 hours, antibiotics are associated with significantly decreased chorioamnionitis and endometritis</td>
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Delivery versus Expectant Management
Women with PPROM ≥34 weeks should be delivered (see also Chapter 19). Induction of labor with oxytocin has been shown to decrease the rate of maternal chorioamnionitis and postpartum fevers without significantly affecting the rate of cesarean delivery, compared with expectant management [14]. In most randomized controlled trials (RCTs), induction was with oxytocin or prostaglandin, with one trial using homeopathic caulo-phylum. Overall, no differences are detected for mode of birth between induced and expectant groups: cesarean (relative risk [RR] 0.94, 95% confidence interval [CI] 0.82–1.08); operative vaginal birth (RR 0.98, 95% CI 0.84–1.16). Significantly fewer women in the induced compared with expectant management groups have chorioamnionitis (RR 0.74, 95% CI 0.56–0.97) or endometritis (RR 0.30, 95% CI 0.12–0.74). No difference is seen for neonatal infection (RR 0.83, 95% CI 0.61–1.12). However, fewer infants under induced management went to neonatal intensive or special care compared with expectant management (RR 0.72, 95% CI 0.57–0.92, number needed to treat is 20) [17].

Compared with expectant management, induction of labor by oxytocin is associated with a 37% decrease in risk of maternal infection, 28% in endometritis, and 36% in neonatal infection [17]. Based on one trial, women were more likely to view their care positively if labor was induced with oxytocin [14]. Cesarean delivery rates are similar between groups. Oxytocin is associated with more frequent use of pain relief and internal fetal heart rate monitoring. Perinatal mortality rates are low and not significantly different between groups, although the trend is toward fewer perinatal deaths with induction of labor by oxytocin [17].

Compared with placebo or no treatment, both vaginal prostaglandin E2 (PGE2) and E2a (PGF2a) reduce the likelihood of vaginal delivery not being achieved within 24 hours, with no evidence of a difference in cesarean delivery [17]. Induction of labor by prostaglandins is associated with a decrease by 23% in risk of chorioamnionitis and by 21% in admission to neonatal intensive care. Induction by prostaglandins is associated with a more frequent maternal diarrhea and use of anesthesia and/or analgesia [17]. There is insufficient data to make meaningful conclusions for the comparison of vaginal PGE2 and PGF2a. PGE2 tablet, gel, and pessary appear to be as efficacious as each other. Lower-dose regimens appear as efficacious as higher dose regimens [17].

Medications for Induction
Immediate induction of labor with intravenous oxytocin has been shown to decrease the rate of maternal infection without increasing the rate of cesarean section [14]. Please see above data on individual agents for induction compared with placebo or no treatment.

Currently, there have been no studies that demonstrate misoprostol or PGE2 are superior to intravenous oxytocin even in women with an unfavorable cervix [18]. However, when misoprostol is compared with PGE2, length of labor is shorter but there is increased uterine tachysystole [18].

Abbreviations: GBS, group B streptococcus; PPROM, preterm premature rupture of membranes; PROM, premature rupture of membranes.
Compared with prostaglandins, induction with oxytocin is associated with decrease in maternal nausea and/or vomiting, numerous vaginal examinations, chorioamnionitis and neonatal infections, neonatal antibiotic therapy, and admission to neonatal intensive care unit (NICU), but increase in epidural analgesia and internal fetal heart rate monitoring, Cesarean delivery, endometritis, and perinatal mortality are not significantly different between the groups [17]. Cost is less with oxytocin induction.

Vaginal misoprostol in doses above 25 µg every 4 hours is more effective than PGE2, intracervical PGE2, and oxytocin (or obviously expectant management) in achieving vaginal delivery [17,19]. Compared with oxytocin, misoprostol was associated with a decrease in cesarean section from 51% to 20% in a small trial in women with an unfavorable cervix and from 14% to 11% in other women [17,19]. If a dose of 50 µg of vaginal misoprostol is used, in 85% of cases only one dose is needed for induction with term PROM, but this is associated with a higher rate of uterine tachysystole compared with oxytocin, with similar maternal and neonatal outcomes [20]. However, the studies reviewed were not large enough to exclude the possibility of serious adverse events with misoprostol including neonatal complications from uterine hyperstimulation. Lower doses of vaginal misoprostol were similar to other methods in effectiveness and risk. Oral misoprostol appears to be an effective means of labor induction comparable to oxytocin, but again the studies reviewed were not large enough to exclude the possibility of serious adverse events with misoprostol [21–23].

There is insufficient evidence to assess if the combination of dinoprostone (PGE2) together with oxytocin is superior to oxytocin alone [24].

The safety and efficacy of Foley bulbs or other mechanical means of induction in women with PROM have been evaluated, and Foley appears to be both safe and efficacious for cervical ripening of women with PROM and unfavorable cervical examination [25–27] (see also Chapter 21).

In summary, in patients with PROM at term, once the diagnosis has been confirmed, induction of labor is recommended. Induction should probably occur at least within 6–12 hours of PPROM, or earlier if feasible. If expectant management is performed, there is a significant increase in chorioamnionitis and neonatal infection and time duration increases [28–30]. Currently, oxytocin is the recommended induction medication, as it is safe, effective, and cost-effective. Foley bulb (or other balloon) is also a safe and effective intervention, especially if unfavorable cervical examination (e.g., cervix <3 cm dilated) [25–27]. Vaginal and oral misoprostol are also safe and effective but have not been proven to be superior to oxytocin when appropriate doses (e.g., 25 µg for vaginal misoprostol) are used [18]. With increasing latency from PROM to delivery, there is an increased risk for chorioamnionitis, endomyometritis, and postpartum hemorrhage [13]. Women should be counseled to initiate induction of labor in order to decrease the risk of these complications.

**Antibiotics**

**GBS positive**: In women with a positive GBS culture, antibiotics should be administered according to the CDC guideline [31] (see Chapter 37 in Maternal-Fetal Evidence Based Guidelines).

**Women who are GBS positive with PROM at ≥34 weeks are treated with penicillin in labor.** Ampicillin is a reasonable alternative. If the patient is penicillin-allergic but not at high risk for anaphylaxis, cefazolin is the agent of choice. For women at high risk for anaphylaxis, GBS isolate should be tested for susceptibility to clindamycin and erythromycin. Vancomycin is recommended if isolate is resistant to either clindamycin or erythromycin. Intrapartum treatment for chorioamnionitis is recommended regardless of GBS maternal status.

In these revised recommendations, 4 hours of intravenous antibiotics are considered adequate treatment for a neonate; however, no medically necessary obstetric procedure should be deferred to achieve 4 hours of intravenous antibiotic therapy [31]. If a patient presents with PROM at term, antibiotics can be administered and induction of labor can begin concurrently. In addition, if a woman is scheduled for a repeat cesarean section and presents with PROM, GBS antibiotic prophylaxis should be administered while arranging for the repeat cesarean section [31].

**GBS negative**: There is some evidence to recommend for antibiotics in GBS-negative women with PPROM at ≥34 weeks. A recent meta-analysis evaluated the efficacy of antibiotic prophylaxis in women with term or near-term premature rupture of membranes and concluded antibiotic prophylaxis was not associated with maternal or neonatal benefits. However in women with latency greater than 12 hours, prophylactic antibiotics were associated with significantly lower rates of chorioamnionitis (2.9% vs. 6.1%, RR 0.49, 95% CI 0.27–0.91) and endometritis (0% vs. 2.2%, RR 0.12, 95% CI 0.02–0.62) [32].

**GBS status unknown**: If the patient has no risk factors (previous infant with GBS sepsis, <37 weeks, rupture of membranes >18 hours, chorioamnionitis), antibiotics can still be considered, especially in the woman expected to have a latency to delivery interval >12 hours [20]. Alternatively, in PPROM 34 to 36 6/7 weeks, if the patient has no GBS culture from the previous 5 weeks, antibiotics can be administered per protocol or a rapid GBS test can be performed. If the rapid GBS result is negative, antibiotics can be withheld unless other risk factors develop (chorioamnionitis or rupture of fetal membranes greater than 18 hours) [31].

In summary, for PROM at ≥34 weeks, antibiotics should be given if the woman is GBS+ or infectious complications develop during labor [33], and considered for all women with expected latency from PPROM to delivery >12 hours [32].

**Special Considerations for PPROM at 34 to 36 Weeks’ Gestation**

Management of PPROM from 34/0 to 36 6/7 weeks is based on extrapolation from a few small studies [34–38], and therefore the scientific evidence is limited. These studies were performed before a standardized therapy for PPROM was developed, and corticosteroids and antibiotics were not routinely used. The current The American Congress of Obstetricians and Gynecologists (ACOG) Practice Bulletin states that “At 34 0/7 weeks or greater of gestation, delivery is recommended for all women with ruptured membranes. If expectant management is continued beyond 34 0/7 weeks of gestation, the balance between benefit and risk should be carefully considered and discussed with the patient, and expectant management should not extend beyond 37 0/7 weeks of gestation.” [1].

Recent studies have called attention to the neonatal complications of late PTB, which include respiratory complications, sepsis evaluations, hyperbilirubinemia requiring phototherapy, and death. This combination of potential complications has led to an increase in NICU admissions, increasing length of stay (LOS), urgent hospital visits, and hospital readmissions [39–46]. The effort to decrease late-preterm birth in the United...
States has been successful, with a decrease from 9.15% in 2006 to 7.99% in 2013 (a 13% decrease) [47].

The original research determining the best antenatal approach to PPROM including the late preterm period studied patients with documented fetal lung maturity [35,36]. Spinnato et al. randomized 47 nonlaboring PPROM patients to either delivery or expectant management with hospitalization and intensive antenatal surveillance. There was a statistically significant increase in chorioamnionitis or postpartum endometritis in expectant management group; however, neonatal morbidity and mortality were similar between groups [35]. Mercer et al. also randomized 93 PPROM patients from 32 to 36 weeks’ gestation with fetal lung maturity to immediate induction of labor or expectant management. Women randomized to expectant management had a significant increase in chorioamnionitis and fetal heart rate abnormalities in labor [36]. In addition, neonates from the expectant management group had a significant increase in the initiation and length of antibiotic treatment for presumed neonatal infection [36]. These two trials had significant similarities in that the expectant management group was hospitalized with intensive surveillance and antibiotics were only administered for chorioamnionitis. Overall their results demonstrated that expectant management in PPROM patients, including the late preterm period and with fetal lung maturity, had longer latencies and increased maternal infection without significant neonatal benefit. Naef et al. also performed a randomized trial of 120 PPROM patients in the late preterm period. Women in the expectant management group received GBS prophylaxis during their hospitalization [37]. Even with the addition of GBS prophylaxis, women randomized to the expectant management group had a significant increase in chorioamnionitis and their neonates had a significant increase in NICU admission with a trend toward an increase in culture-proven sepsis [37]. Overall, these studies found that expectant management produced longer latencies; however, there was a significant increase in maternal infection without demonstration of neonatal benefit [35–37]. Unfortunately the meta-analysis of the seven RCTs on this issue is not clinically helpful as studies have been underpowered for significant neonatal and maternal outcomes [48].

Two international RCTs compared delivery versus expectant management in women with PPROM diagnosed at 34–36 6/7 weeks gestation. In the combined analysis of 736 women who were randomized, induction of labor reduced the risk of chorioamnionitis (1.6% vs. 5.3%, RR 0.31, 95% CI 0.1–0.8). There was no statistical difference in neonatal sepsis with (2.7% with induction of labor vs. 4.1% with expectant management; RR 0.66, 95% CI 0.30–1.5). The overall rate of neonatal sepsis was lower than expected, and therefore the studies did not have adequate power to show a statistical difference in neonatal sepsis [49–50].

A recent multicenter randomized controlled trial was performed at 65 centers in 11 countries with 1839 women [51]. Women with ruptured membranes between 34 0/7 and 36 6/7 weeks gestation were randomized to immediate delivery or expectant management looking at the primary outcome of neonatal sepsis. There was no statistical difference in neonatal sepsis (2% in immediate delivery versus 3% in expectant management, RR 0.8, 95% CI 0.5–1.3). The immediate delivery group had increased rates of respiratory distress (RR 1.6, 95% CI 1.1–2.3), mechanical ventilation (RR 1.4, 95% CI 1.0–1.8), and longer intensive care unit days (4 days vs. 2 days, p < 0.01) compared with the expectant management group. However, women in the immediate delivery group had a lower rate of antepartum or intrapartum hemorrhage (RR 0.6, 95% CI 0.4–0.9), intrapartum fever (RR 0.4, 95% CI 0.2–0.9), postpartum antibiotic use (RR 0.8, 95% CI 0.7–1.0), and longer hospital stay (5 days vs. 6 days, p < 0.01). Expectant management performed at this gestational age should weigh the potential maternal benefits with the maternal risks with the consideration of the use of latency antibiotics.

With the additional information from these studies, ACOG continues to recommend delivery for all women with ruptured membranes after 34 0/7 weeks or greater. Although there is concern for maternal risks associated with late preterm birth, neonates born to women with ruptured membranes have a higher rate of adverse outcomes when matched with controls for gestational age and higher complications when chorioamnionitis is present [52,53]. Delivery of all women with ruptured membranes after 34 0/7 weeks or greater should continue to be the standard of care until there is additional data.

There is new evidence that steroids are associated with neonatal benefits also >34 weeks, as three RCTs have been done on steroids 34–36 6/7 weeks in women at risk of PTB, and two RCTs in women at >37 weeks. Infants of women who received antenatal corticosteroids >34 weeks had a significantly lower incidence of respiratory distress syndrome (RDS) (RR 0.76, 95% CI 0.62–0.93), mild RDS (RR 0.40, 95% CI 0.23–0.69), moderate RDS (RR 0.39, 95% CI 0.18–0.89), transient tachypnea of the newborn (RR 0.62, 95% CI 0.50–0.77), severe RDS (RR 0.66, 95% CI 0.53–0.82), use of surfactant (RR 0.61, 95% CI 0.38–0.99), mechanical ventilation (RR 0.62, 95% CI 0.41 to 0.94), significantly lower time on oxygen (mean difference [MD] −2.06 hours, 95% CI −2.17 to −1.95), lower maximum inspired oxygen concentration (MD −0.66%, 95% CI −0.69 to −0.63), lower LOS in NICU (MD −7.64 days, 95% CI −7.65 to −7.64), higher Apgar score at 1 and at 5 minutes (MD 0.06, 95% CI 0.05–0.07) compared with those who did not. Steroids should be considered for decreasing respiratory morbidity and other neonatal outcomes in women ≥34 weeks [54,55]. See Chapter 19 for other details on management for women with PPROM <34 weeks [48].

REFERENCES


Induction of labor

Corina N. Schoen and Anthony C. Sciscione

KEY POINTS

• Complications of induction of labor at term include prolonged first stage of labor, operative delivery and their risks.
• An early (e.g., <20 weeks) ultrasound examination helps determine accurate gestational age and is associated with a reduction in the rates of induction of labor for post-term pregnancy.
• Gestational age should be documented accurately before considering induction.
• Possible indications for induction of labor, and suggested best gestational age for induction, are listed in Table 21.1.
• Without indications, induction before 39 weeks should be avoided. An induction based solely on maternal request should be designated as such (induction for maternal request).
• Contraindications to induction of labor include transverse or oblique fetal lie, umbilical cord prolapse, previous classical uterine incision or transfundal uterine surgery (e.g., from myomectomy), placenta or vasa previa, active genital herpes infection, and any contraindications to vaginal delivery, or indication for cesarean delivery (CD) (Table 21.2).
• Misoprostol should not be used for cervical ripening or labor induction in women with prior uterine incisions, given the >5% risk of uterine rupture. In women with prior uterine incisions, prostaglandin E2 (PGE2) for cervical ripening is associated with approximately 1.4%–2.5% risk of rupture; oxytocin has a 1.1% risk. Cervical ripening with the Foley catheter does not appear to be associated with any additional risk of uterine rupture in patients undergoing a trial of labor after cesarean section.
• A Bishop score of ≥9 is usually associated with the probability of vaginal delivery after labor induction similar to that after spontaneous labor.
• In women with an unfavorable (Bishop score <5, or even <9) cervical examination:
  • Sweeping of membranes at term doubles the rate of onset of labor (to approximately 36%) in the next 48 hours, without major complications.
  • The Foley balloon is associated with less hyperstimulation accompanied by fetal heart rate (FHR) changes, and increased use of oxytocin, but results in the same number of vaginal deliveries and the same number of women delivered within 24 hours as all of the locally applied prostaglandins ([LAPG]; prostaglandin E1 [PGE1] and PGE2). It is therefore one of the preferred methods for labor induction when the cervix is <3 cm dilated.
  • Vaginal misoprostol 25 mg every 3–6 hours is the preferred safe dosage, as effective as PGE2 or any other method; this is also a preferred method of labor induction.
• PGE2 tablet, gel, and insert appear to be as safe and efficacious as each other in terms of tachysystole, CD rates, and neonatal outcomes.
• Other cervical ripening or induction agents are either not sufficiently studied, unsafe, or not as effective as the agents already mentioned.
• In women with favorable cervical examination:
  • Oxytocin is safe and effective for induction of labor in these women.
  • There are insufficient safety data for outpatient use of pharmacologic cervical ripening or induction agents. However, there is emerging evidence that the Foley catheter may be an effective and safe method for outpatient cervical ripening.
• Contraction pressures of ≥200 Montevideo units should be targeted in induction or augmentation of laboring patients to achieve adequate labor. While the definition of a failed induction of labor is elusive, a failed induction should not be diagnosed until after 24 hours of oxytocin after membrane rupture in the active phase (usually ≥6 cm in nulliparous women), assuming reassuring fetal heart pattern.
• Induced labor should be managed, in general, as spontaneous labor.

DEFINITIONS

Induction of labor is the stimulation of uterine contractions prior to spontaneous labor in order to achieve childbirth. Cervical ripening is a process that occurs prior to labor in which the cervix is softened, thinned, and dilated.

INCIDENCE/EPIDEMIOLOGY

In 2013, there were 3,932,181 total births in the United States, of which 903,638 were induced (23%) [1]. From 1990 to 2010, the induction rate has increased for all gestational ages [2]. However, the incidence of preterm inductions has decreased since 2005, and accounted for just 13.9% of all inductions in 2013 [2]. Although previous trends showed an increased rate of inductions without medical indication [3], there is now evidence for a decreasing rate of indicated preterm births (17% decline from 2005 through 2012) [4]. However, it has been shown that using birth certificate data overestimates the number of non-medically indicated inductions by 11-fold [5].

BASIC PATHOPHYSIOLOGY

The cervix functions as the gatekeeper to parturition maintaining a fine balance between the integrity of the pregnancy and delivery. Histologically, the cervix is composed of mostly collagen, with some smooth muscle; the stability of these components and the ability to become dynamic when stressed on
Table 21.1 Common Indications for Possible Induction of Labor or Cesarean Delivery (Suggested Timing of Delivery for Selected Conditions)

<table>
<thead>
<tr>
<th>Placental and Uterine Issues</th>
<th>Gestational Age&lt;sup&gt;a&lt;/sup&gt; for Delivery</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Placenta previa&lt;sup&gt;b&lt;/sup&gt;</td>
<td>36–37 weeks</td>
</tr>
<tr>
<td>• Placenta accreta/increta/percreta with placenta previa</td>
<td>34–35 weeks</td>
</tr>
<tr>
<td>• Prior classical cesarean (upper segment uterine incision)</td>
<td>36–37 weeks</td>
</tr>
<tr>
<td>• Prior myomectomy requiring cesarean delivery</td>
<td>37–38 weeks (Situations with more extensive or complicated procedures may require earlier delivery, similar to prior classical cesarean)</td>
</tr>
<tr>
<td>• Prior uterine rupture</td>
<td>36–37 weeks</td>
</tr>
<tr>
<td>• Abruptio placenta</td>
<td>At detection (see Chapter 26)</td>
</tr>
<tr>
<td>• Intrauterine infection (e.g., chorioamnionitis)</td>
<td>At detection</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Fetal Issues</th>
<th>Gestational Age&lt;sup&gt;a&lt;/sup&gt; for Delivery</th>
</tr>
</thead>
<tbody>
<tr>
<td>• FGR—singleton</td>
<td>38–39 weeks</td>
</tr>
<tr>
<td>• Otherwise uncomplicated, no concurrent findings</td>
<td>34–37 weeks</td>
</tr>
<tr>
<td>• Concurrent conditions (e.g., abnormal umbilical artery Dopplers)</td>
<td>Expedient delivery regardless of gestational age</td>
</tr>
<tr>
<td>• Persistent abnormal fetal surveillance suggesting imminent fetal jeopardy</td>
<td></td>
</tr>
<tr>
<td>• FGR—twin gestation</td>
<td>36–37 weeks</td>
</tr>
<tr>
<td>• Dichorionic-diamniotic twins with isolated FGR</td>
<td>32–34 weeks</td>
</tr>
<tr>
<td>• Monochorionic-diamniotic twins with isolated fetal growth restriction</td>
<td></td>
</tr>
<tr>
<td>• Concurrent conditions (e.g., abnormal umbilical artery Dopplers)</td>
<td>Expedient delivery regardless of gestational age</td>
</tr>
<tr>
<td>• Persistent abnormal fetal surveillance suggesting imminent fetal jeopardy</td>
<td></td>
</tr>
<tr>
<td>• Fetal congenital malformations</td>
<td>34–39 weeks</td>
</tr>
<tr>
<td>• Suspected worsening of fetal organ damage</td>
<td></td>
</tr>
<tr>
<td>• Potential for fetal intracranial hemorrhage (e.g., Vein of Galen aneurysm and NAIT)</td>
<td></td>
</tr>
<tr>
<td>• When delivery prior to labor is preferred (e.g., EXIT procedure)</td>
<td></td>
</tr>
<tr>
<td>• Previous fetal intervention</td>
<td></td>
</tr>
<tr>
<td>• Concurrent maternal disease (e.g., preeclampsia and chronic hypertension)</td>
<td></td>
</tr>
<tr>
<td>• Potential for adverse maternal effect from fetal condition</td>
<td></td>
</tr>
<tr>
<td>• Expedient delivery regardless of gestational age</td>
<td></td>
</tr>
<tr>
<td>• When intervention is expected to be beneficial</td>
<td></td>
</tr>
<tr>
<td>• Fetal complications develop (abnormal fetal surveillance, new onset hydrops fetalis, and progressive/new-onset organ injury)</td>
<td></td>
</tr>
<tr>
<td>• Maternal complications develop (mirror syndrome)</td>
<td></td>
</tr>
<tr>
<td>• Multiple gestations: dichorionic/diamniotic&lt;sup&gt;b&lt;/sup&gt;</td>
<td>38 weeks</td>
</tr>
<tr>
<td>• Multiple gestations: monochorionic/diamniotic&lt;sup&gt;b&lt;/sup&gt;</td>
<td>34–37 weeks</td>
</tr>
<tr>
<td>• Multiple gestations: di/di or mono/di with single fetal death&lt;sup&gt;b&lt;/sup&gt;</td>
<td>If occurs at or after 34 weeks, consider delivery</td>
</tr>
<tr>
<td>• Multiple gestations: monochorionic/monoamniotic&lt;sup&gt;b&lt;/sup&gt;</td>
<td>32–34 weeks</td>
</tr>
<tr>
<td>• Multiple gestations: monochorionic/monoamniotic with single fetal death&lt;sup&gt;b&lt;/sup&gt;</td>
<td>Consider delivery; individualized according to gestational age and concurrent complications</td>
</tr>
<tr>
<td>• Oligohydramnios (MVP &lt;2 cm)—isolated but persistent&lt;sup&gt;b&lt;/sup&gt;</td>
<td>37 weeks</td>
</tr>
<tr>
<td>• Nonreassuring fetal heart testing (e.g., category III FHR tracing)&lt;sup&gt;c&lt;/sup&gt;</td>
<td>At detection (see Chapter 10)</td>
</tr>
<tr>
<td>• Fetal death</td>
<td>At detection (see Chapter 10)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Maternal Issues</th>
<th>Gestational Age&lt;sup&gt;a&lt;/sup&gt; for Delivery</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Chronic hypertension—no medications&lt;sup&gt;b&lt;/sup&gt;</td>
<td>38–39 weeks</td>
</tr>
<tr>
<td>• Chronic hypertension—controlled on medications&lt;sup&gt;b&lt;/sup&gt;</td>
<td>37–39 weeks</td>
</tr>
<tr>
<td>• Chronic hypertension—difficult to control (requiring frequent medication adjustments)&lt;sup&gt;b&lt;/sup&gt;</td>
<td>36–37 weeks</td>
</tr>
<tr>
<td>• Gestational hypertension&lt;sup&gt;c&lt;/sup&gt;</td>
<td>38 weeks</td>
</tr>
<tr>
<td>• Preeclampsia—severe</td>
<td>At diagnosis (recommendations limited to pregnancies at or after 34 weeks)</td>
</tr>
<tr>
<td>• Preeclampsia—mild</td>
<td>37 weeks</td>
</tr>
<tr>
<td>• Eclampsia</td>
<td>At detection</td>
</tr>
<tr>
<td>• Diabetes—gestational well-controlled&lt;sup&gt;d&lt;/sup&gt;</td>
<td>39–40 weeks</td>
</tr>
<tr>
<td>• Diabetes—pregestational with vascular disease</td>
<td>37–39 weeks</td>
</tr>
<tr>
<td>• Diabetes—gestational, poorly controlled</td>
<td>34–39 weeks (individualized to situation)</td>
</tr>
<tr>
<td>• Diabetes—gestational well-controlled on diet&lt;sup&gt;d&lt;/sup&gt;</td>
<td>Induction before 40 weeks not recommended</td>
</tr>
<tr>
<td>• Diabetes—gestational well-controlled on medication&lt;sup&gt;d&lt;/sup&gt;</td>
<td>39–40 weeks</td>
</tr>
<tr>
<td>• Diabetes—gestational poorly controlled on medication&lt;sup&gt;d&lt;/sup&gt;</td>
<td>34–39 weeks individualized to situation</td>
</tr>
<tr>
<td>• Cardiac disease</td>
<td>39 weeks (individualize depending on type and severity)</td>
</tr>
</tbody>
</table>

(Continued)
The risks and complications of induction of labor should be weighed against the possible benefits. A successful induction has been defined in many different ways but usually is one that achieves an uncomplicated vaginal delivery within 24 hours. If active phase is not achieved within 24 hours, this is not a reason per se for CD. Compared with a shorter induction-to-delivery interval, an induction lasting greater than 24 hours is associated with a higher risk for adverse outcomes, with a higher (e.g., 50%) risk of CD [13]. Neonatal outcomes including intensive care unit (ICU) admission, Apgar <7, and arterial cord pH <7 do not appear to increase with a prolonged latent stage as long as fetal status is reassuring [8,9]. A minimum of 24 hours should be allowed after cervical ripening and oxytocin administration (optimally with membranes ruptured) prior to diagnosing a failed induction [13–15]. After appropriate counseling, this time period may be extended assuming a reassuring fetal status.

### Table 21.1  Common Indications for Possible Induction of Labor or Cesarean Delivery (Suggested Timing of Delivery for Selected Conditions) (Continued)

<table>
<thead>
<tr>
<th>Obstetrical Issues</th>
<th>Gestational Age* for Delivery</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Prior stillbirth—unexplained</td>
<td>39 weeks</td>
</tr>
<tr>
<td></td>
<td>Consider amniocentesis for fetal pulmonary maturity if delivery planned before 38 weeks</td>
</tr>
<tr>
<td>• Spontaneous PPROM</td>
<td>34 weeks</td>
</tr>
<tr>
<td>• Spontaneous preterm labor</td>
<td>Delivery at 34–36 weeks if progressive labor or additional maternal/fetal</td>
</tr>
<tr>
<td>Prolonged pregnancy</td>
<td>≥41 weeks</td>
</tr>
</tbody>
</table>

*Gestational age is in completed weeks, thus “34 weeks” includes 34 weeks and 0 days through 34 weeks and 6 days.

Table 21.2  Contraindications to Induction of Labor

- Transverse or oblique fetal lie
- Umbilical cord prolapse
- Previous classical uterine incision or transfundal uterine surgery (e.g., from myomectomy)
- Placenta or vasa previa
- Active genital herpes infection
- Any contraindications to vaginal delivery, or indication for cesarean delivery

### Table 21.3  Contraindications to Induction of Labor (Continued)

- Any contraindications to vaginal delivery, or indication for cesarean delivery

### Table 21.4  Contraindications to Induction of Labor (Continued)

- Any contraindications to vaginal delivery, or indication for cesarean delivery

### PREVENTION

A routine (i.e., performed on every pregnant woman) early (e.g., before 20 weeks) ultrasound examination is associated with a 39% reduction in the incidence of postterm pregnancies and rates of induction of labor for postterm pregnancy, by allowing a more precise estimation of exact gestational age. An ultrasound performed in the first trimester (6–14 weeks) provides the best estimate of gestational age and the most benefit in terms of avoiding induction for postterm pregnancy [16] (see also Chapter 4).

### CRITERIA FOR INDUCTION

**Workup/Counseling**

Gestational age should be documented accurately before considering induction, to avoid inadvertent postterm and preterm deliveries. Indications and contraindications need to be carefully reviewed. Counseling with the patient should include discussion of specific indications, risks (possible complications), and benefits of induction.

### Indications

Once a term gestation has been confirmed, possible **indications for induction and suggested best timing** are shown in Table 21.1 [17,18].

For more details on the indications, see each specific guideline (e.g., Chapter 27 “Postterm Pregnancy,” Chapter 19 “Preterm Premature Rupture of Membranes,” Chapter 14 “Trial of Labor After Cesarean” in this volume; and Chapter 4 “Pregestational Diabetes,” Chapter 5 “Gestational Diabetes,” Chapter 46 “Fetal Macrosomia,” Chapter 44 “Multiple Gestations,” and Chapter 10 “Intrahepatic Cholestasis of Pregnancy” in Maternal-Fetal Evidence Based Guidelines). The term **elective induction** should be avoided, as an induction should be performed upon a precise and accepted indication [19]. An induction based solely on maternal request should be designated as such.
**Gestational Age of Induction**

The gestational age at which the induction is being considered is very important. There are some indications for induction before 39 weeks (Table 21.1), but **without indications induction before 39 weeks should be avoided** [20]. In this chapter, induction and ripening in the third trimester, and usually at or near term, is reviewed. For second-trimester induction and ripening, see Chapter 55 in *Maternal-Fetal Evidence Based Guidelines*. For induction for gestational age ≥41 weeks, see Chapter 27.

**Induction without Medical Indication**

Some have advocated induction at 39 0/7–40 6/7 weeks, even without medical indications. In a meta-analysis including 11 randomized controlled trials (RCTs) and 25 observational studies evaluating pregnancies at 37 0/7–41 6/7 weeks, **compared with “elective” induction, expectant management was associated with higher incidence of CD (odds ratio [OR] 1.22, 95% CI 1.07–1.39), and higher incidence of meconium (OR 2.04, 95% CI 1.34–3.09), but otherwise similar maternal and perinatal outcomes** [11]. These results are driven mostly by data at 41 weeks or more, when induction is indicated (see Chapter 27). Another meta-analysis of 31 trials including 6248 women randomized to induction of labor between 37 0/7 and 41 6/7 weeks confirmed these results, with a reduced risk of CD (OR 0.83, 95% CI 0.76–0.92) and no difference in neonatal outcomes of 5 minute Apgar score <7, neonatal intensive care unit (NICU) admission, or perinatal death [10]. A meta-analysis restricted to five trials including women randomized to induction or expectant management from 39 0/7 to 40 6/7 showed no difference in CD or perinatal outcomes [21]. Another individual patient data (IPD) meta-analysis of advanced maternal age (AMA) women also showed no difference in CD rate, though the number of patients included was low (n = 367) [22]. A RCT of nulliparous women without medical indication for delivery were randomized to induction or expectant management with a primary outcome of increase in CD rate by 50%. Although this increase was not statistically significant, the rate of CD was higher in the induction group (30.5% vs. 17.7%) (relative risk [RR] 1.72, 95% confidence interval [CI] 0.97–3.06) [23]. **There are insufficient data to evaluate the safety and effectiveness of induction without a medical indication at 39–40 6/7 weeks**, though the data from these meta-analyses [21,22] are useful when counseling women considering induction by maternal request. The Maternal-Fetal Medicine Units Network is currently conducting a study randomizing nulliparous women to induction or expectant management at 39 weeks (ARRIVE [NCT01990612]) [24]. Results from this trial may have significant effects on management of these women at term.

**Contraindications**

Induction of labor is **contraindicated** in the situations shown in Table 21.2 [20]. The attending physician should use his or her own discretion in the event of multifetal pregnancy, maternal heart disease, prior low-transverse CD, severe hypertension, and abnormal fetal heart rate (FHR) patterns not necessitating emergent delivery.

**Induction of Labor after Cesarean**

The **risk of uterine rupture** after induction in women with a prior CD deserves special attention (see Chapter 14). Misoprostol induction in women with a prior CD was associated with a 5.6% risk of uterine rupture in one of the largest series [25]. Therefore, **misoprostol should not be used for cervical ripening or labor induction in women with prior uterine incisions except under very unusual circumstances or management of stillbirth in the second trimester** (see Chapter 55 in *Maternal-Fetal Evidence Based Guidelines*). According to retrospective studies, using PGE2 for cervical ripening in women who have a history of previous cesarean also increases the risk of uterine rupture [26–29]. Risk of uterine rupture is approximately **1.4–2.5 with PGE2 use** (with or without oxytocin) [28,29], and about **1.1 with oxytocin alone** [29]. Alternatively, cervical ripening with the Foley catheter is **not associated with any additional risk of uterine rupture in patients undergoing a trial of labor after cesarean section** [30]. A patient who has a prior CD, no previous vaginal delivery, and an unfavorable Bishop score up to 39–40 weeks has more risks (e.g., sepsis, uterine rupture, and hysterectomy) from induction of labor; these women may elect for repeat CD after counseling [26–29] (Chapter 14). However, in a prospective observational trial of 12,676 women undergoing either induction of labor or expectant management after a prior CD, induction at 39 weeks remained associated with a significantly higher chance of vaginal birth after cesarean (VBAC) as well as uterine rupture (OR 1.31, 95% CI 1.03–1.67; and OR 2.73, 95% CI 1.22–6.12, respectively) [31]. Uterine rupture was not statistically different in those women induced at 40 completed weeks compared with expectantly managed women, however there was also no increase in VBAC odds [31].

**Time of Day for Induction**

Spontaneous labor has been shown to have a circadian rhythm with a higher occurrence at night due to a higher concentration of myometrial oxytocin receptors and maternal oxytocin concentrations. A meta-analysis of three RCTs was performed, however results were not pooled secondary to substantial heterogeneity between studies [32]. One RCT comparing morning and evening oxytocin inductions showed no difference in outcomes for women induced in the morning versus the evening, except for slightly increased neonatal admission to the maternity ward, medium care neonatal unit, or ICU (RR 1.48, 95% CI 1.02–2.14) in the morning group, though the authors could not explain a reason for this effect [33]. The remaining two RCTs compared prostaglandin induction timing, and showed no differences in neonatal outcomes or CD, but higher maternal satisfaction in morning inductions [32]. There were less operative vaginal delivery in the morning nulliparas group in one trial, and less multiparas delivered during the evening, without an effect on nighttime deliveries [34].

**Prediction of Successful Induction: Maternal Characteristics**

There are certain maternal characteristics that have been shown to favorably predict successful induction of labor. A secondary analysis of a study investigating vaginal misoprostol versus vaginal dinoprostone showed that prior vaginal delivery, lower maternal BMI, tall maternal stature, and neonatal birth weight <4000 g were associated with a higher likelihood of a vaginal delivery, independent of method. Race was also predictive, with Hispanic race being a positive predictor of successful induction and African-American race being associated with a lower likelihood of successful induction of labor [35].

**Bishop Score**

In 1964, Bishop reported that his pelvic score (Table 21.3) is inversely proportional to the time from examination to time at
which spontaneous labor begins [6]. Unfavorable (<5) Bishop scores at admission for induction of labor are associated with two- to threefold increased risk of CD when compared with spontaneous onset of labor [6,7,36]. Data show a score of ≥9 to predict a short time until onset of spontaneous labor and, therefore, indicate favorability for induction [35]. A simplified Bishop score might also be considered [37]. In a meta-analysis of mostly cohort studies aimed primarily to assess Bishop score as a predictor of cesarean, scores of 4, 5, or 6 were not predictive. A Bishop score of 8 or 9 had a negative predictive value of 96% for cesarean [38]. An additional meta-analysis that included a larger sample of studies and randomized trials, higher Bishop scores were associated with a higher rate of vaginal delivery. A Bishop score of 4 had an OR of 2 for vaginal delivery, where a Bishop of 8 increased the odds of vaginal delivery by 5.5 [39]. A Bishop score of ≥9 is usually associated with a probability of vaginal delivery after labor induction similar to that after spontaneous labor [20].

Transvaginal Cervical Length
 Compared with using a Bishop score <5 for defining an unfavorable cervix, using a transvaginal ultrasound (TVU) cervical length (CL) >27 mm is associated with a reduced need for prostaglandins without affecting the outcome of the induction in one RCT [40]. However, in a systematic review of mostly observational studies, a CL of <30 mm did not predict vaginal delivery; a CL >30 mm did not predict cesarean delivery [41]. When compared with Bishop score, there was no significant difference in prediction of successful labor induction, mode of delivery, vaginal delivery within 24 hours of starting induction, or achievement of the active stage of labor [40,42]. Cervical wedging/ tunneling has been investigated in eight studies including 1139 patients and has a low sensitivity to predict a failed induction [42]. In nulliparas, a CL cut-off of 30 mm had a sensitivity of 70% and specificity of 74% for predicting cesarean [42].

Fetal Fibronectin
 Of five prospective observational trials [43–47], only one showed an association with positive fetal fibronectin and successful induction [43]. As such, fetal fibronectin is not recommended to predict which patient’s labor induction will result in a successful vaginal delivery.

Induction/Ripening Methods
 Induction of labor is one of the most-studied interventions in obstetrics, with hundreds of trials reported. Cervical ripening/induction agents can be functionally divided into two methods: mechanical and pharmacologic. Mechanical methods have been utilized since the days of Hippocrates, 460–360 BC. Many methods have been compared not only to placebo or no treatment but also among themselves, in different populations and clinical settings, making for an extensive experience.

Mechanical Methods
 Mechanical methods include the following: hygroscopic dilators (laminaria, lamicel, or dilapan); balloon (e.g., Foley catheter); and balloon with extra-amniotic infusion. Other methods reviewed under this category are membrane stripping and amniotomy.

Hygroscopic (Osmotic) Dilators (Laminaria, Lamicel, or Dilapan)
 Hygroscopic devices are made either from synthetic or from organic material. The laminaria are the organic hygroscopic devices, which are made from cold water seaweed. Under direct visualization facilitated by a vaginal speculum, laminaria are placed in the cervical canal. Once in the cervix, the device absorbs water from surrounding tissues causing it to slowly swell, dilating the cervix. Generally the laminaria are left in place for 6–12 hours.

Currently, no evidence exists to support using laminaria to decrease either the interval from induction to delivery or the rate of CD. However, their use may cause an increase in maternal endometritis and neonatal sepsis likely due to the inability to reliably ensure sterility in this organic product [48]. In attempt to avoid problems associated with lack of sterility, polyvinyl alcohol polymer–magnesium sulfate (lamicel) and polyacrylonitrile (dilapan) were developed as synthetic hygroscopic devices. Like the laminaria, they are inserted in the same manner and function by attracting water from the surrounding tissue to achieve cervical softening, effacement, and dilation. However, lamicel and dilapan may be delivered sterilely.

Compared with placebo/no treatment, laminaria are associated with similar incidence of CD [48]. Similarly, no differences were noted between lack of treatment and dilapan in one trial [49].

Compared with any prostaglandins (vaginal PGE2, intra-cervical PGE2, or misoprostol), laminaria are also associated with similar incidence of CD, but less tachysystole with FHR changes. Serious maternal or perinatal morbidity is infrequent [48].

Compared with oxytocin, laminaria are associated with similar incidence of CD [48].

Compared with extra-amniotic induction, laminaria are associated with similar outcomes [48].

Compared with prostaglandin alone, the addition of laminaria is not associated with significant benefit [48].

Compared with oxytocin alone, there is insufficient evidence (one trial only) but no evidence of benefit from the addition of laminaria [48].

Balloon Catheter
 In 1853, Kraus first described a balloon device for preinduction cervical ripening. Much like the placement of laminaria, a vaginal speculum is often utilized to insert the Foley catheter in the cervical os; however, placement via a digital examination is becoming more common. Optimally, the catheter is placed at a level above the internal cervical os sometimes with the assistance of forceps or a clamp to guide the catheter. The

<table>
<thead>
<tr>
<th>Score</th>
<th>Dilation (cm)</th>
<th>Effacement (%)</th>
<th>Station</th>
<th>Cervical Consistency</th>
<th>Position of Cervix</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Closed</td>
<td>0–3</td>
<td>−3</td>
<td>Firm</td>
<td>Posterior</td>
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<tr>
<td>1</td>
<td>1–2</td>
<td>40–50</td>
<td>−2</td>
<td>Medium</td>
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<td>2</td>
<td>3–4</td>
<td>60–70</td>
<td>−1,0</td>
<td>Soft</td>
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<td>3</td>
<td>5–6</td>
<td>80</td>
<td>+1,2</td>
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Foley catheter affects cervical ripening in two ways: (1) gradual mechanical dilation and (2) separation of the decidua from the amnion, stimulating prostaglandin release. Many studies have demonstrated the Foley catheter to be an effective tool for achieving a favorable cervix [48,50–55].

There is some evidence to assess the type of Foley catheter to use. Various sizes and balloon capacities have been investigated and used; these include a range from 25- to 80-mL balloons with 14- to 18-F catheters. Filling the balloon to 80 mL versus 30 mL resulted in an increased rate of delivery within 24 hours, increase in vaginal delivery rate, decreased need for oxytocin, and higher rate of postripening dilation of 3 cm or more [56–58].

Once placed above the internal os, the operator uses sterile saline or water to inflate the catheter’s balloon. Correct placement is verified by gentle traction on the catheter until the inflated balloon meets the resistance of the internal os. Once the location is verified, gentle traction is applied by taping the distal end of the catheter to the patient’s inner thigh. Adding weighted traction has not been shown to increase vaginal delivery overall, vaginal delivery within 24 hours, or decrease CD rate [59]. The cervix dilates as the balloon is expelled. After expulsion, a favorable Bishop score is most often achieved and induction may begin. If the Foley is not expelled within 12 hours, consideration should be given to removing it, as leaving it in 24 hours is associated with longer inductions [60].

Compared with no treatment, one trial reported Foley catheters to have no effect in the risk of CD [48].

Compared with all LAPG, including PGE2 and misoprostol, the rate of CD is the same in a meta-analysis of 21 studies (n = 3202) [48]. There was no significant difference in vaginal delivery in 24 hours in the Foley catheter group [48]. LAPG were associated with a higher rate of excessive uterine activity and Foley catheter was associated with an increased need for oxytocin augmentation in labor, according to a meta-analysis of nine trials [48]. Obviously this analysis is not very clinically helpful, as comparison group should not mix PGE1 and PGE2.

Compared with vaginal misoprostol, transcervical Foley catheter has been demonstrated to be equivalent for cervical ripening. There is a similar risk of CD, chorioamnionitis, and other maternal and perinatal outcomes, except a higher incidence of tachysystole associated with misoprostol (RR 2.9, 95% CI 1.39–5.81) [55,61–63]. Foley was associated with a longer time to from induction-to-delivery in two meta-analyses [62,64]. In a network meta-analyses of 51 RCTs, vaginal misoprostol reduced the risk of vaginal delivery not achieved in 24 hours (RR 0.43, CI 95% 0.35–0.67) and cesarean section (RR 0.84, 95% CI 0.71–0.99), but Foley catheter reduced the risk of hyperstimulation (RR 0.15, 95% CI 0.07–0.3) [64].

Compared with dinoprostone (intracervical and vaginal), Foley catheter had similar rates of vaginal delivery not achieved in 24 hours and CD. Foley catheter was significantly less likely to cause hyperstimulation than vaginal dinoprostone (RR 0.27, 95% CI 0.12–0.52), but not intracervical dinoprostone [64].

Compared with use with oxytocin for cervical ripening, Foley use without oxytocin was associated with a lower rate of CD (RR 0.57, 95% CI 0.38–0.88) [48]. There was no difference in tachysystole with or without FHR changes between groups [48]. There is insufficient evidence to support concurrent use of Foley catheter with oxytocin [48,65].

Compared with prostaglandin alone, the addition of a Foley catheter to locally applied PGE1 and PGE2 increases the likelihood of vaginal delivery within 24 hours, in three trials, with a lower likelihood of observing no cervical change when a balloon catheter is used [48]. There was no difference in CD between the groups in a meta-analysis of eight trials [48]. Additional RCTs performed after the previous meta-analysis [48] again showed shorter time to delivery, but again no difference in mode of delivery [66,67].

Foley bulbs were equally effective ripening agents in both outpatient and inpatient settings per the results of one trial [68], but data is insufficient to fully assess outpatient Foley balloon safety and effectiveness.

In women with a singleton gestation who undergo term induction with the intracervical Foley catheter, two trials showed conflicting results regarding length of labor and risk of CD after early amniotomy. There is insufficient evidence to support early amniotomy after Foley induction [69,70].

Theoretical risks associated with Foley catheter use include bleeding, fever, displacement of the presenting part, and premature rupture of membranes (PROM); however no randomized trial has shown an increase in these complications in comparison to other methods. Foley should not be used in women with low-lying placenta. Overall, the Foley catheter is an inexpensive, safe, well-tolerated, and easy tool for cervical dilation [50–54]. In a recent review of over 1200 low-risk women who received the intracervical Foley catheter for cervical ripening, there were no adverse events necessitating delivery in the preinduction ripening period [71]. In a meta-analysis of 26 trials including 5563 women, there was no increased risk of infectious morbidity with Foley catheter use [72]. Foley is as effective as other methods, including misoprostol, and possibly safer than pharmaceutical methods, and should be considered as first line in all inductions (Figure 21.1), including those with PROM (see Chapter 20).

Extra-Amniotic Saline Infusion (EASI)

EASI involves, as the term states, infusing usually saline through the cervix by a Foley balloon. For infusion of PGE2, see section “Extra-Amniotic Prostaglandins.”

Compared with LAPG, EASI showed no difference in risk to delivery >24 hours, CD, or tachysystole (six studies, n = 747) [48].

Compared with PGE2, EASI with an intracervical Foley balloon is associated with shorter time intervals to yield a favorable Bishop score, and similar incidence of CD [48,73–75].

Compared with vaginal misoprostol, EASI with an intracervical Foley balloon and oxytocin is associated with shorter induction-to-delivery interval in one trial, but no difference in another [76]. There was similar incidence of CD in both trials [48,76,77].

Compared with laminaria, EASI with an intracervical Foley balloon and oxytocin is associated with shorter induction-to-delivery interval and less CD for failed induction in one trial [77].

Compared with Foley only, EASI with Foley catheters has been associated with shorter induction-to-delivery interval in one RCT [78] but not in another larger RCT [79].

In summary, there is insufficient evidence to use EASI instead of Foley balloon for cervical ripening.

Double-Balloon Foley Catheters

Double-balloon Foley catheters (Cook® Catheter or Atad catheters) have become more frequently used for mechanical labor

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Double-Balloon Foley Catheters

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The catheter is designed specifically for labor induction, providing an intra-cervical and vaginal balloon to compress the cervix in addition to membrane separation to ripen the cervix. Both balloons in the Cook catheter are approved to fill to a volume of 80 mL in each balloon.

Compared with PGE2, no difference in CD rates were reported in five RCTs [80–83]; one RCT of 126 women found a higher rate of CD in the dinoprostone group [84]. Time from vaginal delivery was shorter in the double-balloon group in two of four RCTs reporting on the outcome [81–84] and delivery within 24 hours was increased for the double-balloon group in two of three trials reporting the outcome [82–84]. Overall maternal and neonatal morbidity was similar in both groups, with the exception of tachysystole being more common with PGE2 [84]. There is insufficient evidence to support the use of double-balloon catheter over PGE2 for labor induction.

Compared with single-balloon Foley catheter, there was no difference in vaginal delivery within 24 hours, CD rate in two trials [81, 85]. Time from induction to delivery was longer in the double-balloon group in one trial [81]. There were more operative deliveries in the double-balloon group in one RCT [79] but not the other [81]. There is a significant cost difference between the catheters: single-balloon Foley catheters may cost up to $12–14 (the United States) for large balloon catheters (75 mL), but are cheaper when 30 mL balloons are used. The double-balloon catheter costs Labor and Delivery units $45-$200 in the United States per unit. There is currently insufficient evidence to support the use of double-balloon catheters over single-balloon catheters for the induction of labor. Until further information is available, a Foley catheter should be used over double-balloon catheter for both efficacy and economic concerns.

Membrane Stripping (or Sweeping)

Membrane stripping is the practice of inserting a finger through the internal os and sweeping to separate the membranes from the lower uterine segment. This technique stimulates prostaglandin release as plasma prostaglandin levels have been observed to increase poststripping. Sweeping the membranes promotes the onset of labor.

Compared with no sweeping, sweeping of the membranes, performed as a general policy in women at term (e.g., weekly starting at 38 weeks), is associated with reduced duration of pregnancy and reduced frequency of pregnancy continuing beyond 41 weeks (RR 0.59, 95% CI 0.46–0.74) and 42 weeks (RR 0.28, 95% CI 0.15–0.50) [86–90]. To avoid one formal induction of labor, sweeping of membranes must be performed in eight women. Rate of CD and maternal or neonatal infection are similar. Discomfort during vaginal examination and other adverse effects (bleeding and irregular contractions) are more frequently reported by women allocated to sweeping, but not associated with complications. Studies comparing sweeping with prostaglandin administration are of limited sample size and do not provide evidence of benefit.

When used as a means for induction of labor, the woman should be counseled that her chance of going to spontaneous labor after one sweeping at term is about 36% in the next 48 hours, versus 17% without sweeping (so doubling the rate of onset of labor) [86,87]. Possible complications such as bleeding, infection, and ruptured membranes are not found to be increased with stripping [86,87].
In nulliparas being induced with PGE2 and oxytocin, the addition of membrane sweeping is associated with shorter induction-to-delivery interval and increased vaginal delivery rates in one trial [88]. There were no differences noted in nulliparas with favorable cervixes or in multiparas.

In women attempting trial of labor after cesarean (TOLAC), weekly membrane sweeping had no effect on duration of pregnancy, spontaneous labor, or CD rate [91].

Amniotomy
Amniotomy—artificial rupture of the membranes—is another technique used in labor induction. There is insufficient evidence to assess the effectiveness of amniotomy alone [92]. No trials compared amniotomy alone with intracervical prostaglandins. If performed without cervical ripening or achieving a favorable cervix, amniotomy may be followed by long intervals before onset of labor. In induced patients, early amniotomy is associated with shorter duration of labor and no increase in CD rates in most studies [65–93-95]. One RCT of 168 patients showed longer duration of labor and CD rate in early amniotomy group [96], but this is contrary to findings in the aforementioned RCTs. The rate of intrapartum fever is mixed in RCTs and warrants additional research [93–97].

Compared with placebo, amniotomy and intravenous (IV) oxytocin are associated with significantly fewer instrumental vaginal deliveries than placebo. Compared with vaginal prostaglandins, amniotomy and IV oxytocin result in more postpartum hemorrhage and more dissatisfaction in women [65,92-97].

Pharmacologic Methods
Pharmacologic methods include the prostaglandins—E1: misoprostol; E2: dinoprostone; and F2α—as well as mifepristone, estrogen, relaxin, oxytocin, etc.

Misoprostol
Misoprostol (Cytotec™) is PGE1, which is an endogenously produced hormone that acts locally on surrounding tissues. Through complex molecular actions, PGE1 stimulates uterine contractions and cervical dilation in a manner akin to the onset of spontaneous labor. More specifically, PGE1 potentiates calcium ion transport across the cellular membrane and regulates cyclic adenosine monophosphate (AMP) within the uterine smooth muscle cells to trigger contractions. Additionally, PGE1 facilitates cervical ripening by stimulating the pathway leading to the activation of collagenases. Collagenases, in turn, break down the structural collagen network of the cervix yielding a softer, thinner cervix.

Although it is currently on the market as a 100-mg tablet to prevent peptic ulcers, misoprostol is available and widely used in an “off label” form for preinduction cervical ripening and induction. Misoprostol should not be used for cervical ripening or labor induction in women with prior uterine incisions (e.g., prior CD) after 28 weeks [20–25]. For use of misoprostol for induction in the second trimester, see Chapter 55 in Maternal-Fetal Evidence Based Guidelines. Misoprostol can be administered vaginally, orally (buccal), or sublingually.

Vaginal misoprostol suppository. Vaginal misoprostol is most commonly administered by placing a tablet in the posterior fornix of the vagina. Several studies have focused on the dose administered. Vaginal misoprostol in doses above 25 µg every 4 hour is more effective (higher success rate for vaginal delivery within 24 hours of induction, decrease need for oxytocin, and decrease induction-to-delivery intervals) than conventional methods of labor induction, but with more uterine hyperstimulation. Lower doses (25 µg) are similar to conventional methods in effectiveness and risks. Lower doses of misoprostol compared with higher doses were associated with more need for oxytocin augmentation and less uterine tachysystole, with and without FHR changes, and less meconium aspiration, and less CD for NRHF [72,98-99]. Therefore, 25 µg of misoprostol (one-quarter of a 100-µg tablet) given not more frequently than every 3–6 hours has been recommended by the American College of Obstetricians and Gynecologists [20].

Compared with placebo, misoprostol is associated with reduced failure to achieve vaginal delivery within 24 hours (RR 0.51, 95% CI 0.37–0.71), and with increased uterine tachysystole without FHR changes (RR 3.52, 95% CI 1.78–6.99) [98,99].

Compared with vaginal PGE2, intracervical PGE2, and oxytocin, vaginal misoprostol is associated with less epidural analgesia use, fewer failures to achieve vaginal delivery within 24 hours, and more uterine tachysystole [64,98].

Compared with vaginal or intracervical PGE2, oxytocin augmentation was less common with misoprostol, and meconium-stained liquor more common [97,98].

Compared with vaginal PGE2 only, a meta-analysis of 11 RCTs showed that vaginal misoprostol was associated with a higher likelihood of vaginal delivery within 12 and 24 hours and lower use of oxytocin augmentation. There was no difference between the groups in relation to rate of CD, or incidence of tachysystole. A second meta-analysis including only singleton pregnancies confirmed these results [98,100]. A network meta-analysis comparing intracervical PGE2 and vaginal misoprostol showed a reduction in hyperstimulation, decreased number of failed vaginal deliveries in 24 hours, and reduced risk of CD [64]. Additionally, the cost of a 25-mg pill of misoprostol is approximately $2 compared with the dinoprostone vaginal insert at approximately $168 [98].

Compared with oral misoprostol solution and all formulation of PGE2, a network meta-analysis including over 48,000 women showed that oral misoprostol solution had higher vaginal delivery rate in 24 hours and reduced risk of CD. High-dose vaginal misoprostol, then low-dose misoprostol, followed as the most effective treatments [99].

In summary, vaginal misoprostol 25 µg every 4–6 hours is a safe, effective means for cervical ripening, more effective than PGE2.

Vaginal misoprostol insert. A phase III clinical trial of a 50 and 100 µg vaginal insert (which is equivalent to a 25 µg tablet fragment being inserted every 6 hours) showed that the 100 µg vaginal insert had equal median times to vaginal delivery as the 10 mg dinoprostone vaginal insert. The 50 and 100 µg misoprostol inserts and the 10 mg dinoprostone insert all had comparable rates of CD and safety profile [101]. Misoprostol inserts of 50, 100, and 200 µg have similar rates of vaginal delivery within 24 hours, but higher rate of NRHF requiring CD and tachysystole in the 200 µg group [102]. A more recent RCT by the same authors compared misoprostol insert 200 µg to dinoprostone 10 mg insert, and showed shorter time to delivery without an increase in CD rate. However, tachysystole was more common in the misoprostol insert group [103]. In a secondary analysis, there was a higher rate of NRHF and CD secondary to NRHF in the high-dose misoprostol insert groups [103,104]. In summary, there is still insufficient evidence for clinical use of misoprostol vaginal insert.
Oral misoprostol. Studies that examine patient satisfaction have shown a definite preference toward oral administration [103,104]. Compared with placebo in women with PROM, women who received oral misoprostol had a higher likelihood to deliver vaginally within 24 hours (RR 0.16, 95% CI 0.05–0.49), needed less oxytocin (RR 0.42, 95% CI 0.37–0.49), and had a lower cesarean section rate (RR 0.72, 95% CI 0.54–0.95) [105] (see Chapter 20).

Compared with vaginal PGE2 prostaglandins, low-dose oral misoprostol (eg, 20 µg) administered every 2 hours had the same rate of vaginal delivery in 24 hours and a significantly lower rate of CD [105,106]. Women given oral misoprostol were also less likely to need a cesarean section (RR 0.88, 95% CI 0.78–0.99). There was some evidence that they had slower inductions, but there were no other significant differences [105].

Compared with oxytocin, oral misoprostol was associated with an increase in meconium-stained liquor in women with ruptured membranes (RR 1.65, 95% CI 1.04–2.6) but lower CD rate (RR 0.77, 95% CI 1.04–2.6) [105].

Compared with vaginal misoprostol, oral misoprostol had similar outcomes in regards to vaginal delivery in 24 hours and CD in a meta-analysis of 37 RCTs [105]. There were fewer babies with low Apgar score, postpartum hemorrhage in the oral misoprostol group, but higher rate of meconium-stained fluid [105]. A network meta-analysis, which allows for ranking treatments across studies through indirect comparisons, found that vaginal misoprostol had the highest probability of performing best among all prostaglandins to achieve vaginal delivery in 24 hours, but oral misoprostol avoided the most CD [99]. When the informal analysis of both achieving vaginal delivery in 24 hours and avoid CD was combined, low-dose (<50 µg) oral misoprostol performed best, followed by high-dose, then low-dose vaginal misoprostol [99]. A later network meta-analysis showed again that vaginal misoprostol avoided failed vaginal delivery in 24 hours the most, but was not different from oral misoprostol for reduced chance of CD [64]. Compared with vaginal misoprostol, low-dose titrated oral misoprostol (20–25 µg) given every 2 hours was shown in a review of two trials to cause fewer incidences of uterine tachysystole with FHR changes and a higher rate of vaginal delivery in 24 hours [106].

Sublingual misoprostol. Based on only three small trials, sublingual misoprostol appears to be at least as effective as when the same dose is administered orally [107].

Compared with vaginal misoprostol, a meta-analysis of five trials found no difference was found in vaginal delivery within 24 hours, tachysystole (when grouped according to dose), or CD [108]. There is inadequate data to assess safety, optimal dose, and side effects.

Prostaglandin E2 (Dinoprostone). PGE2 is an endogenously produced hormone that acts locally on surrounding tissues. Such effect is manifested in the smooth muscle of the uterus and gastrointestinal tract. PGE2 can be used for induction of labor via different routes of administration, such as vaginal, extra-amniotic, oral, and IV. The vaginal route is the most common route of administration of PGE2 for labor induction. It can be given in different forms, such as tablet, gel, and insert.

All PGE2 vaginal forms. Compared with placebo or no treatment, vaginal PGE2 (all forms) is associated with a higher likelihood of vaginal delivery within 24 hours (RR 0.32, 95% CI 0.02–4.83), no difference in CD rates, and an increase in the risk of uterine tachysystole with FHR changes (4.4% vs. 0.5%; RR 3.16, 95% CI 1.67–5.98) [109]. PGE2 tablet, gel, and pessary appear to be as efficacious as each other [109].

PGE2 vaginal gel. Dinoprostone gel (Prepidil) is packaged as a 0.5-mg dose in a 2.5-mL syringe. A shielded catheter is added to the syringe end to facilitate safe injection, usually intracervical. Under direct visualization using a speculum, the syringe contents should be injected into the endocervical canal using sterile technique. The patient should remain supine for 30 minutes to minimize leakage from the canal. An alternative method for administering the gel is to inject into the posterior fornix or intravaginal administration. Until achieving a favorable cervix, dinoprostone 0.5 mg may be repeated every 6 hours up to a maximum dose of 1.5 mg in a 24-hour period. Once the cervix is favorable, oxytocin may be initiated for induction 6 hours after the last dose.

Compared with placebo, intracervical PGE2 is associated with decreased risk of not achieving vaginal delivery within 24 hours (RR 0.61, 95% CI 0.47–0.79). There was a small, and statistically nonsignificant, reduction of the risk of cesarean section when PGE2 was used (RR 0.88, 95% CI 0.77–1.0). The finding was statistically significant in a subgroup of women with intact membranes and unfavorable cervix only (RR 0.82, 95% CI 0.68–0.98). The risk of tachysystole with FHR changes was not significantly increased (RR 1.21, 95% CI 0.72–2.05). However, the risk of tachysystole without FHR changes was significantly increased (RR 1.59, 95% CI 1.09–2.33) [109,110].

Compared with PGE2 tablets, PGE2 gel has similar rate of vaginal delivery not achieved in 24 hours, CD, hyperstimulation causing FHR changes, and oxytocin use [109].

PGE2 vaginal insert. PGE2 vaginal insert (Cervidil) (also called slow-release pessary) is a thin, vaginal insert containing 10 mg of dinoprostone and delivers roughly 0.3 mg of dinoprostone each hour over a 24-hour period. The insert is placed in the posterior fornix of the vagina and left in place until the desired ripening has occurred when the insert is removed. Removal should occur at least 30 minutes prior to starting oxytocin. Cervidil use is indicated for cervical ripening and induction of labor in patients who have a medical indication for induction at or near term.

Compared with PGE2 gel, a PGE2 insert is associated with an increased risk of not achieving a vaginal delivery within 24 hours (RR 1.26, 95% CI 1.12–1.41). There was no change in the risk of cesarean section (RR 1.07, 95% CI 0.93–1.22). The risks of tachysystole with FHR changes (RR 0.76, 95% CI 0.39–1.49) and without FHR changes (RR 0.86, 95% CI 0.56–1.15) were nonsignificantly different with the two methods of PGE2 administration. Only one trial with a small sample size reported on women's views, with no difference between groups [109–113]. The use of sustained-release PGE2 inserts is associated with a reduction in instrumental vaginal delivery rates (RR 0.47, 95% CI 0.32–0.68) when compared with vaginal PGE2 gel or tablet [109]. Compared with intracervical PGE2 low dose, intracervical PGE2 high dose has higher rate of hyperstimulation with FHR changes but no change in CD rate [109].

Compared with Cervidil followed by oxytocin, Cervidil started concurrently with oxytocin is associated with a shorter induction-to-delivery interval and higher incidence of vaginal deliveries within 24 hours in one small trial [114].

Extra-amniotic prostaglandins. There is insufficient evidence to fully assess the effectiveness of extra-amniotic prostaglandins for induction of labor, with enough evidence...
to discourage its use compared with other methods (see also the section “Extra-Amniotic Saline Infusion”). Extra-amniotic placement of prostaglandins was first undertaken in the early 1970s and has been largely replaced with cervical or vaginal placement. Most of the studies used PGE2 (the minority prostaglandin F2a [PGF2a]) and gel preparations. Of the primary outcomes, there were significantly fewer women delivered vaginally within 24 hours among those induced with extra-amniotic PGF2a compared with vaginal misoprostol (RR 2.43, 95% CI 1.42–4.15). No other differences between groups for primary outcomes were found to be statistically significant. Oxytocin was used to initiate or augment labor significantly less frequently with extra-amniotic prostaglandins when compared with placebo (RR 0.51, 95% CI 0.39–0.67) but significantly more frequently when compared with vaginal misoprostol (RR 1.73, 95% CI 1.20–2.49). When extra-amniotic PGE2 was compared with vaginal misoprostol, women preferred the treatment compared with vaginal PGE2. There were no other significant differences when extra-amniotic prostaglandins were compared with other methods of cervical ripening or induction of labor. Although this could suggest that extra-amniotic prostaglandins are as effective as other agents, the findings are difficult to interpret because they are based on very small numbers and may lack the power to show a real difference [115].

Oxymetazoline. Compared with placebo or no treatment, PGE2 is associated with a 54% decrease in CD. Otherwise, there were no significant differences between PGE2 and other interventions for this outcome [116].

Compared with vaginal prostaglandins, there is insufficient evidence, but no gross differences in three small trials [116].

Compared with all oxytocin treatments, oral PGE2 is associated with a trend for a lower incidence of vaginal delivery not achieved within 24 hours (RR 1.97, 95% CI 0.86–4.48). Oral prostaglandin was associated with vomiting across all comparison groups. There are no clear advantages to oral prostaglandin over other methods of induction of labor.

IV prostaglandins. IV prostaglandins should not be used for induction or cervical ripening, as they are no more efficient than IV oxytocin for the induction of labor, but their use is associated with higher rates of maternal side effects and uterine tachysystole. Compared with oxytocin, IV prostaglandins are associated with higher rates of uterine tachysystole both with and without changes in the FHR, and similar incidence of vaginal delivery [117]. Use of IV prostaglandins is also associated with significantly more maternal side effects (gastrointestinal, thrombophlebitis, and pyrexia). No significant differences emerged from subgroup analysis or from the trials comparing combination oxytocin/PGF2a and oxytocin or extra-amniotic versus IV PGE2 [117]. There is insufficient information to assess a combination of PGF2α and oxytocin compared with oxytocin alone or extra-amniotic and IV PGE2.

Oxytocin

In 1948, the posterior pituitary extract, oxytocin, was first used for labor induction via IV drip. Oxytocin was then synthesized by du Vigneaud and associates in 1953; this accomplishment won the Nobel Prize in chemistry in 1955. Oxytocin is now widely utilized worldwide. Oxytocin is routinely utilized as it is the drug of choice also for augmentation of labor. While induction of labor is the stimulation of contractions before the spontaneous onset of labor, augmentation is the stimulation of contractions in the face of inadequate contractions following the spontaneous onset of labor.

By increasing intracellular calcium concentration, oxytocin stimulates the smooth muscle cells of breast, vessels, and, moreover, the uterus. Receptors for oxytocin are expressed in cells of the endometrium, liver, pancreas, and breast tissue. After the 13th week of gestation, myometrial cells express oxytocin receptors as well. Peak expression by the myometrium and endometrium occurs at term. Oxytocin increases both the amplitude and frequency of contractions, making labor effective. When continuously administered IV, oxytocin affects uterine response within 1 minute. Steady-state plasma concentrations are obtained within 40 minutes.

Overall, comparison of oxytocin with either intravaginal or intracervical PGE2 reveals that the prostaglandin agents probably increase the chances of achieving vaginal birth within 24 hours. Oxytocin induction may increase the rate of interventions in labor. Women with ruptured membranes induction can be recommended by either method, and in women with intact membranes there is insufficient information to make firm recommendations [108].

Compared with expectant management, oxytocin alone inductions are associated with fewer women failing to deliver vaginally within 24 hours (8% vs. 54%; RR 0.16, 95% CI 0.10–0.25), with the CD rate slightly increased (RR 1.17, 95% CI 1.01–1.35). There is a significant increase in the number of women requiring epidural analgesia (RR 1.10, 95% CI 1.04–1.17) [119].

Compared with vaginal prostaglandins, oxytocin alone is associated with an increase in unsuccessful vaginal delivery within 24 hours (70% vs. 21%; RR 3.33, 95% CI 1.61–6.89), irrespective of membrane status, but there was no difference in cesarean section rates [118].

There was a small increase in epidurals when oxytocin alone was used (RR 1.09, 95% CI 1.01–1.17). Most of the studies included women with ruptured membranes, and there was some evidence that oxytocin decreased infection in mothers (chorioamnionitis; RR 0.66, 95% CI 0.47–0.92) and babies (use of antibiotics; RR 0.68, 95% CI 0.53–0.87). These data should be interpreted cautiously as infection was not prespecified in the original review protocol [118].

Compared with intracervical PGE2 prostaglandins, oxytocin alone is associated with an increase in unsuccessful vaginal delivery within 24 hours (50% vs. 35%; RR 1.47, 95% CI 1.10–1.96). For all women with an unfavorable cervix regardless of membrane status, the cesarean section rate is increased (19% vs. 13%; RR 1.37, 95% CI 1.08–1.74) [118].

Compared with vaginal prostaglandin F2α, there was no difference in CD rate (RR 1.19, 95% CI 0.65–2.18). The outcome of failing to delivery vaginally in 24 hours was not reported in any of the three RCTs included in the meta-analysis [118].

Oxytocin seems to be as effective as prostaglandins in women with PROM [118] (see Chapter 20).

Compared with low-dose oxytocin regimens (<100 mU in the first 40 minutes and <600 mU in the first 2 hours), there were no differences in rates of vaginal delivery not achieved within 24 hours (RR 0.94, 95% CI 0.78–1.14) or CD (RR 0.96, 95% CI 0.81–1.14). There was no difference in serious maternal or neonatal morbidity. No trials reported on the number of women who had uterine hyperstimulation
with fetal heart rate changes. When high bias studies were removed from the meta-analysis, there was a significant reduction of induction to delivery interval and increase in hyperstimulation without specifying fetal heart rate changes. Table 21.4 shows examples of each regimen (see also Chapter 7). There is insufficient evidence to support the use of high-dose oxytocin over low-dose protocols [119].

**Prostaglandin F2α**

**Compared with placebo,** vaginal PGF2α has a similar CD rate (RR 0.59, 95% CI 0.31–1.14). Vaginal delivery within 24 hours was not reported in the three trials included in a meta-analysis [109]. There were insufficient data to make meaningful conclusions for the comparison of vaginal PGE2 and PGF2α [109]. There is therefore insufficient data to assess the safety and efficacy of PGF2α for induction.

**Mifepristone**

**Compared with placebo,** mifepristone-treated women were more likely to be in labor or to have a favorable cervix at 48 hours (RR 2.41, 95% CI 1.70–3.42), were less likely to need augmentation with oxytocin (RR 0.80, 95% CI 0.66–0.97), and less likely to undergo cesarean section (RR 0.74, 95% CI 0.60–0.92), but more likely to have an instrumental delivery (RR 1.43, 95% CI 1.04–1.96) [120].

**Compared with oxytocin,** women treated with mifepristone for PROM after 36 weeks were less likely to have a vaginal delivery within 24 hours (RR 0.30, 95% CI 0.1–0.88) and their babies had an increased likelihood of neonatal adverse outcomes with more NICU admissions (RR 4.83, 95% CI 1.20–19.44), and abnormal fetal heart rate patterns (RR 5.63, 95% CI 1.11–28.52) in one trial [120].

There is insufficient information available from clinical trials to support the use of mifepristone to induce labor.

**Estrogen**

Several studies have shown that estradiol given via a variety of routes has the ability to achieve some degree of improved cervical ripening with minimal myometrial stimulation [121]. However, on a whole there were insufficient data to draw any conclusions regarding the efficacy of estrogen as an induction agent, given small, differing trials with different controls and different outcomes reported [122].

**Relaxin**

There is insufficient evidence to assess the safety and efficacy of relaxin as an intervention for induction of labor. There are no reported cases of uterine tachysystole with NRFHT in any of the four small trials [93]. Compared with placebo, relaxin is not associated with differences in CD [124].

**Dehydroepiandrosterone Sulfate**

Dehydroepiandrosterone sulfate (DHEAS) is converted to estrogen by the fetoplacental unit, and it was investigated as a possible mechanism for cervical ripening without myometrial contractions. There is insufficient evidence on its efficacy as one trial that investigated its use showed poor results when compared with placebo [123].

**Dexamethasone Sulfate**

There is insufficient evidence to assess the safety and efficacy of steroids as induction agents. Compared with oxytocin alone, dexamethasone IM and oxytocin are not associated with significant effects in maternal and perinatal outcomes in one small RCT [124]. Compared with Foley catheter, Foley catheter plus extra-amniotic infusion of dexamethasone solution had a significantly shorter time from induction to delivery in one small RCT (11.9 ± 3 hours vs. 14.5 ± 4.8 hours, p < .01 [125]).

**Hyaluronidase**

There is insufficient evidence to assess the safety and efficacy of intracervical injections of hyaluronidase for cervical ripening. It is not common practice, and it is an invasive procedure that women may find unacceptable in the presence of less invasive methods. In one RCT, when compared with placebo for cervical ripening, intracervical injections of hyaluronidase resulted in women receiving significantly fewer cesarean sections (RR 0.37, 95% CI 0.22–0.61), less need for oxytocin augmentation (RR 0.10 vs. 0.47%, RR 0.20, 95% CI 0.10–0.41), and increased cervical favorability after 24 hours (60% vs. 98%, RR 0.62, 95% CI 0.52–0.74). No side effects for mother or baby were reported in this trial [126]. Compared with Foley catheter, hyaluronidase was associated with a higher CD rate, and no difference in oxytocin use [127].

**Nitric Oxide Donors**

Nitric oxide (NO) donors do not appear currently to be a useful tool in the process of induction of labor. A meta-analysis including 10 studies compared NO donors with placebo, vaginal PGE2, intracervical PGE2, and vaginal misoprostol. There are very limited data available to compare NO donors to any other induction agent. There is no evidence of any difference between failed vaginal delivery within 24 hours, CD, or hyperstimulation [128]. There was an increase in nonserious maternal side effects (nausea, headache) [128].

**Other Methods**

**Acupuncture**

There is insufficient and conflicting evidence to assess the efficacy of acupuncture for induction of labor. Compared with a sham procedure, the use of acupuncture prior to a scheduled induction is not associated with any difference in the need for induction agents, CD, but had greater cervical change [129].

**Breast Stimulation**

**Compared with no intervention,** breast stimulation is associated with a significant reduction in the number of women not in labor at 72 hours (63% vs. 94%; RR 0.67, 95% CI 0.6–0.74). This result is not significant in women with an unfavorable cervix.

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**Table 21.4** Labor Stimulation with Oxytocin: Examples of Low- and High-Dose Oxytocin Protocols

<table>
<thead>
<tr>
<th>Regimen</th>
<th>Starting Dose</th>
<th>Incremental Increase (mU/minute)</th>
<th>Dosage Interval (Minutes)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low dose</td>
<td>0.5–1</td>
<td>1</td>
<td>30–40</td>
</tr>
<tr>
<td></td>
<td>1–2</td>
<td>2</td>
<td>15</td>
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<tr>
<td>High dose</td>
<td>–6</td>
<td>–6</td>
<td>15</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>6*</td>
<td>20–40</td>
</tr>
</tbody>
</table>

*The incremental increase is reduced to 3 mU/minute in presence of tachysystole and reduced to 1 mU/minute with recurrent tachysystole.*
The rate of postpartum hemorrhage is reduced (0.7% vs. 6%; RR 0.16, 95% CI 0.03–0.87). There is no significant difference in the cesarean section rate, in the rate of meconium staining or uterine tachysystole. The three perinatal deaths were associated just with breast stimulation (1.8% vs. 0%; RR 8.17, 95% CI 0.45–147.77) [130].

**Compared with oxytocin alone,** breast stimulation had similar rates of women not in labor after 72 hours, cesarean section rates, and meconium staining. Three of the four perinatal deaths were in high-risk women in the breast stimulation group (17.6% vs. 5%; RR 3.53, 95% CI 0.40–30.88) [130]. Until safety issues have been fully evaluated, breast stimulation should not be used in high-risk women [130].

**Castor Oil**

Castor oil should not be used for induction of labor [131]. There was no evidence of differences in caesarean section rates in the two trials reporting this outcome (RR 2.04, 95% CI 0.92–4.55). There were no data presented on neonatal or maternal mortality or morbidity. All women who ingested castor oil felt nauseous [131].

**Enemas and Baths**

There are no trials on enemas or baths for induction of labor.

**Homeopathy**

There is insufficient evidence to recommend the use of homeopathy (e.g., with caulophyllum) as a method of induction. No benefits were seen in the two small, poor-quality trials [132].

**Sexual Intercourse**

There is insufficient evidence to assess the efficacy of sexual intercourse for induction of labor. A “coital diary” prospective study showed no difference in the incidence of spontaneous labor in the “advised-coitus” group despite 1.5 times as many participants in that group reporting intercourse. There was also no difference in the rate of cesarean section or adverse outcomes [133]. One RCT is too small for meaningful guidance [134]. In another RCT, compared with no such advice, advice for coitus was associated with more coital activity (60% vs. 40%; RR 1.5, 95% CI 1.1–2.0), but similar rates of spontaneous labor (56% vs. 52%; RR 1.1, 95% CI 0.8–1.4) [135]. In a trial of 1175 women randomized to coitus-advised versus no advice, there were no differences in intervention to delivery, gestational age at delivery, or need for induction [136]. A limitation is that only 144 women returned their coital diary.

**ANTEPARTUM TESTING DURING CERVICAL RIPENING**

Fetal heart monitoring during cervical ripening depends on the agent used. There are no trials to assess the effectiveness and best modality for monitoring. In general, a nonstress test (NST) should be obtained before any induction or cervical ripening agent is used to assure fetal well-being. After administration of PGE2 gel or tablet, the fetal heart can be monitored continuously for about 0.5–2 hours, although the proper amount of time for monitoring is unclear. After administration of PGE2 insert, the fetal heart can be monitored continuously for the duration of the insertion [137]. After administration of misoprostol, the fetal heart should be monitored continuously, given the higher chance of contractions, and uterine tachysystole with related NRFHT.

**OUTPATIENT VERSUS INPATIENT**

Induction of labor in outpatient settings appears feasible, but the evidence is still insufficient for routine use. Important adverse events are rare [138]. There is insufficient evidence to know which induction agents are most effective and safe to use in outpatient setting. Studies examined vaginal and intracervical PGE2, isosorbide mononitrate (for these three agents there is most evidence of safety, n ≥500 each), Foley [71], vaginal and oral misoprostol, mifepristone, estrogens, and acupuncture (very limited evidence for these other agents) [139]. There is insufficient evidence to assess the safety of outpatient misoprostol for induction of labor. While effective in decreasing the length of gestation and induction-to-delivery interval, the safety of this approach, even at low (25 mg) doses, is still unproven in the three small trials [139–141]. There is insufficient data to assess the safety of outpatient dinosprostone for induction of labor. In a RCT including 827 women, the outpatient dinoprostone group had more NRFHT and was unable to be discharged home. Over half of the randomized women did not receive the allocated intervention secondary to labor or not requiring ripening [142].

There was no strong evidence that agents used to induce labor in outpatient settings had an impact (positive or negative) on maternal or neonatal health. There was some evidence that, compared with placebo or no treatment, induction agents reduced the need for further interventions to induce labor and shortened the interval from intervention to birth. There was no evidence that induction agents increased interventions in labor such as operative deliveries. In one cost analysis performed alongside a prospective RCT, inpatient prostaglandin gel did not have a significantly increased cost compared with outpatient Foley catheter, although outpatient Foley catheter was associated with fewer prelabor hours spent on the birthing unit [143]. A survey conducted along with an outpatient induction RCT found that women were willing to accept an extra 1.42 trips to hospital (2.42 trips total) and a travel time of 30.6 minutes per trip (73.3 minutes total) to be able to return to their own home while waiting for the priming to work, and overall preferred outpatient priming when the option was to return to their own home (OR 1.8, 95% CI 1.4–2.1, p < .0001) [144].

**Labor Management with Induction**

The patterns by which labor progresses in spontaneous labor and electively induced labor are significantly different [36]. Latent and early active phases proceed slower than a spontaneous labor in induced labor in which cervical ripening was necessary. Induction not requiring cervical ripening may be associated with a quicker labor course from 4 to 10 cm [36]. The risk of CD is increased during the first stage of labor of an induction needing cervical ripening, mainly because of dystocia. Induction without need for cervical ripening may have no or only a minor effect on the risk of cesarean [36]. Applying the same standards of spontaneous labor curves (e.g., Friedman's curve) to induced patients may lead to an increase cesarean section rate in induction (see Chapters 7 and 8).

When administering oxytocin, the target is to stimulate uterine activity that is sufficient to effect cervical change as well as fetal descent without compromising the fetus. Minimal criteria for effective uterine activity are 3 contractions per 10 minutes averaging greater than 25 mmHg above baseline, with 5 contractions in 10 minutes. The Montevideo unit was created in 1957 to describe the summation of the amplitudes of all contractions in a 10-minute window. Uterine tachysystole is defined as >5 contractions in 10 minutes. During induction with oxytocin, 91% of patients delivered vaginally achieved 200 Montevideo
units without neonatal morbidity in one retrospective study [145]. Contraction pressures of ≥200 Montevideo units should be targeted in induction or augmentation of laboring patients to achieve adequate labor [145,146]. Induced labor should be managed, in general, as for spontaneous labor (see Chapters 7 and 8). If active phase is not achieved within 24 hours, this is not a reason per se for CD. A failed induction should not be diagnosed until after 24 hours of oxytocin after membrane rupture in the active phase (usually 6 cm in a nulliparous patient), assuming reassuring fetal heart pattern [14,15,36].

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Intra-amniotic infection
Elizabeth Liveright and Sara Campbell

KEY POINTS
• Intra-amniotic infection (IAI) is diagnosed by maternal fever, and one or more other clinical criteria: maternal tachycardia (>100 beats per minute), leukocytosis, fundal tenderness, foul-smelling amniotic fluid, and/or fetal tachycardia (>160 beats per minute), in the absence of other causes of fever.
• Rates of IAI increase with duration of labor and rupture of membranes (ROM) greater than 24 hours.
• Rates of cesarean delivery are higher in women with a diagnosis of IAI.
• The clinical diagnosis of IAI may not correlate with histologic findings in the placenta.
• IAI is associated with increased maternal and neonatal morbidity.
• Antibiotics (e.g., ampicillin and single dose gentamicin) should be given at the time of intrapartum IAI diagnosis.
• Antibiotics treatment can in general be stopped with vaginal delivery. For cesarean, anaerobic coverage (clindamycin or metronidazole) should be added intrapartum, and one more dose of triple antibiotics given postpartum.
• Neonates delivered after IAI should be evaluated and treated as per Centers for Disease Control and Prevention (CDC) and American Academy of Pediatrics (AAP) guidelines. In general, if IAI happens before the birth of a neonate <34 weeks, workup and antibiotic treatment are recommended. In neonates ≥34 weeks born with IAI, workup and treatment with antibiotics are recommended only for ill-appearing babies, while most can be instead closely observed in the nursery. Antibiotics are usually continued for about 48 hours, and stopped when culture results return as negative.
• There are several preventive techniques shown to decrease rates of IAI. Some of these include, among others, antibiotic prophylaxis in premature rupture of membranes, as well as limiting the length of labor and the number of vaginal examinations.

DEFINITIONS/DIAGNOSIS
A wide range of terminology is used in the literature to describe intrapartum infection. The term “chorioamnionitis” has traditionally been used to describe infection of the amniotic fluid, placenta, membranes, and/or umbilical cord; however IAI may be a more accurate representation of the clinical picture. The clinical and histological presentation of IAI is heterogeneous, and its association with maternal and neonatal morbidity varies widely. Most studies utilize a combination of maternal fever, tachycardia (>100 beats per minute), leukocytosis, fundal tenderness, foul-smelling amniotic fluid, and fetal tachycardia (>160 beats per minute) to diagnose IAI (Table 22.1). Maternal fever is the most important and identifiable sign [1]; however, isolated maternal fever may be from an extra-uterine source, or may not be infectious in etiology. A well accepted definition is maternal fever (>100.4°F) with at least one more of these criteria: maternal tachycardia (>100 beats per minute), leukocytosis, fundal tenderness, foul-smelling amniotic fluid, and/or fetal tachycardia (>160 beats per minute), in the absence of other causes of fever.

Ancillary signs have been identified and quantified in a number of studies. The presence of maternal tachycardia is reported in 50%–80% of cases of IAI, and fetal tachycardia in 40%–70% of cases [1]. Uterine tenderness and foul-smelling amniotic fluid are present in 70%–90% of cases, but tenderness may be masked by epidural anesthesia [2].

Traditionally, the presence of one to three of these clinical features, in addition to fever, is used for diagnosis, but there have been no universally accepted diagnostic criteria published in the literature to date. In fact, studies report that these clinical criteria used for IAI diagnosis correlate poorly (accuracy of each or the combination of clinical signs was 47%–58%) with positive bacterial polymerase chain reaction (PCR) or culture by amniocentesis [3].

EPIDEMIOLOGY
This chapter focuses on IAI at term (≥37 weeks). The incidence of IAI in preterm births may be as high as 40% in deliveries under 27 weeks’ gestation [2]. In term deliveries, however, the reported incidence is from 2% to 4%. The rate is higher in cesarean section, with as many as 12% of cesarean deliveries having a clinical diagnosis of IAI [4].

In women with premature (or prelabor) rupture of membranes (PROM), the reported incidence is higher. The Term PROM study showed a rate overall of 7% in women with rupture of membranes prior to active labor [5]. In women with PROM greater than 24 hours, the rates of infection may be as high as 40% [6].

PATHOPHYSIOLOGY
IAI is largely considered to be an acute inflammation that is due to an ascending polymicrobial infection after membrane rupture. In this setting, the most common bacteria implicated are ureaplasma, urealyticum, and mycoplasma. Other bacteria often isolated in amniotic fluid cultures are Gardnerella, bacteroides, group B streptococcus, and Escherichia coli [7]. Rarely, hematogenous spread is the source, as in the case of Listeria infection [8].

IAI can be diagnosed histologically based on placental evaluation; however, a clinical diagnosis may not be confirmed by histological studies. It has been reported that up to one-third of clinical diagnoses may not have corresponding histologic findings [9]. A number of grading systems have been proposed to describe the histologic severity of chorioamnionitis,
and they include depth and location of neutrophil infiltration of the placenta. There are a number of inflammatory markers that may be associated with the clinical syndrome and histologic diagnosis of IAI, including cytokines, such as interleukin-1 alpha (IL-1alpha), IL-1beta, IL-6, and IL-8, among others. It is suggested that microorganisms invading the amniotic space stimulate these inflammatory cytokines, which favors the migration of neutrophils and ultimately produces the histologic diagnosis of chorioamnionitis [10].

**RISK FACTORS**

Risk factors for developing IAI include: _longer duration of labor, prolonged rupture of membranes, meconium-stained amniotic fluid, use of internal fetal monitoring, group B strep colonization, nulliparity, bacterial vaginosis, and increased number of vaginal examinations_ [2,5,11–18]. For example, the International Multicentre Term Prelabor Rupture of Membranes Study (Term PROM) demonstrated that increasing number of digital vaginal examinations, longer duration of labor, and meconium stained amniotic fluid were the most commonly identified risk factors for developing infection [5]. More recent studies suggest that some of these previously identified risk factors are eliminated by controlling for confounders. Additionally, a meta-analysis of several randomized controlled trials (RCTs) demonstrated that the use of transcervical Foley catheters for induction of labor was not associated with increased infectious morbidity [19].

**COMPLICATIONS: MATERNAL**

Intrauterine infection is associated with increased risk of cesarean delivery and endomyometritis, as well as postpartum hemorrhage, _need for blood transfusion, intensive care unit (ICU) admission, and need for hysterectomy_ (Table 22.2). The duration of chorioamnionitis has been shown to be associated directly with blood transfusion and ICU admissions [4]. Fortunately, infection rarely lasts greater than 24 hours after delivery, particularly when given a dose of antibiotics postpartum (see “Management” section below). Treatment failure, defined as persistent fevers after receiving one postpartum dose of antibiotics, is reported to be from 2% to 6% [20].

The risk of cesarean section is approximately two- to threefold higher for a woman meeting clinical criteria for IAI [21,22]. The increased risk is thought to be due to decreased uterine contractility and subsequent dysfunctional labor. Decreased uterine contractility may also lead to higher rates of postpartum hemorrhage due to atony; risk of postpartum hemorrhage after vaginal delivery may be up to 80% more likely and 50% more likely after cesarean section [21,23].

**COMPLICATIONS: NEONATAL**

IAI is associated with increased rates of _bacteremia, sepsis, and mortality in neonates, as well as an increased risk of cerebral palsy_ [4,24]. Intrapartum fever has been shown to increase transient neonatal adverse effects as well as neonatal seizures and encephalopathy [25–28]. These risks are present in both term and preterm infants, though the relative risk is greater in preterm neonates. The overall relative risk for cerebral palsy in term neonates born to mothers with IAI is reported as 4.7 [29]. It is thought that the fetal inflammatory response is a contributing factor to cerebral palsy in infants of mothers with IAI. Studies show that elevated levels of inflammatory cytokines were present in the amniotic fluid of mothers of children with cerebral palsy [30].

**MANAGEMENT**

**Treatment of IAI**

Initiation of antibiotics at the time of diagnosis is the standard of care for treatment of IAI (Table 22.3). This recommendation is supported by several studies demonstrating decreased maternal and fetal morbidity with administration of intrapartum antibiotics. These studies most consistently have shown decreased rates of neonatal bacteremia and sepsis, and also decreased rates of maternal febrile morbidity and length of postpartum stay [8,25,31,32]. These data are exclusively from cohort studies; there have been no RCTs to date comparing antibiotics to no treatment or placebo for women with IAI [33].

The source of IAI is thought to originate in the vaginal flora; coverage is therefore recommended to target beta-lactamase producing aerobes and anaerobes. _Ampicillin and gentamicin_ are the antibiotics most commonly used when the diagnosis of IAI is made in labor, based largely on clinical consensus, in the absence of data supporting other regimens [33,34]. _Daily dosing of the aminoglycoside_ is equally effective as 8-hour dosing with the advantages of increased bactericidal effects, decreased nephrotoxicity, as well as decreased cost [35]. The two RCTs identified in a Cochrane Review showed no statistically significant difference in maternal or neonatal outcomes when clindamycin was added to ampicillin and gentamicin [31,36].

_for patients undergoing cesarean section, however, anaerobic coverage through antepartum addition of clindamycin or metronidazole to ampicillin and gentamicin is associated with fewer wound infections postpartum_[37].

_Antipyretics_ should be considered when maternal fever is present.

**Delivery**

IAI itself does not necessitate cesarean section, unless otherwise motivated by standard obstetric indications, though there is an increased rate of cesarean delivery in the setting of IAI, as described above. Time from initiation of antibiotic treatment to
delivery has not been shown to affect outcomes; it is therefore not recommended to expedite delivery for IAI alone [4,31].

**POSTPARTUM**

For women who had a vaginal delivery, antibiotics do not need to be continued postpartum, as shown by RCTs comparing antibiotics versus placebo [33]. The cure in these cases is the delivery itself.

In women who had a cesarean delivery, at least a single dose of antibiotics after delivery is recommended to decrease postpartum maternal morbidity, but prolonged treatment has not been shown to improve outcomes [20,38,39].

**NEONATAL MANAGEMENT**

Maternal diagnosis of IAI can be associated with serious implications even for well-appearing term infants. The Centers for Disease Control and American Academy of Pediatrics state that infants born to mothers with either suspected or clinically proven IAI require antibiotic treatment, as well as laboratory testing [40,41]. This often results in separation from mother, admission to neonatal intensive care units (NICU), and exposure to drug-resistant bacteria in the NICU setting. A “sepsis calculator” has been suggested as a tool to calculate need for treatment based on maternal risk factors (GBS status, intrapartum antibiotic treatment), as well as continuous variables (peak febrile temperature, duration of ruptured membranes) and infant’s clinical appearance (clinically ill, equivocal, or well-appearing) [42,43]. This model reduced the proportion of unnecessary laboratory testing and antibiotic treatment, while not missing any cases of culture positive sepsis [43].

In general, if IAI happens before the birth of a neonate <34 weeks, workup and antibiotic treatment are recommended. In neonates ≥34 weeks born to mothers with IAI, workup and treatment with antibiotics are recommended only for ill-appearing babies, while most can be instead closely observed in the nursery. There is limited evidence regarding the length of neonatal antibiotic therapy. Antibiotics are usually continued for about 48 hours, and stopped when culture results return as negative.

**PREVENTION**

There are several interventions associated with prevention of IAI.

Antibiotics prophylaxis for prevention of IAI in preterm, premature rupture of membranes (PPROM) has been consistently shown to yield benefit for the neonate [44] (see Chapter 19). Even in women with term PROM, antibiotics prophylaxis, especially in women expected to have latency from PROM to delivery >12 hours, has been associated with significant reduction in IAI and endometritis [44]. (Chapter 20)

Modifiable risk factors for IAI in term pregnancies present the opportunity for prevention, particularly length of ruptured membranes and length of labor. Techniques used to shorten active labor have been shown to decrease maternal infectious morbidity [45]. Decreasing length of labor inherently decreases other risk factors associated with IAI, such as increased number of digital examinations [11,18].

Perineal hygiene, in addition to prolonged supine positioning and digital examinations, are suggested to be contributing factors to IAI [46]. A Cochrane review, however, showed no evidence for the prevention of IAI and only a statistically insignificant trend toward a decrease in postpartum endometritis [47]. Prophylactic antibiotics to reduce vaginal infections have been shown to lower rates of bacterial vaginosis and trichomonas, without a decrease in risk of IAI or preterm birth [48]. Intrapartum antibiotic amniinfusion has not been shown to provide consistent improvement in maternal or neonatal outcomes [49].

**REFERENCES**


Meconium
Alessandro Ghidini

KEY POINTS
• Fetal passage of meconium is common (about 12%) at term and postterm, while rare (<1%) in the preterm period.
• Meconium-stained amniotic fluid (MSAF) is due to a combination of increased fetal bowel motility and decreased clearance of meconium by the fetus or placenta. In a minority of cases such effects are due to fetal hypoxia.
• Meconium aspiration syndrome (MAS) is frequently a misnomer, as the respiratory compromise in the fetus is often due to chronic processes (e.g., infection or hypoxia, acute or chronic) which also stimulate colonic activity and fetal gasping.
• Prevention of meconium passage and of MAS may be accomplished by reducing the rate of postterm deliveries.
• Amnioinfusion for MSAF is associated with improvements in perinatal outcome only in settings where facilities for perinatal surveillance are limited. Under standard perinatal surveillance, compared with no amnioinfusion, amnioinfusion for MSAF is not associated with significant reductions in MAS, perinatal mortality, or combined outcome perinatal death or severe morbidity.
• Oro- and nasopharyngeal suctioning before delivery of the shoulder does not decrease the incidence of MAS, need for mechanical ventilation for MAS, any other associated morbidities, or neonatal mortality.
• Routine endotracheal intubation at birth in meconium-stained neonates who are otherwise vigorous does not improve neonatal outcomes over routine resuscitation.

HISTORIC NOTES
Meconium is a term derived from the Greek “mekoni,” which means poppy juice or opium. Confirming previous clinical impressions, meconium passage was formally recognized to be associated with increased perinatal morbidity and mortality in the 1975 Collaborative Study of Cerebral Palsy.

DIAGNOSES/DEFINITIONS
Meconium is the intestinal content of the fetus and is variably composed of mucopolysaccharides, blood by-products, hair, and squamous cells. Diagnosis of MSAF is made clinically on the basis of appearance (greenish or brownish staining) or by histopathologic examination of the placenta. Particularly in the preterm (<33-week) gestation, a clinical impression of MSAF may be false and instead reflect staining by another mechanism (i.e., hemosiderin). The diagnosis of MAS is respiratory distress requiring supplemental oxygen usually in the first 4 hours of life in the presence of meconium in a neonate without other causes of respiratory distress, and classified as shown below.

EPIDEMIOLOGY/INCIDENCE
The incidence of MSAF increases with gestational age: It is <1% before 37 weeks, about 10% of 39- to 41-week gestations, 18% of >41-week gestations [1]. Meconium in placenta macrophages is documented in about 17%-19% of term placentas; clinical evidence of MSAF is present in about 12% (7%-22%) of term deliveries [1]. However there is a poor concordance between histologic and clinical evidence of meconium presence, probably due to persistence of old meconium in macrophages after its clearance from the amniotic fluid, as well new release of meconium by the fetus before uptake by placenta macrophages [2]. Moreover, there is discordance among histopathologists in the definition of presence of placental meconium [3]. MAS occurs in about 5% of MSAF cases, and, of these, approximately 4% die [4].

ETIOLOGY/BASIC PATHOPHYSIOLOGY
Although regular fetal bowel movements occur since the early second trimester [5], meconium does not become green (i.e., contains biliverdin pigments) until 22–24 weeks. In term and postterm fetuses, physiologic increased motility levels or pathologic increases mediated by fetal stress from hypoxia, infection, or cord compression may lead to meconium passage. MSAF may also be caused by decreased clearance of meconium by the fetus or placenta in the presence of hypoxia [1].

Aspiration of meconium can lead to respiratory compromise by causing a chemical pneumonitis, associated with inhibition of surfactant function, inflammation, and obstruction. However, respiratory compromise in the presence of meconium (i.e., MAS) is more commonly due to other processes (such as chronic or acute asphyxia, or intrauterine infection) than damage from meconium aspiration itself [6]. For example, hypoxia may stimulate colonic activity and fetal gasping, leading to meconium aspiration. In such cases, meconium is not causative of the respiratory compromise, but rather a manifestation of underlying chronic or acute processes leading to fetal compromise.

SYMPTOMS
Symptoms of neonatal MAS include respiratory compromise, with tachypnea, cyanosis, and reduced pulmonary compliance. In some cases, pulmonary hypertension develops [4].

CLASSIFICATION (OF MECONIUM ASPIRATION SYNDROME)
• Mild: Supplemental oxygen <40% for <48 hours
• Moderate: Supplemental oxygen ≥40% or for ≥48 hours
• Severe: Need for intubation (or primary pulmonary hypertension)
RISK FACTORS

Postterm pregnancy; acute fetal acidemia; intrauterine infection (as suggested by higher rates of histologic acute chorioamnionitis, clinical chorioamnionitis and endometritis, and neonatal sepsis in the presence of MSAF); placental dysfunction leading to chronic fetal hypoxia (e.g., fetal growth restriction, preclampsia, oligohydramnios); uterine hyperstimulation (e.g., with misoprostol); long labors (every 2-hour increase in duration of labor is associated with a 30% increase in risk of MSAF) [7,8].

The multiplicity of risk factors, and the possible coexistence of independent risk factors (e.g., meconium and oligohydramnios) may explain why MSAF has inconsistently been associated with lower umbilical artery pH at birth.

COMPLICATIONS

Because the intrauterine processes underlying accumulation of meconium can jeopardize the fetus, MSAF and MAS can be associated with increased risk of fetal acidemia, neonatal seizures, neonatal intensive care unit (NICU) admission, neonatal sepsis, respiratory distress, neonatal encephalopathy, cerebral palsy, and neonatal death [1,4,9–11].

PREGNANCY MANAGEMENT

Meconium at Genetic Amniocentesis

As discolored amniotic fluid before 24 weeks (i.e., at genetic amniocentesis) is not due to MSAF but rather to intra-amniotic bleed or infection, evaluation for such causes of discolored amniotic fluid may be considered. These may include for example microbiologic studies on amniotic fluid, Kleihauer-Betke test on maternal blood, careful sonographic examination for evidence of retroplacental or intra-amniotic bleeds; assessment of placental implantation by evaluating placental thickness, dimensions, echotexture, and cord insertion.

Meconium in Late Preterm and Early Term Fetus <39 Weeks

MSAF in the late preterm and term fetus <39 weeks should prompt evaluation for infection and fetal hypoxia as the finding is quite uncommon and unlikely to be physiologic at this gestational age.

Meconium at ≥39 Weeks

MSAF in the full-term or postterm fetus may reflect normal physiology and maturation of the gastrointestinal tract, but one should first exclude the possibilities of infection or hypoxia as etiologies. Progression in meconium consistency in labor from no/little meconium to presence of thick meconium, or occurrence of meconium in the setting of category II fetal heart rate tracings, should elicit particular concern as this is associated with higher rates of fetal acidemia and lower Apgar scores at 5 minutes [12–14].

PREVENTION

Prevention of meconium passage and of MAS may be accomplished by reducing the rate of postterm deliveries. Early ultrasound dating, stripping of membranes at ≥38 weeks, and induction of labor at 41 weeks decrease the incidence of postterm pregnancies (see Chapter 27).

MANAGEMENT TECHNIQUES USED IN THE SETTING OF MSAF

Fetal

Amnioinfusion

The efficacy of amnioinfusion to “dilute” meconium and reduce associated neonatal morbidity has historically been controversial. The most recent meta-analysis includes 14 total randomized controlled trials (RCTs) [15]. Results are reported separately for sites with standard versus those with limited perinatal surveillance. The main outcome in the RCTs was usually occurrence of MAS.

Under standard perinatal surveillance, compared with no amnioinfusion, amnioinfusion for MSAF was not associated with a reduction in MAS (relative risk [RR] 0.52, 95% confidence interval [CI] 0.26–1.06) or perinatal deaths (RR 1.00, 95% CI 0.29–3.45), or the combined outcome perinatal death or severe morbidity (RR 1.13, 95% CI 0.88–1.47). There was considerable heterogeneity among studies for several secondary outcomes, such as presence of heavy meconium staining, cesarean section for non-reassuring fetal testing, occurrence of fetal heart rate decelerations, 5-minute Apgar score less than 7, presence of meconium below the vocal cords, need for neonatal ventilation or NICU admission.

Under limited perinatal surveillance, compared with no amnioinfusion, amnioinfusion for MSAF was associated with a significant reduction in MAS (RR 0.17, 95% CI 0.05–0.52) and perinatal mortality (RR 0.24, 95% CI 0.11–0.53), as well as reductions in cesarean section for non-reassuring fetal testing, 5-minute Apgar score less than seven, neonatal ventilation or neonatal intensive care unit admission, and neonatal encephalopathy.

In summary, amnioinfusion for MSAF is associated with improvements in perinatal outcome in settings where facilities for perinatal surveillance are limited, but not in settings with standard perinatal surveillance. In general, amnioinfusion is offered at >34 weeks. There are many variations of the amnioinfusion technique, but a “typical” protocol calls for infusion via an intrauterine pressure catheter (obviously in a woman with dilated cervix and ruptured membranes) of 500 mL of normal saline over a period of 30 minutes, see also Chapter 10. For amnioinfusion in presence of variable decelerations, see Chapter 10; for amnioinfusion for oligohydramnios without preterm premature rupture of membranes (PPROM), see Chapter 57 in Maternal-Fetal Evidence Based Guidelines; for amnioinfusion for PPROM, see Chapter 19.

Antibiotics

There is insufficient evidence to assess the effectiveness of antibiotics for women with meconium in labor. A meta-analysis on the subject (inclusive of two studies, both of which utilized ampicillin-sulbactam) has shown that compared with normal saline, antibiotic prophylaxis in women with MSAF is associated with no statistically significant reduction in the incidence of neonatal sepsis (RR 1.00, 95% CI 0.21–4.76), NICU admission (RR 0.83, 95% CI 0.39–1.78), and postpartum endometritis (RR 0.50, 95% CI 0.18–1.38), but a significant decrease in the risk of chorioamnionitis (RR 0.36, 95% CI 0.21–0.62) [16]. No serious adverse effects have been reported.

Oro- and Nasopharyngeal Suctioning

Suctioning of the oro- and nasopharynx before delivery of the shoulder or the “first cry” does not decrease the incidence of MAS, need for mechanical ventilation for MAS, any other associated morbidities, or neonatal mortality [17,18].
Neonatal Endotracheal Intubation
A policy of routine endotracheal intubation at birth in meconium-stained babies who are otherwise vigorous does not improve neonatal outcomes over routine resuscitation [19]. For depressed or nonvigorous newborns, endotracheal intubation and suctioning may still be performed in infants born through MSAF [19].

REFERENCES
Malpresentation and malposition
Alexis Gimovsky

KEY POINTS
• Malpresentation is associated with uterine anomalies, fibroids, placenta previa, grand multiparity, contracted maternal pelvis, pelvic tumors, prematurity (the earlier the gestational age (GA), the higher the incidence of malpresentation), multiple gestation, polyhydramnios, short umbilical cord, fetal anomalies (e.g., anencephaly, hydrocephalus), abnormal fetal motor ability, and prior breech delivery.
• Complications of breech presentation include congenital anomalies, preterm birth (PTB), birth trauma, low Apgar scores and low pH, regardless of mode of delivery. Cord prolapse, head hyperextension, and head or arm entrapment are more common with vaginal breech delivery.
• External cephalic version (ECV) is a safe and effective intervention for malpresentation. Urgent cesarean delivery (CD) for nonreassuring fetal heart rate tracing (NRFHT) and placental abruption occur in <0.5% of ECV.
• ECV is to be avoided with any contraindications to vaginal delivery such as placenta previa, or prior classical uterine incision. ECV is relatively contraindicated in rupture of membranes, oligohydramnios, known uterine or fetal anomaly, unexplained uterine bleeding, or active phase of labor.
• ECV reduces the incidences of noncephalic birth by 54% and CD by 37%. Because ECV is associated with a very low incidence of adverse events and with a significant decrease in CD, all women at or near term with nonvertex presentations should be offered an ECV. Success rates range between 50% and 70%. Success is increased with higher parity, transverse or oblique lie, nonengaged presenting part, relaxed uterus, palpable fetal head, and maternal weight less than 65 kg.
• There is insufficient evidence to assess the best GA at which to perform ECV. Compared with ECV at term, ECV before term (e.g., 34–35 weeks) reduces noncephalic presentation at birth but does not reduce the rate of CD, and may be associated with an increase in the incidence of PTB. About 36 weeks is generally considered to be the optimal time for attempted ECV.
• Tocolytics with betamimetics prior to attempt at ECV is associated with fewer failures of ECV and less CDs.
• Anesthetic dose neuraxial blockade (usually with spinal) is associated with a 44% increase in the success rate of ECV.
• ECV should be performed, after appropriate counseling and consent, in a facility with ready availability for ultrasound and for emergency CD.
• There is inconsistent evidence that moxibustion at point BL67 alone or in combination with acupuncture, is associated with higher success rates of ECV, especially when performed in China.
• Compared with planned vaginal delivery, planned CD for the term breech fetus is associated with a decrease in perinatal death and a reduction in serious neonatal morbidity, but no difference in death or neurodevelopmental delay at 2 years after delivery.
• There is insufficient evidence to assess whether outcomes of the preterm breech presenting fetus are affected by mode of delivery.
• There is insufficient evidence to assess the best mode of delivery for the nonvertex second twin. Vaginal delivery of the second nonvertex twin by breech extraction is a reasonable management option by expert operator, possibly by breech extraction.
• There is insufficient evidence to assess any intervention for malposition.

DEFINITIONS
• Presentation: Fetal body part that is in the lower uterine segment (lowest in the uterus and closest to the cervix).
• Malpresentation: Fetus presenting with the fetal head not in the lower uterine segment.

MALPRESENTATION
Symptoms
Maternal impression of fetal presentation based on fetal movement is suggestive but overall unreliable for predicting fetal presentation.

Epidemiology/Incidence
Breech presentation complicates 3%–4% of all pregnancies at term (≥37 weeks) [1]. Its incidence is inversely proportional to GA, with an incidence of about 25% at 28 weeks, 11% at 32 weeks, and 5% at 34 weeks [2]. In 2003, 87% of all breech presentations resulted in CD; this is similar to 1990, (90%), but much increased from 1970 (12%) [3,4]. Breech as an indication for CD accounts for 15% of all cesarean deliveries and adds 1.4 billion dollars to U.S. obstetrical costs [4].

Classifications
Breech
Fetus presents in longitudinal lie with head not in the lower uterine segment.
Fetal breech presentation is further classified as follows:
• Complete—Flexion of the fetal hips and knees
• Incomplete—Extension of one or both hips (includes footling)
• Frank—Flexion at the hips and extension at the knees
Transverse
The fetal longitudinal axis is perpendicular to the long axis of the uterus. The fetus can either present “back up” (fetal small parts present to the cervix) or “back down” (fetal spine or shoulder present to the cervix).

Oblique
The fetal longitudinal axis is diagonal to the long axis of the uterus.

Face
The fetal head is hyperextended so that the fetal occiput is in contact with the fetal back and the mentum (chin) is presenting. The fetal chin may be anterior or posterior relative to the maternal pubic symphysis.

Brow
The presenting part is the portion of the fetal head between the orbital ridge and the anterior fontanelle. The fetal head is positioned midway between full flexion and extension.

Compound
Simultaneous presentation of a prolapsing fetal extremity and the presenting part.

Risk Factors/Associations
Both maternal and fetal factors can lead to malpresentation, including uterine anomalies, fibroids, placenta previa, grand multiparity, contracted maternal pelvis, pelvic tumors, prematurity (the earlier the GA, the higher the incidence of malpresentation), multiple gestation, polyhydramnios, short umbilical cord, fetal anomalies (e.g., anencephaly, hydrocephalus), abnormal fetal motor ability, and prior breech delivery. There is a 9% risk of recurrence of malpresentation in subsequent pregnancies following a prior breech delivery.

Complications
Incidence of congenital anomalies (up to 6%), PTB, birth trauma, low Apgar scores, and low pH are increased with breech presentation compared with vertex presentation, regardless of mode of delivery. Breech presentation may be both a sign and a consequence of fetal compromise, regardless of delivery mode. Incidence of cord prolapse is the same with frank breech as with vertex presentations (<1%), 5% with complete breech, 15% with footling breech, and is inversely proportional to GA. Head hyperextension (associated with spinal cord injury) and head or arm entrapment are associated with breech presentation, particularly with vaginal breech delivery. Presentation at birth does not seem to affect adult intellectual performance. Cesarean or vaginal delivery for breech presentation do not differ in terms of long-term adult intellectual performance [5].

Workup
Fetal presentation should be assessed by Leopold’s maneuvers at each prenatal visit starting at ≥ 34 weeks of gestation. If the clinician is unsure, a vaginal examination, or even better, if still unclear, an ultrasound may be indicated to assess fetal presentation.

External Cephalic Version
Definition
Procedure performed by application of pressure and maneuvers to the maternal abdomen with the goal to turn the fetus to a cephalic presentation, thus increasing the likelihood of vaginal delivery (Figure 24.1) [1].

Complications
While the rate of short-term fetal bradycardia can be as high as 20%, the rate of need for urgent CD for NRHFT after an ECV is about 1/600 [6]. Placental abruption (<1%) and onset of labor are uncommon complications. Rare fetal deaths following attempts at version have not been determined to be a result of the procedure [1]. Femur fracture has been reported. In a meta-analysis, there was a risk of 4.7% for transient abnormal cardiocotography, 0.21% risk of abnormal cardiocotography leading to emergency CD (with good neonatal outcomes), and 0.35% risk of emergency CD. Other risks included 0.24% risk of stillbirth, 0.18% risk of placental abruption, 0.18% risk of cord prolapse, and 0.19% risk of fetal death. These complications were not found to be directly related to the ECV procedure. Vaginal bleeding related to ECV occurred in 0.34% of patients and rupture of membranes related to ECV occurred in 0.22% of patients [7].

Contraindications
Any contraindications to vaginal delivery such as placenta previa or prior classical uterine incision are considered contraindications to ECV. There are no trials on ECV in multiple gestations, so the safety and efficacy of this procedure cannot be assessed in this population. Relative contraindications are rupture of membranes, oligohydramnios, known uterine or fetal anomaly, unexplained uterine bleeding, or active phase of labor [1,8]. ECV in women with prior cesarean deliveries are associated with comparable success rates to those of women without prior cesarean deliveries, but there is insufficient data to assess the safety of this management [9].

Efficacy
Compared with no ECV, ECV at term is associated with a statistically significant and clinically meaningful 58% reduction in noncephalic birth (relative risk [RR] 0.42, 95% confidence interval [CI] 0.29–0.61) and a 42% decrease in CD (RR 0.58, 95% CI 0.40–0.82) [10]. There are no significant differences in Apgar score ratings <7 at 5 minutes (RR 0.63, 95% CI 0.29–1.36), low umbilical artery pH levels (RR 0.65, 95% CI 0.17–2.44), neonatal admission (RR 0.80, 95% CI 0.48–1.34), perinatal death (RR 0.39, 95% CI 0.09–1.64), or time from enrollment to delivery, compared with no ECV [10]. Because ECV is associated with a very low incidence of adverse events and with a significant decrease in CD, all women at or near term with nonvertex presentations should be offered an ECV.

Success rates of ECV average 58%, with a range of 25% to 80% [1]. Success is increased with higher parity and with transverse or oblique lie (vs. breech). Lower amniotic fluid volume, anterior placenta, and high maternal BMI might decrease the success rate [1]. A meta-analysis showed that multiparity, non-engagement of the breech, a relaxed uterus, a palpable fetal head, and maternal weight less than 65 kg were predictors of successful ECV [11]. There is no scoring system to accurately predict the probability of success of ECV. After successful ECV, the chance of spontaneous version to breech is low, but understudied. The chance of spontaneous version after failed ECV is about 6.6% in one study [12].

Timing of Version
Compared with no ECV, ECV before term reduces noncephalic births [13–16].
Compared with ECV at term, ECV before term reduces noncephalic presentation at birth, but does not reduce the rate of CD, and may be associated with an increase in the incidence of PTB. There is insufficient evidence to assess the best GA at which to perform ECV. In general, the later the GA, the lower the success rate, but there are reports of successful ECV in women in term labor. About 36 weeks is generally considered to be the optimal time for attempted version. At 36 weeks, there is felt to be adequate room to turn the fetus while minimizing the risk of return to breech presentation following a successful ECV. Additionally, if delivery becomes necessary, a 36-week infant has a low rate of respiratory distress syndrome or other complications of prematurity compared with a fetus of <36 weeks GA. Compared with ECV at 37 0/7–38 0/7 weeks, ECV at 34 0/7–36 0/7 weeks is associated with nonsignificant trends for slightly lower (57% vs. 66%) noncephalic presentation at birth and slightly lower (65% vs. 72%) CD [13–16]. In the largest randomized controlled trial (RCT) to date, compared with ECV at >37 weeks, ECV at 34 0/7–35 6/7 weeks was associated with a decrease in the incidence of noncephalic presentation at birth (41% vs. 49%; RR 0.84, 95% CI 0.75–0.94), but no difference in rates of CD (52% vs. 56%; RR 0.93, 95% CI 0.85–1.02) or risk of PTB (6.5% vs. 4.4%; RR 1.48, 95% CI 0.97–2.26) [17].

**Tocolysis**

Tocolysis with betamimetics prior to attempting ECV is associated with 30% fewer ECV failures, a 32% reduction in noncephalic presentations at birth, and 23% less CD [18]. A common tocolytic used is terbutaline with a dose of 25 mg subcutaneously once, 10–15 minutes before ECV. Other different betamimetics have been used with no evidence as to the best one or its dosage/timing [18]. One RCT of adjusted doses of intravenous salbutamol tocolysis prior to ECV increased success rates, decreased CD rate, and was well tolerated [19]. Tocolysis can also be used with success in a second ECV attempt after a first ECV attempt has failed [20].

**Nifedipine** as a uterine relaxant for ECV has also been evaluated. In a randomized, double-blind, placebo-controlled trial of 320 participants, nifedipine did not significantly improve the success of ECV [21]. In two RCTs evaluating oral nifedipine versus subcutaneous terbutaline tocolysis for ECV there was higher ECV success with terbutaline (52.2%–58.1% for terbutaline vs. 34.1%–39.5% for nifedipine), less CDs and no difference in neonatal outcomes, although more side effects were noted (maternal palpitations and tachycardia) [22–24].
Nitroglycerin has been studied as an agent to improve ECV success rates. In four small trials, sublingual nitroglycerin was associated with significant side effects and was not found to be effective [18]. One RCT demonstrated that treatment with intravenous nitroglycerin increased the rate of successful ECV in nulliparous women (24% compared with 8%, p = .04) but not in multiparous women [25]. In one trial terbutaline 25 mg subcutaneous 5 minutes prior to ECV was found to have a significantly higher ECV success rate than nitroglycerin [26].

**Fetal Acoustic Stimulation**
Fetal acoustic stimulation to the fetal head for 1–3 seconds in midline fetal spine positions is associated with fewer failures of ECV at term in a very small study [18,27]. Eleven of 12 ECVs were successful following stimulation [27]. The crossover arm (patients who failed version without stimulation) were then stimulated and 8 of 10 patients were successfully verted, for a total of 19 of 22 successful versions (86%) [26]. The success rate in the control group in this study was lower than expected (8%) [26]. The evidence is limited and is insufficient to make a recommendation.

**Anesthesia**
There is evidence that regional anesthesia affects ECV success. ECV failure, non-cephalic births, and cesarean deliveries were reduced in two trials with epidural but not in three trials with spinal anesthesia [18]. ECV success rates increased from 33% to 59% with epidural in one study and from 32% to 69% in another [4,28]. Potential bias in both studies lies in that the care provider was not blinded to placement of epidural [4,28]. It is important to note that the control groups had lower success rates than expected, as the average success rate in the literature, which is mostly without anesthesia, is about 58%. All patients in both studies received terbutaline prior to attempt at ECV. It has been postulated that large volume preloading with epidural may increase the amniotic fluid volume [18]. The use of spinal anesthesia has not been associated with any benefit in the success of ECV in some RCTs [8,18]. However, a meta-analysis including seven RCTs, of which five used spinal and two used epidural anesthesia for ECV versus controls, demonstrated anesthetic dose neuraxial blockade (usually spinal) is associated with a 44% increase in the success rate of external fetal version [29]. There are no trials to evaluate the potential effects of hydration or transabdominal amnioinfusion on the success rate of spontaneous version or ECV.

**Systemic Opioids**
There is limited evidence that systemic opioids improve success rates for ECV. One RCT of 60 women showed that the frequency of CD was similar in the opioid (remifentanil) group versus placebo (RR 0.9, 95% CI 0.20–4.27) [18]. In one RCT comparing spinal anesthesia, systemic opioid (remifentanil) and control (no anesthesia/analgésis) for ECV success, ECV was most successful in the spinal anesthesia group (83%) versus opioid (64%) and control (64%). Pain relief was highest with spinal anesthesia, followed by opioid and then control. Incidence of CD for fetal bradycardia was similar amongst groups [30].

**Hypnosis**
There is one RCT on hypnosis versus neurolinguistic programming for pain relief during ECV and both groups reported a similar degree of relief [18].

**ECV Procedure**
Given the possible complications, it is prudent to perform ECV in a facility with ready availability of emergency CD. Consent should be obtained after counseling regarding possible complications, alternatives (CD), prognosis and explanation of the actual procedure. A nonstress test should be performed before and after ECV. Anesthesia is usually not necessary and has not been absolutely proven to benefit outcomes. Betamimetic prophylactic tocolysis should be given (e.g., terbutaline 25 µg subcutaneously 5–10 minutes prior to procedure). There are no trials comparing other technical aspects of ECV. One or two operators can be used. Frequent if not continuous ultrasound guidance to assess for fetal well-being and presentation is suggested. Rh-negative women should receive anti-D immunoglobulin. There is no evidence to support immediate induction after successful ECV.

**Moxibustion and/or Acupuncture**
Moxibustion is a form of traditional Chinese medicine that uses heat generated by burning herbs, most often Artemisia vulgaris, to stimulate the acupuncture point BL67 (Zhiyin in Chinese) [31–35]. There is inconsistent evidence to assess whether the use of moxibustion converts a breech to a cephalic presentation. Differences in interventions (e.g., moxibustion alone or with acupuncture) make it difficult to perform a satisfactory meta-analysis. Moxibustion may reduce the need for ECV by 53% and reduce the incidence of nonvertex presentation at term by 35%–70%, in Chinese trials [32,33]. In two trials performed in Italy, moxibustion was not well tolerated by 22% of women and therefore not effective [34]; or effective when used with acupuncture [35]. Moxibustion may decrease the use of oxytocin before or during labor for women who had vaginal deliveries and might reduce non-cephalic presentation at birth and CD compared with acupuncture [31]. A meta-analysis performed compiling data from six studies of both Western and Chinese databases shows that moxibustion at point BL67, alone or in combination with acupuncture, is associated with higher rates of cephalic version of 72.5%, compared with 53.2% in the control group (RR 1.36, 95% CI 1.17–1.58). This data should be viewed with caution given the high degree of heterogeneity of the studies [36]. Also, there were no significant differences found in safety of moxibustion compared with other techniques. A recent RCT of 328 women showed no beneficial effect of moxibustion to facilitate ECV compared with controls when looking at the percent of fetuses in breech presentation at 37 weeks (72.0% vs. 63.4%; RR 1.13, 95% CI 0.98–1.32 [37]. It might be that acupuncture and not moxibustion (especially not at home) is beneficial [35].

**Maternal Change in Posture**
Maternal changes in position such as knee–chest and others has been suggested as a means to correct breech presentation in pregnancy. There is insufficient evidence from the small trials reported so far to support the use of postural management for breech presentation [38]. Meta-analysis could not be undertaken since study designs and outcomes measured were different [39]. Postural management is not associated with a significant effect on the rate of noncephalic births, either for the subgroup in which no ECV was attempted, or for the group overall (RR 0.98, 95% CI 0.84–1.15). No differences were detected for cesarean deliveries (RR 1.10, 95% CI 0.89–1.37). To date there is no solid evidence for this practice [38,39].

**Delivery Outcomes**
It is important to note that the rate of CD after ECV is still about double that of pregnancies presenting with spontaneous cephalic presentation due to higher incidences of dystocia and NRFHT after successful ECV [40].
MODE OF DELIVERY

Singleton

**Term Breech**

Three RCTs [41], including one large study (the Term Breech Trial) [42], have compared a policy of planned CD to a policy of planned trial of labor to attempt a vaginal delivery. CD occurs in about 45% of women allocated to a vaginal delivery protocol and >90% in those allocated to a CD protocol.

At 4–6 weeks after delivery, compared with planned vaginal delivery, planned CD is associated with a 67% decrease in perinatal or neonatal death (RR 0.33, 95% CI 0.19–0.56) (excluding fetal anomalies) or serious neonatal morbidity [41]. This reduction is less for countries with high national perinatal mortality rates [42]. Planned CD is associated with a 71% reduction (from 1.15% to 0.26%) in perinatal or neonatal death (excluding fetal anomalies) [41]. This reduction was similar for countries with low and high national perinatal mortality rates. One death could be prevented for every 112 CDs planned [41]. A secondary analysis [43] of the short-term outcomes of the Term Breech Trial [42] looked at factors associated with adverse outcomes. The lowest morbidity was found in patients with planned CD prior to the onset of labor. In a planned vaginal breech delivery labor augmentation and a second stage greater than 60 minutes are associated with less optimal outcomes [43]. Factors not shown to affect outcome included induction, parity, use of continuous electronic fetal monitoring, or epidural [43]. A “skilled clinician” at the delivery was associated with lower adverse outcomes. “Skilled clinician” was best described by the clinicians themselves rather than by years of experience or being a licensed obstetrician [43].

Three months after delivery, women allocated to the planned CD group reported 38% less urinary incontinence; 89% less abdominal pain; and 68% less perineal pain [44].

Two years after delivery, there was no difference in the combined outcome “death or neurodevelopmental delay.” Of 463 vaginal delivery patients followed at 2 years, there were only six deaths and seven children with neurodevelopmental delay (2.8%), compared with 2 and 12 of 457 patients (3.1%) in the CD group [45]. The authors postulate that there was no difference seen at the 2-year follow-up (vs. immediate neonatal outcome) because the study was underpowered, the predictive value was low for an association of measures of early morbidity with later death and adverse neurodevelopmental outcomes, and because planned CD is perhaps only reducing the risk of perinatal mortality/morbidity associated with fetal hypoxemia [45]. Maternal outcomes at 2 years were also very similar, with constipation significantly more common in the CD group (27% vs. 20%), while self-reported incontinence was nonsignificantly different (18% vs. 22%) [46]. Incontinence was different (16% vs. 25%) if comparing women who actually planned and had a CD versus those who planned and had a vaginal delivery [46].

These results are mostly from the Term Breech Trial [42] and its secondary analyses and follow-up [42–45]. These outcomes are based on deliveries done by “clinicians who were regarded as experienced at vaginal breech delivery” [42–46]. As the number of vaginal breech deliveries decreases, physician skill will continue to diminish, with the potential to make vaginal delivery less safe. While it is estimated that >90% of babies presenting nonvertex are currently delivered by CD, there might still be a small role for vaginal delivery for the woman who declines scheduled CD or who presents in advanced labor. The American College of Obstetricians and Gynecologists (ACOG) in 2014 supported the decision that the mode of delivery depends on the experience of the health care provider [47]. CD is the preferred mode due to the limited experience of most physicians, but planned vaginal delivery of a term singleton fetus may be considered under specific hospital protocols. All women with breech presentation with a large fetus (>3500 g estimation), unfavorable pelvis, hyperextended head, incomplete or footling breech presentation, NRFHT, severe fetal growth restriction, or lack of experienced obstetrical and anesthesiological operators should have a CD.

**Technical aspects**

**Cesarean breech delivery:** There are no trials to assess technical aspects of breech (or other malpresentation) CD. There is insufficient evidence to assess whether intra-abdominal version during CD before uterine incision affects outcomes.

**Vaginal breech delivery:** There are several technical suggestions for assisting a vaginal breech delivery; none are based on trials. There is insufficient evidence to assess whether clinical/radiologic pelvimetry affects outcomes in the management of breech presentation. A double setup is suggested: a vaginal delivery should be organized in the operating room and ready for possible CD.

Some other suggestions are as follows: minimal intervention until the abdomen up to the umbilical cord, is delivered; prevention of head extension, with prophylactic Mauriceau maneuver and proper use of Pipers forceps, if necessary. There is not enough evidence to evaluate the effects of expedited vaginal breech delivery (breech delivery from umbilicus to delivery of the head within one contraction) on perinatal outcomes [48].

The same management options exist for transverse/oblique lie as for breech. Fetal ECV, and CD if persistent malpresentation, are the standards of care, with limited trial evidence.

**Preterm Breech**

There is insufficient evidence to assess whether outcomes of the preterm breech presenting fetus are affected by mode of delivery. Very little prospective data, mostly nonrandomized, exists regarding vaginal versus CD of the premature breech infant [1]. Two trials aimed to assess this question failed to randomize the planned sample sizes. After 17 months of patient recruiting at 26 different hospitals, only 13 women had been randomized making it impossible to confer any conclusions in one study [49].

The Iowa premature breech trial was somewhat more successful, recruiting 38 patients over 5 years, but had insufficient data for meaningful conclusions [50]. Outcomes in premature breech infants are mainly related to prematurity and/or fetal anomaly, with unclear effect of mode of delivery [51].

**Twins**

**Breech Second Twin**

(See also Chapter 44 in *Maternal-Fetal Evidence Based Guidelines*)

Pregnancies at ≥35 weeks with vertex/breech presentation in twin gestations <7 cm dilation have similar Apgar scores and incidence of neonatal morbidity in the second twin delivered by vaginal or cesarean birth [52]. There was no incidence of birth trauma or intraventricular hemorrhage in any of the 27 breech deliveries of the second twin [52]. Maternal febrile morbidity and length of stay was increased in the CD group [52]. In the largest RCT that randomized women with twin pregnancies to planned CD or planned vaginal delivery, there was a large number of women with non-cephalic second twin. There was no statistically significant difference seen in composite fetal or neonatal morbidity, even when evaluating for interaction with fetal presentation, comparing perinatal morbidity in planned CD versus
planned vaginal delivery (odds ratio [OR] 1.16, 95% CI 0.77–1.74). Morbidity included birth trauma, seizures and assisted ventilation. There were also no significant differences noted in serious maternal morbidity (OR 0.86, 95% CI 0.65–1.13) [53]. **Vaginal delivery of the second nonvertex twin is a reasonable management option.** Attempt at vaginal twin delivery of the second twin, especially for a second twin with an estimated fetal weight (EFW) of >1500 g, should be performed with adequate experience of the obstetrician, as well as with continuous availability of expert anesthesia, in or very close to, an operating room. **Total breech extraction of the second twin** is associated with shorter maternal stay, lower neonatal pulmonary disease, infection, and intensive care nursery stay compared with cephalic version [49,54]. There are no trials for twins presenting with first twin nonvertex (about 26% of twins). Recommendation for CD under this circumstance is based mostly on data from singleton gestations. Because vaginal delivery of triplets is usually associated with an increased risk for stillbirth or neonatal and infant deaths, as compared with CD, **CD is the route of choice**, even if some small series have recently reported similar outcomes for trial of labor or CD for triplets.

**Preterm Twins**

There are no RCTs of planned CD versus vaginal delivery of preterm twins less than 34 weeks with malpresentation of the second twin. There is insufficient data to make a recommendation.

**MALPOSITION**

**Definitions**

**Position:** Relationship of presenting part (usually occiput for head) to pelvic outlet.

**Malposition:** Fetal position that is not occiput-anterior (or sacrum anterior).

Sincipital of the fetal head is the specific malposition in which neither of the parietal bones precedes the sagittal suture into the maternal birth canal. The situation when one of the parietal bone precedes the sagittal suture is called *asynclitism* [55].

**Epidemiology/Incidence**

Occiput posterior (OP) is the most common fetal malposition. Only 5% of term fetuses are OP at time of delivery, but about 23% were OP at the beginning of labor [56].

**Risk Factors/Associations**

OP position is associated with African American race, advanced maternal age, post term pregnancy, birth weight >4000 g, and epidural anesthesia [57]. In a meta-analysis evaluating epidural versus no-epidural/analgesia in labor, malposition was higher in women in the epidural group (RR 1.40, 95% CI 0.98–1.99), although this was not statistically significant [58].

**Diagnosis**

Fetal malposition can be detected by provider digital examination as well as by transabdominal ultrasound. One prospective RCT found a difference in digital vaginal examination and transabdominal ultrasonographic examination of fetal head position during the second stage of labor in 20% of the cases evaluated, promoting the use of transabdominal ultrasound to locate the position of the fetal head [59].

**Management**

**Hands and Knees Posture**

There is insufficient evidence for assessing the effect of hands and knees posture to correct malposition. This has been evaluated both before (for prevention) and during labor.

There are two trials on the effect of hands and knees posture before labor [60]. Compared with a sitting position, 10 minutes in the hands and knees position is associated with a lower likelihood of malposition of the presenting part of the fetus in a small trial, but assuming the hands and knees posture for 10 minutes twice daily in the last weeks of pregnancy had no effect on the baby’s position at delivery or any of the other pregnancy outcomes measured in a larger trial [60].

One study evaluated the use of hands and knees position in labor involving 147 women in labor at term who assumed the hands and knees position for a period of at least 30 minutes. There was no significant reduction in occiput-posterior or occiput-transverse positions at delivery or reduction of operative deliveries. However, there was a significant reduction in back pain [61].

**Manual Rotation**

Persistent fetal OP position is a risk factor for prolonged labor and higher rate of CD. There is insufficient evidence evaluating the use of manual (also called digital) rotation in reducing the prevalence of persistent occiput-posterior position and its consequences. In a retrospective cohort study, in women with a fetus in persistent occiput posterior or transverse position in the second stage of labor, manual rotation was associated with lower rates of CD, perineal lacerations, postpartum hemorrhage, and chorioamnionitis, but a higher rate of cervical lacerations [62]. A prospective (but not randomized) study of singletons with occiput-posterior position reported an increase in fetuses delivered in occiput-anterior position (93% vs. 15%) and in spontaneous vaginal delivery (77% vs. 27%) compared with no manual rotation using historic controls [63] (see Chapter 8).

**Operative Delivery from Malposition (OP)**

There is insufficient evidence to make a recommendation.

**Ultrasound**

In a recent RCT that evaluated the influence of ultrasound on diagnosis of fetal head malposition and subsequent mode of delivery, using transabdominal ultrasound in active labor did not improve management of labor, increased the incidence of operative delivery (CD + operative vaginal delivery) (RR 1.24, 95% CI 1.08–1.43) and did not decrease maternal and neonatal morbidity [64].

**ANESTHESIA**

For vaginal breech delivery, an anesthesiologist skilled at the pharmacology of uterine relaxation (e.g., nitric oxide) should be present.

**REFERENCES**

18. Hofmeyr GJ. Interventions to help hard to turn breech babies to head first presentation when using external cephalic version. Cochrane Database Syst Rev. 2015;(3):CD00184. [17 RCTs c tocolysis, n = 1876; 1 acoustic stimulation, n = 114]


Shoulder dystocia

Sean C. Blackwell

KEY POINTS

- Risk factors for shoulder dystocia include prior shoulder dystocia (recurrence risk ~1%–15%); maternal diabetes mellitus, obesity, postterm, labor induction, epidural anesthesia, labor abnormalities (e.g., prolonged second stage), operative vaginal delivery, and fetal macrosomia (Table 25.1).
- In 50% of shoulder dystocia cases, no risk factors are identified. Therefore, clinicians should be ready for possible shoulder dystocia at every delivery.
- In 40%–60% of shoulder dystocia cases, birth weight is <4000 g.
- Maternal complications of shoulder dystocia include serious vaginal laceration (third or fourth degree) and postpartum hemorrhage.
- A perinatal complication of shoulder dystocia is brachial plexus impairment (BPI), which occurs in 4%–40% of shoulder dystocia cases; over 90% are transient. Approximately one-half of cases of BPI occur without documented shoulder dystocia.
- Other perinatal complications of shoulder dystocia include fractures, hypoxic–ischemic encephalopathy, long-term neurologic disability, and even death.
- Preconception prevention of maternal obesity and diabetes, as well as prenatal prevention of excessive weight gain, decrease the incidence of shoulder dystocia, mainly by prevention of fetal macrosomia.
- In women with prior shoulder dystocia, clinicians should review recurrence risk (7%–15%), which risk factors (Table 25.1) are present in the current pregnancy, and also possible complications. If several significant risk factors are still present, the woman may opt for cesarean delivery. If risk factors are not present (except for prior shoulder dystocia), the woman may decide after counseling for either trial of labor or cesarean delivery.
- Induction of labor at term for suspected fetal macrosomia (estimated fetal weights [EFW] ≥4000 g) in women without diabetes is associated with a similar incidence of cesarean or operative vaginal delivery compared with expectant management, a significant decrease in fetal fractures, and higher incidences of hyperbilirubinemia and phototherapy. Induction at ≥38 weeks would prevent the neonatal complications, and can be considered.
- In pregnancies with EFW > 5000 g in nondiabetic and >4500 g in diabetic women, The American Congress of Obstetricians and Gynecologists (ACOG) suggests planned cesarean delivery (without labor attempt) may be considered to avoid shoulder dystocia.
- In women with EFW ≥3800 g, prophylactic McRobert’s maneuver and suprapubic pressure are associated in one randomized controlled trial (RCT) with significantly fewer instances of shoulder dystocia.
- Initial responses to shoulder dystocia involve asking for help, and, as first maneuvers, use of McRobert’s maneuver and suprapubic pressure (Figure 25.1).
- Second line maneuvers include direct fetal manipulation such as delivery of the posterior arm, or then rotation of fetal shoulders. One-third of shoulder dystocia cases require use of more than two maneuvers. See Figure 25.1 for suggested management of shoulder dystocia.

DIAGNOSIS/DEFINITION

Shoulder dystocia is diagnosed when additional obstetrical maneuvers are required following failure of gentle downward traction on the fetal head to effect delivery of the shoulders [1]. Shoulder dystocia pertains only to vertex presentation.

A quantitative definition of shoulder dystocia has been proposed as a prolonged head-to-body delivery time >60 seconds [2]. This definition categorized 10% of vaginal deliveries as having shoulder dystocia while only 25%–45% of these cases were diagnosed as such by the delivery provider [3]. However, these diagnostic criteria have not been adopted and shoulder dystocia remains a subjective clinical diagnosis by the delivering provider. This may account for some of the variability in the frequency of reported shoulder dystocia across various populations and different studies.

SIGNS

1. Retraction of the delivered fetal head against the maternal perineum (turtle sign).
2. Inability to deliver the fetal shoulders with routine traction in the “axial” direction.

EPIDEMIOLOGY/INCIDENCE

The incidence of shoulder dystocia is about 1% (range 0.6%–1.4%, depending on definition used).

ETIOLOGY/BASIC PATHOPHYSIOLOGY

Shoulder dystocia results from size discrepancy between the fetal shoulders and the pelvic inlet that results in impaction of the anterior shoulders behind the maternal pubis symphysis or the posterior shoulder on the sacral promontory (uncommon), or both (rare) [4]. Persistent anterior–posterior location of the fetal shoulders at the pelvic brim occurs most often due to: increased resistance between the fetal skin and vaginal walls (e.g., fetal macrosomia), a large fetal chest relative to the biparietal diameter (e.g., infants of diabetic mothers), and when truncal rotation does not occur (e.g., precipitous labor).
Table 25.1  Factors Associated with Shoulder Dystocia

<table>
<thead>
<tr>
<th>Antepartum</th>
<th>Intrapartum</th>
</tr>
</thead>
<tbody>
<tr>
<td>Multiparity</td>
<td>Labor induction and/or augmentation</td>
</tr>
<tr>
<td>Postterm gestation</td>
<td>Labor abnormalities</td>
</tr>
<tr>
<td>Maternal obesity</td>
<td>Prolonged first stage</td>
</tr>
<tr>
<td>Maternal diabetes</td>
<td>Short second stage</td>
</tr>
<tr>
<td>Prior shoulder dystocia</td>
<td>Prolonged second stage</td>
</tr>
<tr>
<td>Prior macrosomic child</td>
<td>Epidural</td>
</tr>
<tr>
<td>Excessive gestational weight gain</td>
<td>Operative vaginal delivery (forceps or vacuum)</td>
</tr>
<tr>
<td>Fetal macrosomia</td>
<td></td>
</tr>
</tbody>
</table>

Source: American College of Obstetricians and Gynecologists, Obstet Gynecol, 100(5 Pt 1), 1045–1050, 2002.

Table 25.2  Relationship between Increasing Birth Weight and Maternal Diabetes and the Development of Shoulder Dystocia

<table>
<thead>
<tr>
<th>Birth Weight (g)</th>
<th>No Diabetes (%)</th>
<th>Maternal Diabetes (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>4000–4250</td>
<td>5.25</td>
<td>12.2</td>
</tr>
<tr>
<td>4250–4500</td>
<td>9.1</td>
<td>16.7</td>
</tr>
<tr>
<td>4500–4750</td>
<td>14.3</td>
<td>27.3</td>
</tr>
<tr>
<td>4750–5000</td>
<td>21.1</td>
<td>34.8</td>
</tr>
</tbody>
</table>


Figure 25.1  Suggested management algorithm for shoulder dystocia.

RISK FACTORS/ASSOCIATIONS

Table 25.1 lists antepartum and intrapartum risk factors associated with shoulder dystocia. These risk factors have poor predictive value (whether in isolation or in combination with other factors), making them not very useful for clinical decision making. In 50% of cases of shoulder dystocia, no risk factors are identified.

The two risks factors for shoulder dystocia most studied are maternal diabetes and fetal macrosomia. The frequency of shoulder dystocia increases with higher birth weight [5] (Table 25.2).

Although there is a consistent relationship regarding birth weight and the risk for shoulder dystocia, it should also be remembered that 40%–60% of cases occur with birth weight <4000 g, and of deliveries with birth weight >4000 g only 3.3% develop shoulder dystocia [4,6,7]. Although recognized as a risk factor associated with shoulder dystocia, multiple studies including those using advanced statistical models and techniques have been unable to discriminate labor patterns (e.g., prolonged first and/or second stage) in a manner that improves intrapartum management to predict and avoid shoulder dystocia [8–12]. Dyachenko et al. studied 498 cases of shoulder dystocia (including 90 cases with neonatal injury) versus 622 controls with vaginal delivery assessing for whether a combination of maternal and fetal factors could reasonably predict outcomes. The statistical model that best predicted shoulder dystocia with injury included maternal height and weight, gestational age, parity, and birth weight identified 50.7% of cases with a false positive rate of 2.7%.

COMPLICATIONS

Maternal
Although maternal complications can occur, these are most often mild and not associated with long-term morbidities. The most common complications involve postpartum blood loss and vaginal laceration (Table 25.3).

Perinatal
The most common major complication of shoulder dystocia is neonatal BPI which occurs in 4%–40% of cases of shoulder dystocia. In one of the largest series, the rate was 16.8% (Table 25.3) [13]. Most cases of BPI are unilateral (right > left) and involve the C5–C6 roots (Erb–Duchenne’s palsy) (Table 25.4). Most bony injuries (clavicle or humerus), although always concerning, are not serious and heal promptly without long-term sequelae. Approximately 1/3 BPIs have a concomitant bony fracture. Fortunately <10% of BPIs are permanent. Most fully recover with physical therapy and, in some situations, respond to neurosurgical treatment. Permanent BPI is very rare and occurs in 1–2 out of every 10,000 births. Signs and symptoms are a limp or unmoving paralyzed arm, lack of muscle control in the arm, hand, or wrist, and lack of feeling or sensation in the arm or hand. Neonatal BPIs can occur without shoulder dystocia. In fact, it is estimated that one half of all BPIs occur in the absence of shoulder dystocia and up to 4% of BPIs are associated with cesarean delivery [14,15]. Two studies provide evidence that BPI (both transient and permanent) can occur with cesarean delivery. Gherman et al. [6,16] reported the finding of 17 cases of BPI (six with permanent) associated with cesarean delivery with longer term follow-up (12–29 months). Chang et al. described 30 cases of BPI with cesarean delivery (out of total 387 cases) where 60% were persistent at 1 year [17]. In a recent
and 47% outside the United States were not associated with dystocia, approximately 45% of NPP cases in the United States literature review of neonatal BPP with and without shoulder dystocia, mean umbilical pH was not clinically different than cases without shoulder dystocia and there was no association between head-to-body delivery intervals and umbilical pH, base excess, or 5-minute Apgar scores [22]. However, another study done by Leung et al. of 200 cases of shoulder dystocia noted that the umbilical artery pH dropped 0.01 per minute for the head-to-body delivery interval (HBDI). They found an overall 2.5% rate of hypoxic ischemic encephalopathy (the risk was 0.5% if HBDI was <5 minutes compared with 23.5% if HBDI ≥5 minutes) [23].

In a review of 56 cases of fatal shoulder dystocia reported to the Confidential Inquiry into Stillbirths and Deaths in Infancy between 1994 and 1995 from England, Northern Wales, and Ireland the approximate incidence of fatal shoulder dystocia is 0.025 per 1000 deliveries. In 47% of these cases, the head-to-body ratio was less than 5 minutes and greater than 10 minutes in 20% [24].

### MANAGEMENT

#### Principles

Shoulder dystocia is an obstetrical emergency and cannot be reliably predicted. Thus prompt recognition and response are essential to optimizing outcomes once it develops. Obstetricians should be trained and ready to manage this complication at every delivery. Therefore, it is imperative that shoulder dystocia training occurs early, and that at least one clinician experienced in the management of shoulder dystocia is present at every delivery.

#### Prevention

**Preconception**

Preconception prevention of maternal obesity and diabetes, as well as prenatal prevention of excessive weight gain, decrease the incidence of shoulder dystocia, mainly by prevention of fetal macrosomia.

**Antepartum**

Since clinical risk factors (Table 25.1) do not accurately predict who will develop shoulder dystocia, few preventive strategies have been proven effective. Nonetheless, avoidance, prevention or proper control of maternal obesity, diabetes, excessive weight gain, and fetal macrosomia would prevent the increased risk of shoulder dystocia associated with these conditions. Women with prior shoulder dystocia have about a 1%–15% [25,26] risk of recurrence. In women with a prior shoulder dystocia, counseling should include review of prior delivery events including severity of injury of prior child, which risk factors (Table 25.1) are present in the current pregnancy, and also possible short-term and long-term complications of cesarean delivery. If several significant risk factors are still present, the woman may opt for cesarean delivery. However, given at least 85% of women will not have a recurrent shoulder dystocia and cesarean delivery does have certain surgical risks (especially in setting of maternal comorbidities) a woman with a prior shoulder dystocia after counseling may choose to pursue vaginal delivery [1].

Strategies to perform cesarean delivery without a labor attempt, due solely to suspected fetal macrosomia, are neither cost-effective nor practical due to poor prediction abilities [27,28]. This is complicated by the known limitations of prenatal ultrasound to accurately estimate fetal weight, especially with increased fetal size [29,30]. Only at very high estimated fetal weights (>5000 g in nondiabetic and >4500 g in diabetic women) does ACOG suggests planned cesarean delivery (without labor attempt) may be considered to avoid shoulder dystocia [29].

**Labor induction** for women with suspected macrosomia as an intervention to decrease the rate of shoulder dystocia has been tested in four clinical trials, including 1190 women [31–34,35] Women who were randomized to labor induction had a similar incidence of cesarean delivery (26.6% vs. 29.4%);

### Table 25.3 Maternal and Newborn Complications of Shoulder Dystocia Reported at Single Large Tertiary Care Center

<table>
<thead>
<tr>
<th>Complication</th>
<th>Frequency (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Postpartum hemorrhage</td>
<td>11</td>
</tr>
<tr>
<td>Fourth degree laceration (either extension of episiotomy or extension of laceration)</td>
<td>3.8</td>
</tr>
<tr>
<td>Vaginal lacerations</td>
<td>19.3</td>
</tr>
<tr>
<td>Cervical tear</td>
<td>2</td>
</tr>
</tbody>
</table>

#### Perinatal

<table>
<thead>
<tr>
<th>Complication</th>
<th>Frequency (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Any fetal injury</td>
<td>24.9</td>
</tr>
<tr>
<td>Brachial plexus injury</td>
<td>16.8</td>
</tr>
<tr>
<td>Clavicular fracture</td>
<td>9.5</td>
</tr>
<tr>
<td>Humeral fracture</td>
<td>4.2</td>
</tr>
</tbody>
</table>

### Table 25.4 Types of Brachial Plexus Impairments

<table>
<thead>
<tr>
<th>Palsy</th>
<th>Description</th>
<th>Level</th>
<th>Frequency (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Erb’s palsy (also known as Erb–Duchenne’s palsy)</td>
<td>Paralysis of flexion, abduction, internal and external rotation of the forearm</td>
<td>C5, C6</td>
<td>&gt;98</td>
</tr>
<tr>
<td>Klumpke’s palsy</td>
<td>Paralysis of the thumb, fingers, and pronation of the forearm</td>
<td>C8, T1</td>
<td>&lt;1</td>
</tr>
<tr>
<td>Complete brachial plexus palsy</td>
<td>Both Erb’s and Klumpke’s</td>
<td>C5, T1</td>
<td>&lt;1</td>
</tr>
</tbody>
</table>


### Table 25.3 Maternal and Newborn Complications of Shoulder Dystocia Reported at Single Large Tertiary Care Center

<table>
<thead>
<tr>
<th>Complication</th>
<th>Frequency (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Postpartum hemorrhage</td>
<td>11</td>
</tr>
<tr>
<td>Fourth degree laceration (either extension of episiotomy or extension of laceration)</td>
<td>3.8</td>
</tr>
<tr>
<td>Vaginal lacerations</td>
<td>19.3</td>
</tr>
<tr>
<td>Cervical tear</td>
<td>2</td>
</tr>
</tbody>
</table>

relative risk [RR] 0.90, 95% confidence interval [CI] 0.75–1.09),
operative vaginal delivery (13.0% vs. 15.2%; RR 0.86, 95% CI
0.65–1.13), spontaneous vaginal delivery (60.3% vs. 55.4%; RR
1.09, 95% CI 0.99–1.20), shoulder dystocia (2.4% vs. 4.2%; RR
0.57, 95% CI 0.30–1.08), intracranial hemorrhage (0.6% vs. 0.4%;
RR 1.48, 95% CI 0.20–12.57), brachial plexus palsy (0.0% vs.
0.3%; RR 0.21, 95% CI 0.01–4.28), Apgar score < 7 at 5 minutes (0.7% vs.
0.5%; RR 1.51, 95% CI 0.25–9.02), cord blood pH < 7 (0.2% vs.
0.4%; RR 0.44, 95% CI 0.06–2.97), and mean birth weight (mean
difference [MD] –134.41 g, 95% CI –317.27 to 48.46) compared
with women expectantly managed. Induction group had a signifi-
cantly lower time to delivery (MD –7.55 days, 95% CI –8.20
to –6.89), birth weight ≥4000 g (30.7% vs. 61.8%; RR 0.50, 95% CI
0.42–0.59), birth weight ≥4500 g (3.2% vs. 14.8%; RR 0.21, 95% CI
0.11–0.39), incidence of fetal fractures (0.3% vs. 2.0%; RR 0.17,
95% CI 0.03–0.79) and a significantly higher incidence of hyper-
bilirubinemia (8.8% vs. 2.9%; RR 3.03, CI 1.60–5.74) and photo-
therapy (11.0% vs. 6.6%; RR 1.68, CI 1.07–2.66) compared with
operative vaginal delivery (13.0% vs. 15.2%; RR 0.86, 95% CI
0.90, 95% confidence interval [CI] 0.75–1.09), and mean birth weight
(95% CI 0.01–2.73) [31].

In summary, induction of labor at term for suspected
fetal macrosomia (EFW ≥4000 g) in women without diabetes
is associated with a similar incidence of cesarean or oper-
ative vaginal delivery compared with expectant management,
a significant decrease in fetal fractures, and higher inci-
dences of hyperbilirubinemia and phototherapy. Induction at
≥38 weeks would prevent the neonatal complications, and can
be considered.

Data regarding the effects on labor induction in women
with diabetes to reduce shoulder dystocia are limited to one
trial involving 200 women. There was no statistically signifi-
cant differences in the rate of shoulder dystocia between labor
induction and expectant management (0/100 vs. 3/100; RR 0.14
[95 CI 0.01–2.73]) [31].

Intrapartum

Prophylactic McRobert’s maneuver, prior to the development
of shoulder dystocia, has not been shown to decrease subse-
quent shoulder dystocia in women at low-risk for this com-
plexion [36]. In a small RCT, in 40 women, compared with
lithotomy position, the use of the McRobert’s maneuver was
associated with the same incidence (71% vs. 77%) of shoulder
dystocia in both prophylactic and lithotomy groups. The forced
used in traction of the fetal head during vaginal delivery was
the same in each group [36].

In another RCT, in 185 women likely to give birth to a large
baby (EFW >3800 g), compared with no prophylactic maneu-
vers, the McRobert’s maneuver and suprapubic pressure were
associated with a trend for a lower rate of shoulder dystocia
compared with controls (9% vs. 21%; RR 0.44, 95% CI 0.17–1.14).
There were significantly more cesarean sections in the pro-
phyllactic group, and when these were included in the results,
significantly fewer instances of shoulder dystocia were seen
in the prophylactic group (RR 0.33, 95% CI 0.12–0.86). In this
study, 13 (18%) women in the control group required therapeutic
maneuvers after delivery of the fetal head compared with 3 (5%)
in the treatment group (RR 0.31, 95% CI 0.09–1.02). One infant in
the control group had a brachial plexus injury (RR 0.44, 95% CI
0.02–10.61), and one infant had a 5-minute Apgar score less than
seven (RR 0.44, 95% CI 0.02–10.61) [37,38].

Training

Three nonrandomized studies have shown benefits of shoulder
dystocia training. In a single center “before and after” study
conducted in Jamaica, NY, between 2003 and 2006 (pre) to 2006
and 2009 (post), shoulder dystocia training decreased the rate of
BPI associated with shoulder dystocia from 30% (pre) to 10.67%
(post) even after adjusting for confounding factors [39]. Another
“before and after” study in the United Kingdom showed a reduc-
tion in neonatal injury at birth after shoulder dystocia: (9.3% vs.
2.3%; RR 0.25; 95% CI 0.11–0.57) [40]. Finally, in a university
hospital setting in Chicago, IL, the effects of a shoulder dystocia
protocol was studied in three time periods (6 months prior to
training, 6 months during training, and 6 months posttraining).
Over the three time periods complete and consistent document-
ation improved (14%, to 50%, to 92%; p < .001), decrease BPI at
delivery (10.1%, to 4%, to 2.6%; p = 0.03) and BPI at neonatal dis-
charge (7.6%, to 3%, to 1.3%; p = .04). Both the Joint Commission
in the United States and the Royal College of Obstetricians and
Gynecologists recommend such training.

Recognition

After diagnosis of shoulder dystocia, whether done via rec-
ognition of the “turtle sign” or lack of fetal progression with
gentle traction, the delivery provider should “call for help.”
This includes utilizing the assistance of other care givers
present in the delivery room as well as alerting other obstet-
rical, pediatric and/or anesthesia providers on the labor and
delivery unit.

Therapy

There are no interventional trials in human subjects that com-
pare the safety or effectiveness of various shoulder dystocia
therapeutic maneuvers [38]. Thus guidelines are based on
observational data and/or expert opinion. The most important
feature of the response to shoulder dystocia is that it should
be a coordinated and orderly application and progression of
obstetrical maneuvers (Figure 25.1) [1, 41]. Table 25.5 describes
maneuvers employed to relieve shoulder dystocia [4, 42].

Most authorities recommend McRobert’s as the initial
maneuver since it is easy to perform, is effective, and has a low
complication rate. McRobert’s alone is effective in 40%–90% of
cases. Suprapubic pressure is often done in direct conjuction by
nursing personnel. In a series of 134 cases, more than one third
of patients required more than two maneuvers to relieve the
shoulder dystocia [22]. Another series of 231 shoulder dystocia
cases, 57% of cases responded to McRobert’s and suprapubic
pressure alone and had a median duration of 29 seconds [43].

If McRobert’s combined with suprapubic pressure are
not successful, direct fetal manipulation is attempted next. In a
very large series, delivery of the posterior shoulder was asso-
ciated with the highest rate of delivery (84%) compared with
all other maneuvers (24%–72%) [44]. These authors have sug-
gested considering using delivery of the posterior arm if deliv-
ery is not achieved with McRobert’s maneuver and suprapubic
pressure. The need of additional maneuvers is associated with
higher rates of neonatal injury; 10% with three maneuvers, 16%
with four maneuvers, and 23% with five or more [44].

Since shoulder dystocia is a bony dystocia, routine
episiotomy is not advised. In a systematic literature review
including 14 studies of 9769 cases of shoulder dystocia, there
was no reported benefit of episiotomy to prevent or assist
with the management of shoulder dystocia [45]. In situations
where delivery of the posterior arm is being tried, episiotomy
or perineal episiotomy may give the delivery attendant addi-
tional room to grab the posterior fetal arm. Other helpful fetal
manipulation interventions are either the Wood’s corkscrew
(Figure 25.2) or Rubin’s maneuver (Figure 25.3).
Table 25.5 Description of Various Obstetrical Maneuvers to Resolve Shoulder Dystocia

<table>
<thead>
<tr>
<th>Maneuver</th>
<th>Maternal or Fetal Manipulation</th>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>McRobert's</td>
<td>Maternal First line</td>
<td>Hyperflexion and abduction of the maternal hips (knee to abdomen position). This causes cephalad rotation of the symphysis pubis, and flattening of the</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>lumbar lordosis, freeing the impacted shoulder. Pressure directed posteriorly in an attempt to force the anterior shoulder under the symphysis pubis. It also may include lateral pressure from either side of the maternal abdomen or alternating between sides using a rocking pressure.</td>
</tr>
<tr>
<td>Suprapubic pressure</td>
<td>Maternal First line</td>
<td></td>
<td>It works by reducing the fetal bisacromial diameter and rotating the anterior shoulder into oblique pelvic diameter.</td>
</tr>
<tr>
<td>Delivery of posterior arm</td>
<td>Fetal Second line</td>
<td></td>
<td>The delivery attendant places his/her hands in the vagina and applies pressure at the antecubital fossa in order to flex the fetal forearm. The arm is subsequently swept out over the infant's chest and delivered over the perineum. Rotation of the fetal trunk to bring the posterior arm anteriorly is sometimes required.</td>
</tr>
<tr>
<td>Rubin's</td>
<td>Fetal Second line</td>
<td></td>
<td>Fracture of the humerus can occur with grasping and pulling directly on the fetal arm, as well as application of pressure onto the midhumeral shaft.</td>
</tr>
<tr>
<td>Wood's</td>
<td>Fetal Second line</td>
<td></td>
<td>It works by rotating the fetal shoulder into the oblique pelvic diameter.</td>
</tr>
<tr>
<td>Gaskin (knee–chest or all fours)</td>
<td>Maternal Second line</td>
<td></td>
<td>Pressure is applied onto the anterior surface of the posterior shoulder in order to abduct the posterior shoulder.</td>
</tr>
<tr>
<td>Intentional fracture of clavicle</td>
<td>Fetal Third line</td>
<td></td>
<td>It works by rotating the fetal shoulder into the oblique pelvic diameter. Patient is rolled from her existing position onto her hands and knees.</td>
</tr>
<tr>
<td>Zavanelli (cephalad replacement)</td>
<td>Third line</td>
<td></td>
<td>It works through the downward force of gravity and/or a favorable change in pelvic diameters, which causes disimpaction of the fetal shoulder.</td>
</tr>
<tr>
<td>Symphysiotomy</td>
<td>Maternal Third line</td>
<td></td>
<td>Surgical or traumatic separation of the pubis symphysis to facilitate disimpaction of the fetal shoulders.</td>
</tr>
</tbody>
</table>

Figure 25.2 Wood's corkscrew maneuver. Initial pressure exerted on the anterior surface of the posterior shoulder facilitates rotation of the posterior shoulder anteriorly (upper). With concurrent synchronized downward pressure, the shoulder “screws” though the maternal pelvis, disimpacting the previously impacted shoulder (lower). (From Ramsey PS et al., J Reprod Med, 45, 85–88, 2000. With permission.)

Figure 25.3 Rubin's maneuver. (a) and (b) Pressure is exerted on the posterior surface of the most accessible part of the shoulder to facilitate abduction and disimpaction of the anterior shoulder. (c) Further rotation and adduction toward the fetal chesteduces the bisacromial diameter and results in the movement of the shoulders in a transverse position, facilitating passage of the anterior side of the shoulder beneath the pubic arch. (From Ramsey PS et al., J Reprod Med, 45, 85–88, 2000. With permission.)
Shoulder Dystocia Delivery Note Addendum

1. Antepartum documentation:
   a. Prior shoulder dystocia? Yes no
   b. Estimated fetal weight

2. Mode of vaginal delivery
   a. Spontaneous
   b. Vacuum
   c. Forceps

3. How was the shoulder dystocia diagnosed? Turtle sign
   Failure to deliver shoulders with gentle downward pressure
   Other _________________________________________________

4. Time fetal head delivered? _____________________

5. Time body delivered? _____________________

6. Total duration of shoulder dystocia (in minutes and seconds): _____________________

7. What shoulder was under the pubis symphysis (anterior) at delivery: Left Right Unknown

8. Please describe the obstetrical maneuvers attempted, their order, and who performed:

<table>
<thead>
<tr>
<th>Maneuver</th>
<th>Order</th>
<th>By whom</th>
</tr>
</thead>
<tbody>
<tr>
<td>McRobert’s (Hip flexion)</td>
<td>1 2 3 4 5 6 7 8</td>
<td></td>
</tr>
<tr>
<td>Suprapubic</td>
<td>1 2 3 4 5 6 7 8</td>
<td></td>
</tr>
<tr>
<td>Delivery of posterior arm</td>
<td>1 2 3 4 5 6 7 8</td>
<td></td>
</tr>
<tr>
<td>Rubin’s (Anterior scapula)</td>
<td>1 2 3 4 5 6 7 8</td>
<td></td>
</tr>
<tr>
<td>Wood’s (Posterior scapula)</td>
<td>1 2 3 4 5 6 7 8</td>
<td></td>
</tr>
<tr>
<td>Episiotomy</td>
<td>1 2 3 4 5 6 7 8</td>
<td></td>
</tr>
<tr>
<td>Extension of Episiotomy</td>
<td>1 2 3 4 5 6 7 8</td>
<td></td>
</tr>
<tr>
<td>Gaskin’s (Knee-chest, all fours)</td>
<td>1 2 3 4 5 6 7 8</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>1 2 3 4 5 6 7 8</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>1 2 3 4 5 6 7 8</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>1 2 3 4 5 6 7 8</td>
<td></td>
</tr>
</tbody>
</table>

   Additional delivery comments: ____________________________________________________________________________________________________________
_______________________________________________________________________________________________________________________________________

9. Was fundal pressure preformed: Yes No
   If yes, by whom and what was reason: ______________________________________________________________________________________________________

10. Who were the health care providers present at delivery?

   Primary delivery attendant: _____________________________
   CNM: _____________________________
   Resident(s) [include PGY year]: _____________________________
   Nurse 1: _____________________________
   Nurse 2: _____________________________
   Nurse 3: _____________________________
   Pediatrician: _____________________________
   Anesthesiology: _____________________________
   Other providers (please add name and role): _____________________________

11. Birth weight (grams)_____________________________

12. Was baby moving his/her extremities: Left Right Comment: ____________________________________________________

13. Was X-ray ordered for evaluation any bony fractures? Yes No

14. Were umbilical cord gases ordered? Yes No

15. Confirm that notes (e.g. medical, nursing, anesthesia, etc.) are consistent Yes No

16. Has delivery note been dictated? Yes No

17. Family counseled Yes No

18. Debriefing with appropriate personnel involved Yes No

19. Follow-up about condition of neonate in nursery Yes No

Signatures: _____________________________  _____________________________  _____________________________
Primary delivery provider  Primary delivery nurse  Other care providers in attendance

Figure 25.4  Suggested checklist for the documentation of delivery complicated by shoulder dystocia. CNM, certified nurse midwife.
If these steps all fail, one can repeat the sequence of initial maneuvers and/or attempt placement of the patient in the “all fours” position (Gaskin maneuver) (Figure 25.1). More aggressive, “heroic” maneuvers, which have very high frequency of maternal and fetal complications, may be considered after repeat failure of these first and second line interventions. Third-line maneuvers include intentional fracture of the fetal clavicle, Zavanelli (cephalic replacement), and symphysiotomy. In cases of intractable shoulder dystocia, a technique called posterior axilla sling traction (PAST) has been described. The PAST technique makes use of a sling (e.g., suction catheter or firm urinary catheter) that is placed around the posterior axilla and then used to use traction to either deliver the posterior shoulder or rotate it [46]. Fundal pressure should always be avoided in shoulder dystocia, as it may worsen impaction of fetal shoulder(s).

Counseling and Documentation

After completion of all necessary medical interventions (e.g., repair of maternal lacerations or treatment of hemorrhage), the delivery provider should sit down with patient and her family to review the delivery events, the management steps performed, and need for close newborn follow-up to evaluate for any neonatal injuries especially bony fracture and symptoms of BPI. Open and honest communication with the mother and family after delivery is recommended.

It is important to document in the medical record all maneuvers used in detail, including which shoulder was anterior, and the time from delivery of head to completion delivery of the body (Figure 25.4) [47, 48]. Introduction of a shoulder dystocia protocol has been associated not only with better documentation, but also with less incidence of brachial plexus palsy [47].

Anesthesia

An anesthesiologist should be present during cases of shoulder dystocia to ensure adequate analgesia, and prompt preparation for cesarean delivery if needed.

Neonatal/Infant Follow-Up

Table 25.4 describes most common palsies associated with shoulder dystocia. It is important to underscore that causation has never been proven between shoulder dystocia and palsies, and many palsies develop in babies delivered without shoulder dystocia. Erb’s palsy is by far the most common, and the one with the best prognosis. Occasionally BPI occurs on the posterior shoulder. Over 90% of neonates with Erb’s palsy with shoulder dystocia recover within 1 year, with most recovery already evident at 3–6 months. Erb’s palsy without shoulder dystocia has only a 60% recovery rate, and in about 68% of cases involves the posterior shoulder. Only about 40% of babies with Klumpke’s palsy recover by 1 year. If permanent injury (5%–8%) occurs, there is insufficient evidence to assess if surgery is effective.

REFERENCES


Postpartum hemorrhage, retained placenta, and uterine inversion

Jennifer Salati and Jorge E. Tolosa

KEY POINTS
- The most common complications of the third stage of labor are postpartum hemorrhage (PPH), retained placenta, and uterine inversion. Timely diagnosis, adequate physical examination, intravenous (IV) access, anesthesia, adequate use of blood products, nursing support and team work are important for the management of third-stage complications.
- In cases of PPH, one should perform uterine massage, explore for and repair any vaginal or cervical lacerations, and manually explore the uterus.
- **Oxytocin** 20–80 international units (IU) in 500–1000 cc normal saline given as a fast IV infusion, or 10 IU intramuscular (IM), should be used as a primary uterotonic for PPH.
- **Methergine** (except in women with hypertension) and **prostaglandin F2α** (except in women with asthma) are beneficial in the treatment of PPH. **Misoprostol** is also helpful as an adjunctive treatment of primary PPH.
- There is insufficient evidence with randomized controlled trials (RCTs) to assess all other interventions for treatment of PPH.
- There is insufficient evidence with RCTs to recommend pharmacologic agents for the treatment of retained placenta.

POSTPARTUM HEMORRHAGE

Definitions
The American College of Obstetrics and Gynecology (ACOG) defines PPH as an estimated blood loss (EBL) of ≤500 mL for a vaginal delivery, or ≤1000 mL for a cesarean delivery. Primary PPH is defined as blood loss within 24 hours of delivery exceeding 500 cc for vaginal deliveries and 1000 cc for cesarean delivery. Secondary PPH is defined as excessive blood loss after 24 hours and <12 weeks postpartum. There is no single definition for PPH that has been agreed upon internationally, and multiple guidelines from various national and expert authorities exist (Table 26.1). Recent initiatives have focused on reassessing the existing PPH definitions in an effort to improve clinical management and outcomes [1,2].

Incidence
Approximately 5%–15% of deliveries are complicated by PPH.

Etiology
Common:
- Uterine atony—lack of efficient uterine contraction (most common, accounts for 80%)
- Retained placenta
- Genital tract trauma—vaginal, cervical, and vulvar

Less common:
- Uterine rupture
- Uterine inversion
- Consumptive coagulopathy
- Inherited or acquired bleeding disorders

Risk Factors
At term, the uteroplacental circulation has a blood supply of 600–800 cc/minute. The blood volume expansion of 1.5–2.0 L in pregnant women provides a physiologic reserve for blood loss at delivery. Cessation of placental site blood flow is due mostly to effective myometrial contractions. Most major hemorrhage occurs within the first hour postpartum. Risk factors that contribute to decreased myometrial contractions include the following:
- Maternal factors:
  - First pregnancy
  - Maternal obesity
  - Previous PPH—10% recurrence risk
  - High parity
  - Coagulopathy
  - Thrombocytopenia
- Antepartum bleeding—placenta previa, abruption
- Chorioamnionitis
- Uterine fibroids
- Overdistension—macrosomia, multiple gestation, polyhydramnios
- Uterine tocolytics—magnesium, nitroglycerin, anesthetic gas
- Labor factors:
  - Precipitous delivery
  - Prolonged labor—first stage >24 hours
  - Prolonged oxytocin use, and/or maximum of ≥40–50 cc/minute (prolonged labor)

Complications
The most reliable estimates of global mortality for mothers in childbirth are reported to be between 250,000 and 300,000 annually [3]. This number has decreased significantly over the past two decades. Many of these deaths result from complications of the third stage of labor [3]. Ninety-nine percent of maternal deaths occur in low and middle-income countries where maternal mortality remains unacceptably high. In sub-Saharan Africa, the risk of maternal death is very high at 1:31, with at least 25% of those deaths due to hemorrhage [4], while in high income countries a woman’s lifetime risk of death during or following pregnancy is 1:4300. Secondary PPH is a significant contributor to maternal death mainly, but not only, in
low income countries. PPH has other serious complications including hypovolemic shock, disseminated intravascular coagulopathy (DIC), renal failure, hepatic failure, and adult respiratory distress syndrome (ARDS).

### Diagnosis

Regardless of the PPH definition being utilized, the appropriate management of PPH relies upon the clinician's assessment of risk for PPH (Table 26.2) and timely recognition of excessive blood loss. However, the assessment of peripartum blood loss has classically been determined by a provider's subjective visual estimation of blood loss. Visual EBL has consistently been shown to result in underestimation of large volume blood loss (>1000 mL) by up to 30%–50%, and overestimation of small volume of blood loss [5,6]. These types of EBL errors have been shown to occur similarly among providers at all levels of training [6]. Formal training in visual blood loss estimation improves accuracy for a period of time, but skills tend to decline after several months [7].

Some obstetric centers have adopted gravimetric methods for quantifying blood loss, including weighing of pads and sponges, and the use of calibrated under-buttock drapes. One study demonstrated that visual assessment of blood loss underestimated that calculated by weighing of pads and sponges by approximately 30% [8]. Another RCT showed that in a simulated environment, visual blood loss estimation using noncalibrated drapes resulted in underestimation by 15%–40% (greater error with larger volumes), whereas use of calibrated drapes resulted in <15% error at all volumes [9].

Blood loss calculators can be utilized by care teams to facilitate accurate quantification of blood loss after both cesarean and vaginal deliveries. Examples of calculation tools that can be integrated into an electronic medical record system can be found through the California Maternal Quality Care Collaborative (CMQCC) website (www.cmqcc.org/resources-tool-kits/toolkits/ob-hemorrhage-toolkit) and obstetrics (OB) hemorrhage toolkit resources.

### Management

**Primary PPH.** The management of PPH begins with diagnosis of PPH and obtaining help in a multidisciplinary fashion, including obstetricians, nurses (including charge nurse), anesthesia, and the blood bank (Figure 26.1). The early identification of risk factors is the first step to facilitating increased surveillance, adequate preparation and a more rapid response to hemorrhage if it does occur (Table 26.2). Risk factors should be evaluated in the prenatal period, and reassessed throughout the intrapartum and postpartum course.

A common accepted approach to the treatment of primary PPH includes:

**Logistics:**

- Obtain help
- Ensure adequate access with large bore IV (may need two sites)
- Monitor vital signs serially
- Consider placing Foley catheter to empty bladder and monitor urine output
- Obtain laboratory tests—complete blood count with platelets, blood type, antibody screen, fibrinogen, fibrin split products, prothrombin time, and partial prothrombin time

### Table 26.1 Definitions for Postpartum Hemorrhage

<table>
<thead>
<tr>
<th>Guidelines</th>
<th>Postpartum Hemorrhage Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACOG reVITALize Initiative (2014)</td>
<td>Blood loss ≥1000 mL or blood loss with sign/symptoms of hypovolemia in 24 hours after birth (includes intrapartum loss)</td>
</tr>
<tr>
<td>World Health Organization (2012)</td>
<td>Blood loss of ≥500 mL in 24 hours after birth</td>
</tr>
<tr>
<td>American College of Obstetricians and Gynecologists (2013)</td>
<td>Severe PPH-blood loss ≥1000 mL in 24 hours after birth</td>
</tr>
<tr>
<td>Royal Australian and New Zealand College of Obstetricians and Gynaecologists (2014)</td>
<td>Blood loss ≥500 mL following vaginal delivery or ≥1000 mL following cesarean delivery</td>
</tr>
<tr>
<td>Royal College of Obstetrician and Gynaecologists (2011)</td>
<td>Primary PPH-blood loss of 500–1000 mL in the absence of clinical signs of shock</td>
</tr>
<tr>
<td>Society of Obstetricians and Gynaecologists of Canada (2009)</td>
<td>Major PPH-blood loss of ≥1000 mL</td>
</tr>
</tbody>
</table>

**Abbreviations:** ACOG, American College of Obstetricians and Gynecologists; PPH, postpartum hemorrhage.

### Table 26.2 Example of Stratification System by Maternal Risk Factors for PPH

<table>
<thead>
<tr>
<th>PPH Risk</th>
<th>Risk Factors</th>
<th>Preparation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low</td>
<td>No previous CD</td>
<td>Clot only or type and screen</td>
</tr>
<tr>
<td></td>
<td>Singleton pregnancy</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Three prior VD</td>
<td></td>
</tr>
<tr>
<td></td>
<td>No bleeding disorder</td>
<td></td>
</tr>
<tr>
<td></td>
<td>No prior PPH</td>
<td></td>
</tr>
<tr>
<td>Medium</td>
<td>Prior CD</td>
<td>Type and screen or type and cross-match</td>
</tr>
<tr>
<td></td>
<td>Multiple gestation</td>
<td></td>
</tr>
<tr>
<td></td>
<td>≥4 prior VD</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Chorioamnionitis</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Polyhydramnios</td>
<td></td>
</tr>
<tr>
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<td>Macrosomia</td>
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<td>Abruptio</td>
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<tr>
<td></td>
<td>Prolonged labor (&gt;24 hours)</td>
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<tr>
<td></td>
<td>Prior PPH</td>
<td></td>
</tr>
<tr>
<td>High</td>
<td>Large uterine fibroids</td>
<td>Type and cross-match</td>
</tr>
<tr>
<td></td>
<td>Placenta previa</td>
<td>Consider alerting blood bank</td>
</tr>
<tr>
<td></td>
<td>Suspected placenta accrete</td>
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<tr>
<td></td>
<td>Coagulopathy</td>
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<tr>
<td></td>
<td>Platelets &lt;50,000</td>
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<tr>
<td></td>
<td>Multiple risk factors (see above)</td>
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</tbody>
</table>


**Abbreviations:** CD, cesarean delivery; VD, vaginal delivery.
POSTPARTUM HEMORRHAGE, RETAINED PLACENTA, AND UTERINE INVERSION

**Figure 26.1** Suggested management protocol for primary postpartum hemorrhage. Notes: CBC, complete blood count; IM, intramuscular; INR/PTT, international normalized ratio/partial thromboplastin time; IU, international units; IV, intravenous; T&S, type and screen.

Little evidence exists on which to base a protocol with respect to the order of administration of uterotonics for the treatment of PPH following vaginal delivery. It is also important to note that the literature on management of intraoperative and postpartum hemmorhage at the time of cesarean section is limited compared with vaginal delivery. Findings have been generalized to cesarean deliveries based on the assumption that they are likely to be beneficial. Traditionally, oxytocin and ergot alkaloids (e.g., methergine) are used as first-line agents, due to their pharmacokinetic profile and speed of action, with prostaglandin F2α and misoprostol employed as adjunctive therapies. There are few high-quality RCTs directly comparing oxytocin, ergot alkaloids, and prostaglandin F2α, to either placebo or to each other for treatment of PPH. Based on a recent systematic analysis, the addition of misoprostol to conventional injectable uterotonics conferred no added benefit with an increase in side effects [10]. Compared with oxytocin, misoprostol is less effective in primary treatment of PPH. Studies comparing misoprostol to ergot alkaloids have demonstrated conflicting results and the studies are small and heterogeneous.
Thus, there is no compelling evidence to suggest an alteration in uterotonic management at this time. However, it is important to note that in certain circumstances, rectal misoprostol may be a useful “first-line” therapy for the treatment of PPH. It may be a more appropriate first-line agent in the setting of no IV access, patients with contraindications to other uterotonic, or in resource-poor settings.

Other possible treatments include:
- Uterine curettage if retained placental products cannot be removed manually
- Transfusion of blood and blood products as necessary (Table 26.3)
- Hemostatic drugs
  - Factor VIIa—A single RCT demonstrated that recombinant human FVIIa may reduce the need for surgical intervention for severe PPH that has failed medical management [11]. This RCT was small and not placebo-controlled or blinded. Thus, larger blinded and placebo-controlled trials are needed to assess efficacy and safety
  - Tranexamic acid—Evidence for the use of tranexamic acid for treatment of PPH is limited and mixed. One randomized, open-label trial demonstrated decreased blood loss and morbidity associated with high-dose tranexamic acid [12]. Another historically controlled study did not demonstrate benefit [13]. Neither of these studies was large enough to assess effect on mortality, need for hysterectomy or safety
- If stable, consider pelvic arterial embolization done by interventional radiology (IR)
- If unstable (there are no high-quality RCTs investigating the following techniques)
  - Uterine tamponade—gauze packing, Bakri tamponade balloon catheter, or Foley catheter
  - Consider massive transfusion protocol (see Figure 26.2)
  - Laparotomy—uterine artery ligation, B-Lynch or square suture for compression, internal iliac artery ligation
  - Bimanual uterine compression
  - Aortic compression (temporary) and pelvic packing
  - Hysterectomy
  - Nonpneumatic antishock garment (NASG) is a noninvasive intervention. However, a recent systematic analysis incorporating 16 RCTs and 1683 patients identified no differences in the need for manual placental extraction with oxytocin (intraumbilical), prostaglandins (e.g., P2x, E2, or misoprostol) or nitroglycerin [16]. The greatest number of

### RETAINED PLACENTA

#### Definition
Placenta undelivered ≥30 minutes after infant delivery.

#### Incidence
Between 0.5% and 3%.

#### Complications
Hemorrhage, infection, and genital tract trauma.

#### Etiology
- Preterm birth—inversely proportional to gestational age.
- Cord avulsion—incidence up to 3% with controlled cord traction, especially by inexperienced operators.
- Placenta accreta (see Chapter 28).

#### Management
- Obtain adequate anesthesia
- Attempt manual placental extraction
  - One hand is placed over the abdomen to stabilize the uterus fundus, while the other hand is placed through the cervix into the uterus. The placenta is gently separated from the uterus starting at the superior placental edge. If possible, the placenta should be removed intact
  - Consider ultrasound to ascertain all placental tissue has been removed
  - The ultrasonographic appearance of the uterus immediately following delivery is variable. The sensitivity and positive predictive value of ultrasound for the detection of retained products of conception relatively low (44% and 58%, respectively). The specificity and negative predictive value are better (92% and 87%, respectively) [15]
  - Palpate and massage the fundus until firm
  - Proceed to operating room for curettage under ultrasound guidance if unsuccessful extraction or excessive bleeding
  - Consider diagnosis of placenta accreta

#### Pharmacologic Agents
In the past, pharmacologic interventions have been used in the management of retained placenta. The rationale was that stimulating uterine contractions with oxytocin or prostaglandins, or cervical relaxation with nitroglycerin, would facilitate spontaneous placental delivery and avoid further invasive interventions. However, a recent systematic analysis incorporating 16 RCTs and 1683 patients identified no differences in the need for manual placental extraction with oxytocin (intraumbilical), prostaglandins (e.g., P2x, E2, or misoprostol) or nitroglycerin [16]. The greatest number of

### Table 26.3 Blood Products

<table>
<thead>
<tr>
<th>Product</th>
<th>Volume (mL)</th>
<th>Contents</th>
<th>Effect (Per Unit)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Packed red blood cells</td>
<td>250–400</td>
<td>RBC, WBC, plasma</td>
<td>Increase Hgb by 1g/dL or Hct by 3%</td>
</tr>
<tr>
<td>Platelets</td>
<td>50</td>
<td>Platelets, RBC, WBC, plasma</td>
<td>Increase platelet count by 5,000–10,000 mm³</td>
</tr>
<tr>
<td>Fresh frozen plasma</td>
<td>200–250</td>
<td>Fibrinogen, Antithrombin III, clotting factors, plasma</td>
<td>Increase fibrinogen by 5–10 mg/dL</td>
</tr>
<tr>
<td>Cryoprecipitate</td>
<td>20–40</td>
<td>Fibrinogen, factor VIII, vWF, factor XIII</td>
<td>Increase fibrinogen by 5–10 mg/dL</td>
</tr>
</tbody>
</table>

**Abbreviations:** Hct, hematocrit; Hgb, hemoglobin; RBC, red blood cells; WBC, white blood cells; vWF, Von Willebrand factor.
studies [10] investigated intraumbilical administration of oxytocin. They concluded that there is no evidence supporting the use of intraumbilical oxytocin for retained placenta, and that there has not been adequate evaluation of other pharmacologic interventions by RCT. Therefore, there is insufficient evidence to recommend pharmacologic agents for the treatment of retained placenta.

**Antibiotics**
There is insufficient evidence to assess the effects of prophylactic use of antibiotics following manual removal of the placenta [17]. The World Health Organization (WHO) recommends single-dose penicillin or first-generation cephalosporin based on one retrospective study [18,19].

**UTERINE INVERSION**
**Definition**
The collapse of the uterine fundus into the uterine cavity.

**Incidence**
Approximately 1 in 2500 deliveries.

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**Figure 26.2** Suggested massive transfusion protocol. FFP, fresh frozen plasma; pRBC, packed red blood cells.

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**Risk Factors**
- Excessive umbilical cord traction
- Fundal pressure
- Fundal cord insertions
- Abnormal placentation

**Management**
There are no randomized trials to determine the optimal management for uterine inversion. Suggested management may include the following:
- Obtain assistance from other obstetricians or experienced providers and nursing staff
- Obtain adequate anesthesia
- Consider immediate transfer to the operating room
- Obtain large bore IV access
- IV fluid therapy
- If placenta has not yet delivered, avoid separation to reduce bleeding
- Consider uterine relaxing agents
  - Nitroglycerin 50–500 mg orally or inhaled per anesthesia—usually preferred as first line
• Magnesium sulfate 4–6 grams IV bolus
• Terbutaline IV 0.25 subcutaneously (SQ)

• Manual manipulation of the uterus
  • Manually reduce the uterine fundus by placing the hand through the cervix into the uterine cavity and applying gentle pressure to the inverted fundus. Cup the uterus in palm with fingers posteriorly and thumb anteriorly (no use of fist).
  • Once the fundus has been manually replaced, administer uterotonic therapy (oxytocin) followed by methergine to maintain uterine tone and prevent reinversion.

• Surgical intervention—laparotomy (rarely needed)
  • Huntington procedure—clamps are placed on the round ligaments 2 cm deep in the inversion and gentle upward traction applied. Repeat clamping as necessary.
  • Haultain procedure—vertical incision is made in the posterior portion of the inversion ring to increase its size and to reposition the uterus.
  • Uterotonic agents when uterus repositioned.

• If present, treat PPH or retained placenta as described above.

REFERENCES
Late-term and postterm pregnancies

Rebecca Jackson

KEY POINTS
• Postterm pregnancy is defined as a pregnancy that has lasted until ≥42 weeks, or ≥294 days.
• Late-term pregnancy is defined as a pregnancy that has lasted 41 0/7–41 6/7 weeks of gestation.
• Late-term and postterm pregnancies have twofold increase in risk of macrosomia, and subsequent increase in labor dystocia, operative vaginal delivery, cesarean delivery, perineal injury, and shoulder dystocia.
• Postterm neonates are at increased risk for seizures, meconium aspiration, 5-minute Apgar scores less than 4, and admission to neonatal intensive care unit. Postterm pregnancies are also at increased risk for intrauterine infection, oligohydramnios, nonreassuring fetal heart testing (NRFHT), low umbilical artery pH, and perinatal mortality.
• Pregnancies with risk factors such as maternal (e.g., hypertension and diabetes) and fetal (growth restriction) diseases should be delivered at term or as described in the pertinent guidelines.
• Prevention of postterm pregnancy can be achieved with accurate dating by routine first-trimester ultrasound and with serial membrane stripping starting at 38 weeks.
• There is insufficient evidence to assess the efficacy of antepartum testing for pregnancies after their due date. Twice-weekly fetal testing with nonstress test (NST), or NST and maximum vertical pocket (MVP) of amniotic fluid volume (AFV), or biophysical profile (BPP), has been proposed, starting at 41 weeks.
• Routine induction of labor at ≥41 weeks reduces perinatal mortality, due to a decrease in fetal and neonatal deaths. Routine induction of labor in this setting is also associated with a significant 11% decrease in the incidence of cesarean delivery, and a significant 50% decrease in meconium aspiration syndrome. Therefore, pregnancies reaching 41 weeks should be recommended induction at 41 0/7–41 6/7 weeks, unless contraindications for induction exist and/or a cesarean is indicated.
• In women with a prior cesarean delivery, induction of labor is associated with an increase in uterine rupture in comparison to an unscarred uterus and in comparison to spontaneous labor. Nonetheless, induction of labor after appropriate counseling is feasible (e.g., with balloon catheter) in pregnancies ≥41 weeks with a prior cesarean.

DIAGNOSIS/DEFINITION
Postterm pregnancy is defined as pregnancy that has lasted until ≥42 weeks, or ≥294 days, or ≥14 days after the due date (estimated date of confinement [EDC]) [1]. Late-term pregnancy can be defined as a pregnancy that has lasted until ≥41 weeks, or ≥287 days, or ≥7 days after the due date (EDC) [1]. The term “postdates” can signify a pregnancy that lasted until ≥40 weeks, or ≥280 days, but is often defined differently in the literature and should be avoided [1]. All these definitions may have slightly different meanings in the literature, so it is important to be clear when using these terms. These definitions and this guideline pertain to singleton gestations. For multiple gestations, please refer to Chapter 44 in Maternal-Fetal Evidence Based Guidelines.

EPIDEMIOLOGY/INCIDENCE
In 2014, in the United States, the incidence of late-term pregnancy was 6.31% and of postterm pregnancy was 0.41% [2]. These incidences have significantly decreased in the last few years.

ETIOLOGY/BASIC PATHOPHYSIOLOGY
The most frequent cause of postterm pregnancy is an error in dating [1,3]. See Chapter 4 for accurate dating criteria and their benefits, as well as the following sections.

RISK FACTORS/ASSOCIATIONS
Poor (wrong) dating; prior postterm pregnancy; obesity; nulliparity; long (>28 days) cycles without early ultrasound; placental sulfatase deficiency; anencephaly; and male fetus. Obesity appears to be one preconceptionally modifiable risk factor for postterm pregnancy [4].

COMPLICATIONS
Perinatal
Many epidemiological studies have shown that fetal and neonatal complications increase with gestational age after 41 weeks. Meconium aspiration, intrauterine infection, oligohydramnios, macrosomia, NRFHT, low umbilical artery pH, and low 5-minute Apgar score have all been associated with postterm pregnancy. Perinatal mortality (fetal and neonatal deaths) is twice as high at ≥42 weeks and six times as high at ≥43 weeks compared with the rate for pregnancies delivered between 39 and 40 weeks [1,5].

Two markers of immediate neonatal morbidity, umbilical artery pH <7 and base excess ≥12, have been shown to increase in a continuous manner in pregnancies beyond 40 weeks, and increase by odds ratio (OR) 1.65 for pH <7 for pregnancies between 41 and 42 weeks [6].

Neonatal mortality of normal-weight infants appears to be higher for infants born between 41 and 0 days and 41 weeks and 6 days compared with infants born between 38 weeks and 40 weeks and 6 days (OR 1.37, 95% confidence interval [CI] 1.08–1.73) [7].

These data are, however, mixed and may vary based on whether risk factors such as intrauterine growth restriction (IUGR) are included in the analysis. In a review of epidemiological studies published between 1990–2004, 7 out of 11 studies demonstrated an increased stillbirth risk with postterm pregnancies and four suggested that other issues such as IUGR
or fetal anomalies had a greater contribution to the stillbirth rate than did prolonged pregnancy [8].

Postmaturity syndrome is present in about 10%–20% of neonates born postterm. These fetuses have decreased subcutaneous fat and lack of vernix and lanugo. Postmature neonates are also more likely to have meconium staining of the amniotic fluid, skin, membranes and umbilical cord. [1].

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Maternal
Women giving birth late-term and postterm are at increased risk of perineal injury, infection, postpartum hemorrhage, and cesarean delivery [1].

PREGNANCY CONSIDERATIONS
Every woman should be counseled early in pregnancy that up to 50% of gestations, especially in nulliparous women, last until past the due date. This is physiologic and natural for humans. The incidence of fetal death is significantly higher than that of neonatal death at ≥283 days (≥40 weeks and 3 days) [9]. In large series, delivery at 38 weeks is associated with the lowest risk of perinatal death, but this risk is low at <1 to 2/1000 up to 41 weeks and 6 days [10].

It is important to identify maternal (e.g., hypertension and diabetes) and fetal (e.g., growth restriction) risk factors that would necessitate special management, as described in the specific chapters in Maternal-Fetal Evidence Based Guidelines. The OR for stillbirths and neonatal mortality increases in gestations from 40 weeks onward, and this increase is compounded 7- to 10-fold when the fetus is growth restricted [11].

Prevention
Routine Early Ultrasound to Reduce Postterm Pregnancies
Accurate assessment of gestational age is extremely important in improving perinatal morbidity and mortality.

First-trimester ultrasound is the most accurate for pregnancy dating. Early ultrasound also reduces the number of postterm inductions.

Over 40% of women randomized to undergo first-trimester screening had their gestational age adjusted based on the ultrasound measurement of crown–rump length, whereas the corresponding number was only 10.9% for women assigned to second-trimester ultrasound. Furthermore, 4.8% in the first-trimester screening group compared with 13% in the second-trimester ultrasound group had labor induced for a postterm pregnancy (relative risk [RR] 0.37, 95% CI 0.14–0.96), which is a 63% reduction in the induction rate for postterm pregnancy [11,12]. Compared with no routine early ultrasound, routine early pregnancy (<20 weeks) ultrasound reduces by 32%–39% the incidence of postterm pregnancy and of induction for postterm pregnancy [4,12,13] (see also Chapter 4).

MANAGEMENT (FIGURE 27.1)
Preconception Counseling
Women with prior postterm pregnancy are at increased risk for recurrent postterm pregnancy. Prevention strategies including weight loss for obese women, appropriate weight gain during the pregnancy, early initiation of prenatal care, and early ultrasound screening should be discussed [3].

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Figure 27.1  Management of late-term pregnancy. Notes: 2x/week, twice per week; AFV, amniotic fluid volume; EFW, estimated fetal weight; FGR, fetal growth restriction; NST, nonstress test. *Risk factor examples include hypertension, diabetes mellitus, FGR, multiple gestation, etc. Please see guideline pertinent to specific risk factor for management. *Suggest induction for all women between 410 and 416 weeks. †See Chapter 21 for effective induction management.
Stripping/Sweeping of Membranes
Compared with no sweeping, sweeping of the membranes, performed weekly as a general policy in women at term (e.g., weekly starting at 38 weeks), is associated with reduced duration of pregnancy and reduced frequency of pregnancy continuing beyond 41 and 42 weeks [14,15]. To avoid one formal induction of labor, sweeping of membranes must be performed in eight women. Risk of cesarean section, maternal, or neonatal infection is similar. Serial sweeping of membranes starting at 41 weeks every 48 hours also decreases the risk of postterm pregnancy from 41% to 23%, with efficacy both in nulliparous and multiparous women [16]. Discomfort during vaginal examination and other adverse effects (bleeding and irregular contractions) are more frequently reported by women allocated to sweeping, but its safety has been confirmed in multiple studies [14–17] (see also Chapter 20).

Breast and Nipple Stimulation to Reduce Postterm Pregnancies
Breast and nipple stimulation daily starting at 39 weeks has not been sufficiently studied to ascertain safety, but it does appear to reduce the incidence of postterm pregnancy by 48% [18,19].

Antepartum Testing
There are insufficient data to assess the best mode of fetal monitoring after the due date, as there are no trials to assess the effect of antepartum testing on these pregnancies compared with no testing. Since fetal death rates incrementally increase after the due date, it seems reasonable to test fetuses to assure well-being, especially at ≥41 weeks [1,4,9]. The most used options include NST (also called cardiotocography or CTG), BPP, and modified BPP. Modified BPP includes NST and ultrasound measurement of MVP of AFV. Others have been described, with even less evidence for efficacy. Doppler of any vessel, including the umbilical artery, is not effective in the management of postterm pregnancy. Compared with fetal monitoring using NST and AFV, BPP (computerized cardiotocography, MVP, fetal breathing, fetal tone, and fetal body movements) is associated with increased incidence of inductions and similar outcomes in a small trial in women ≥42 weeks [20]. At ≥41 weeks, twice-weekly testing (e.g., with NST and MVP) is recommended [1], but not based on trials (see also Chapter 56 in Maternal-Fetal Evidence Based Guidelines). In addition, an effort should be made to assess fetal weight, either clinically or, if this assessment is limited, by ultrasound, as the known increase in fetal and neonatal mortality in pregnancies beyond their due date is even higher in IUGR fetuses [11,21].

Interventions
Routine Induction of Labor at ≥41 Weeks
A policy of labor induction at 41 completed weeks (41–41 6/7) or beyond is associated with significantly fewer perinatal deaths (1/2814 vs. 9/2785; RR 0.30, 95% CI 0.09–0.99) compared with expectant management, with induction not before 42 weeks [21]. If deaths due to congenital abnormality are excluded, no deaths remain in the labor induction group and eleven deaths remain in the no induction group. There were fewer cesarean sections in the induction group (RR 0.89, 95% CI 0.81–0.97). Labor induction at 41 weeks also significantly reduces the risk of perinatal meconium aspiration syndrome compared with expectant management (RR 0.50, 95% CI 0.34–0.73). Other maternal and perinatal outcomes are similar between the groups. It is important to recognize, however, that in general the outcomes are very good with both expectant management and induction, with the absolute perinatal death rate/1000 ongoing pregnancies no higher than 1.2/1000 at 42 weeks, increasing up to 6/1000 ongoing pregnancies at 43 weeks [22]. About 410 inductions (95% CI 322–1492) would need to be done at 41 weeks in order to prevent one perinatal death [21].

Routine induction is more cost-effective than expectant management [23]. Several studies have shown that patient satisfaction is higher with induction of labor [21–23]. Iatrogenic prematurity should be avoided by careful assessment of gestational age. This risk should be minimal with the advent of widespread early ultrasound use to confirm pregnancy dating. Even planned induction without medical indications at 39 0/7–40 6/7 weeks has been shown not to be associated with an increase with cesarean delivery. [24] See also Chapter 21 for other induction risks and benefits, as well as management of induction.

Women with a favorable cervix. No trials have focused on or included in sufficient numbers pregnancies ≥41 weeks with a favorable cervix (i.e., Bishop score ≥9 or transvaginal ultrasound cervical length [TVU CL] <15 mm). As the complications of induction, particularly a failed induction and unnecessary subsequent cesarean section, are low in women with a favorable cervix, and there appears to be a trend toward decreased perinatal deaths and a statistically significant reduction in meconium aspiration, it seems reasonable to also recommend induction at 41 0/7–41 6/7 weeks (41 and 0/7 days) for these women [21,23].

Women with an unfavorable cervix. Cervical ripening is recommended for women with unfavorable cervix prior to labor induction. If cervical ripening is used, there is no difference in the incidence of cesarean section in women who were induced or expectantly managed [1,23,25,26]. A discussion about induction of labor after 41 weeks versus expectant management should occur with the patient, and management should include consideration for patient preference and access to antenatal testing (see also Chapter 21). Induction should absolutely be recommended for all patients before 42 0/7 weeks. One hundred and ninety-five inductions would prevent one perinatal death in this group [27].

Women with a prior cesarean delivery. In women with a prior cesarean delivery, induction is associated with a higher incidence of uterine rupture, especially in the nulliparous woman with an unfavorable cervix. The rate of uterine rupture is about 0.4%–0.5% for spontaneous labor, 0.8% for labor induced without prostaglandins, and 2.2% for prostaglandin induced labor [28]. Prostaglandins should, in general, not be used for cervical ripening in a woman with a prior cesarean section. If the woman desires vaginal birth after cesarean (VBAC), it seems reasonable to wait until 40–41 weeks for spontaneous labor. There are insufficient data to assess the risks of trial of labor after cesarean section in a postterm gestation, but if the cervix is favorable, induction can be performed after thorough discussion of the risks and benefits. If the cervix is unfavorable, the patient should be apprised of the higher risks of uterine rupture, as well as the risks of a failed induction with an unfavorable cervix. A repeat cesarean section can be offered to decrease these risks [1,29] (see also Chapter 14).

REFERENCES


12. Crowley, P. Interventions for preventing or improving the outcome of delivery at or beyond term. Cochrane Database Syst Rev. 2005;(4):CD000331. [Meta-analysis: 26 RCTs (variable quality), n = >25,000; early ultrasound: 4 RCTs, n = 21,776; induction at ≥41 weeks: 19 RCTs, n = 7925]


Placental disorders include placenta previa, placenta accreta, and vasa previa. For normal or abnormal third stage, including postpartum hemorrhage, retained placenta, and uterine inversion, see Chapters 9 and 26.

**KEY POINTS**

**Placenta Previa**
- Placenta previa is defined as a placenta that covers the internal os.
- Low-lying placenta is defined as a placenta that comes within 0.1–2.0 cm of the internal os but does not cover it.
- Placental location should be assessed when the fetal anatomic survey ultrasound (usually 18–24 weeks) is performed. If a placenta previa is suspected on transabdominal ultrasound, a transvaginal ultrasound (TVU) should be performed to confirm the diagnosis.
- The risk that a placenta previa diagnosed in the antepartum period remains a placenta previa at the time of delivery depends on several factors, especially gestational age (GA) at detection, the placental distance from or extent of overlap of the internal os, and the a priori risk of placenta previa related to other risk factors (e.g., prior cesarean delivery). Most placenta previas diagnosed before the third trimester do not remain placenta previas at the time of delivery.
- Women who have the inferior edge of the placenta ≥1 cm from the internal os at around 20 weeks usually do not require further ultrasounds for placental location.
- All patients with the inferior placenta edge <1 cm from the internal os or overlying the os at around 20 weeks should be rescanned at least once at approximately 32–35 weeks’ gestation to reassess for placental location.
- All patients with prior cesarean delivery and placenta previa that extends anteriorly should have evidence of placenta accreta assessed ultrasonographically.
- Women with a placenta previa should be delivered by cesarean. If the placental lower edge is within 1–19 mm of the internal os, a trial of labor can be attempted, but the risk of significant bleeding during labor may be higher, especially in those with a distance of only 1–10 mm.
- The optimal GA for delivery of an asymptomatic woman (i.e., a woman without acute bleeding) with vasa previa is unknown, but many experts recommend planned cesarean delivery at approximately 36 0/7–37 6/7 weeks.

**Vasa Previa**
- Vasa previa exists when fetal vessels, unprotected by the umbilical cord or placental tissue, run through the membranes and over the internal os. In this circumstance, rupture of the membranes can lead to rupture of these fetal vessels, with a significant possibility of fetal death. Women with risk factors, such as a velamentous cord insertion, resolved placenta previa, or a succenturiate or bilobed placenta in which intervening vessels may cross the cervix, should have a transvaginal ultrasound (TVU) to detect vasa previa.
- The optimal GA for delivery of an asymptomatic woman (i.e., a woman without acute bleeding) with vasa previa is unknown, but many experts recommend planned cesarean delivery at approximately 34 0/7–35 6/7 weeks.

**Placenta Accreta**
- Risk factors for placenta accreta include prior cesarean delivery; placenta previa; prior uterine surgery (e.g., prior myomectomy or prior multiple dilation and evacuations [D&Es]); Asherman’s syndrome; submucous leiomyomata; maternal age ≥35 years; multiparity; and smoking.
- Maternal complications of placenta accreta include hysterectomy, injury to other organs, blood transfusion, disseminated intravascular coagulation (DIC), infection, and death. Fetal complications include preterm birth (PTB) and small for gestational age (SGA).
- There are no randomized trials to assess efficacy of different surgical interventions in or the approaches to the management of placenta accreta. Planned hysterectomy may be beneficial in cases in which the diagnosis is highly suspected, especially for the woman who does not desire further fertility.
- There are no randomized trials to assess the optimal gestation age for delivery of pregnancies with placenta accreta, but The American College of Obstetricians and Gynecologists (ACOG) and Society for Maternal-Fetal Medicine (SMFM) recommend planned cesarean delivery at around 34 0/7–35 6/7 weeks.

**PLACENTA PREVIA**

**Diagnoses/Definitions**

Placenta previa is defined as a placenta that covers the internal os. The diagnosis of placenta previa is established by TVU. The terms partial, marginal, complete, or incomplete placenta previa are rooted in preultrasound physical examinations, have been used to signify different conditions, and therefore should be avoided. Moreover, many studies have used these terms to signify different conditions. When the placenta does not cover the internal os, rather than use ambiguous language, it is best to describe the distance of the tip of the placenta from the internal os. Low-lying placenta is defined as a placenta that comes within 0.1–2.0 cm of the internal os but does not cover it.

**Symptoms**

Approximately two-thirds of women with placenta previas have antepartum vaginal bleeding. Therefore, about 33% do not have any symptoms before delivery.
Epidemiology/Incidence
The incidence of placenta previa at term is approximately 0.5% [1]. The frequency of placenta previa is higher earlier in gestation, but many of these cases will resolve. The likelihood of resolution is inversely proportional to GA at the time of examination (Table 28.1) [2].

Risk Factors/Associations
Prior cesarean delivery (Table 28.2), any other uterine surgery (e.g., myomectomy, D&E, dilation and curettage [D&C], and hysteroscopy) involving the uterine cavity, prior placenta previa, and multiparity [3–5].

Complications
Placenta accreta, antepartum and/or postpartum hemorrhage, PTB, and perinatal death [6].

Management

Principles
Placenta location should be assessed at the time of the fetal anatomic survey (usually 18–24 weeks). If placenta previa is suspected by transabdominal examination, a TVU should be performed for confirmation of the diagnosis (Figure 28.1). A single antenatal ultrasound that detects a placenta previa, however, may not indicate that a placenta previa will be present at delivery, as the relative position of the placenta with respect to the internal os will change as gestation progresses [2]. The reason for this change in relative position is not placental migration but likely due to the growth and development of the lower uterine segment. Atrophy of placental cells overlying the os also has been postulated as a contributing factor to this apparent positional change. This phenomenon has been cited as the reason that massa previa can be seen in this setting (i.e., when an initially diagnosed placenta previa resolves).

Workup

Examination. Because of the reliability of ultrasound for diagnosis of previa, the technique of double setup examination is unnecessary. If employed, double setup examination should be performed in a setting with the ability to proceed in a prompt fashion with cesarean delivery if indicated.

Ultrasound. TVU is the gold standard for the diagnosis of placenta previa (Figure 28.2). It is very safe in these women [3]. The risk that a placenta previa that is diagnosed with ultrasound is present at delivery depends on several factors, including GA at detection, the placental distance from or extent of overlap of the internal os, as well as a woman's a priori risk of placenta previa (e.g., related to a prior cesarean delivery). Most previas diagnosed before the third trimester resolve (Table 28.1) [2,7]. If a placenta previa “resolves” but remains proximate to the internal cervical os, a woman still may have an increased risk of third-trimester bleeding, intrapartum hemorrhage, and Cesarean delivery [8,9].

Women who have the inferior edge of the placenta ≥1 cm from the internal os at around 20 weeks usually do not require further ultrasounds for placental location as they are exceedingly unlikely to have a placenta that is low-lying or a previa at delivery. Women who have the inferior placental edge overlapping the internal os by ≥25 mm at around 20 weeks have been reported to usually have persistence of previa even by term [10,11] (Table 28.1).

All patients with suspected placenta previa in the second trimester should be rescaned at approximately 32–35 weeks' gestation to assess for persistence of placenta previa (Figure 28.1). Additionally, even if a placenta previa is no longer present, measuring the distance from the placental edge to the internal os in the third trimester can help estimate the risk of bleeding with a trial of labor [8,9].

All patients with prior cesarean delivery (or other uterine surgery) and placenta previa that extends anteriorly should have evidence of placenta accreta assessed ultrasonographically [5,6] (see the section “Placenta Accreta”).

Prenatal Care
All patients with placenta previa and antenatal bleeding in the third trimester should be advised about pelvic rest (no vaginal penetration). Pelvic rest recommendations in women with a placenta that is near but not overlapping the internal os should be individualized and based on bleeding during the pregnancy as well as the distance between the placenta and the internal os. There is insufficient evidence to support the

Table 28.1 Persistence of Placenta Previa until Delivery Stratified by GA at Ultrasound Detection, Type of Previa, and Prior Cesarean Delivery

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Incomplete previa,* no prior cesarean</td>
<td>6</td>
<td>11</td>
<td>12</td>
<td>35</td>
<td>39</td>
</tr>
<tr>
<td>Incomplete previa,* prior cesarean</td>
<td>7</td>
<td>50</td>
<td>40</td>
<td>38</td>
<td>63</td>
</tr>
<tr>
<td>Complete previa, no prior cesarean</td>
<td>20</td>
<td>45</td>
<td>56</td>
<td>89</td>
<td>90</td>
</tr>
<tr>
<td>Complete previa, prior cesarean</td>
<td>41</td>
<td>73</td>
<td>84</td>
<td>88</td>
<td>89</td>
</tr>
</tbody>
</table>


Abbreviation: GA, gestational age.

All data presented as percentages.

*Incomplete previa defined as inferior edge of the placenta partially covering or reaching the margin of the internal os.

Table 28.2 Risk of Placenta Previa and/or Accreta Stratified by Number of Prior Cesarean Deliveries

<table>
<thead>
<tr>
<th></th>
<th>First CD</th>
<th>Second CD</th>
<th>Third CD</th>
<th>Fourth CD</th>
<th>Fifth CD</th>
<th>≥Sixth CD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Previa</td>
<td>6.4</td>
<td>1.3</td>
<td>1.1</td>
<td>2.3</td>
<td>2.3</td>
<td>3.4</td>
</tr>
<tr>
<td>Accreta (no previa)</td>
<td>0.03</td>
<td>0.2</td>
<td>0.1</td>
<td>0.8</td>
<td>0.8</td>
<td>4.7</td>
</tr>
<tr>
<td>Accreta (previa)</td>
<td>3.3</td>
<td>11</td>
<td>40</td>
<td>61</td>
<td>67</td>
<td>67</td>
</tr>
</tbody>
</table>


Abbreviation: CD, cesarean delivery.

All data presented as percentage.
use of routine bed rest in women with placenta previa or to be certain who are optimal candidates, when asymptomatic, for hospitalization [12]. Moreover, while pelvic rest, i.e., nothing per vagina, is often suggested, there is no evidence for its effectiveness. There is no evidence to support the use of autologous blood donation/transfusion for placenta previa [13].

**Therapy/Interventions**

*Management at home versus hospitalization.* There has been no evidence of any clear advantage to a policy of home versus routine hospital care, with similar maternal and fetal outcomes demonstrated in the trials that do exist. The only difference is that, compared with hospitalization, management at home, not surprisingly, has been associated with a reduced length of stay in the hospital [12]. There are no data to allow certainty as to who are the optimal candidates for inpatient management. Women who, traditionally, have been more likely to be managed in the hospital even though they are not acutely symptomatic are those who have had recurrent episodes of antepartum hemorrhage (e.g., three or more episodes of bleeding); have other medical complications that increase their risks (e.g., congenital cardiac disease); or are unable to easily access the hospital in an emergency (e.g., lack of telephone at home and far distance from an adequate care facility).
Often, after an initial antenatal bleeding episode related to placenta previa, if several days of no further bleeding have occurred and there is no other indication for hospitalization, women may be managed as outpatients.

Cervical cerclage. Cervical cerclage has not been proven to be an effective intervention for women with placenta previa [12]. Although the meta-analysis in the Cochrane review that evaluated cervical cerclage in the setting of placenta previa revealed that women randomized to cerclage had a reduced length of inpatient hospitalization, as well as reduced risks of an SGA birth or delivery <34 weeks, the benefits were evident in studies with lower methodological quality, and thus there is not yet sufficient evidence to justify this intervention.

Tocolysis. There is insufficient evidence to assess the benefits of tocolysis for treatment of preterm labor (PTL) specifically in women with a placenta previa. It is reasonable to consider tocolysis, as for women without a placenta previa, if PTL is diagnosed and a course of steroids has not yet been administered [14]. Acute bleeding with instability of the mother or fetus is considered a contraindication to tocolysis.

Antepartum testing. There are insufficient data to assess the usefulness of routine antenatal testing in improving outcomes, and this strategy is presently not indicated.

Delivery

Timing. There is insufficient evidence to certain of the optimal GA of delivery for women with placenta previa who are not acutely bleeding, although experts have recommended approximately 36 0/7–37 6/7 weeks [15,16] (Figure 28.1). In the setting of acute bleeding, the timing of delivery depends upon individualization of patient circumstances, taking into account GA, amount of bleeding, and other comorbidities.

Mode. Delivery of women with a placenta previa should be by cesarean. Preoperative ultrasonography to assess placental location is useful in determining the optimal place for the uterine incision (Figure 28.1). Those with a placenta not covering the internal os but in the lower segment may have a trial of labor offered, with individual circumstances (e.g., the presence of antepartum bleeding) and the risk of bleeding at delivery taken into consideration. In a recent series, 26/28 (93%) women who had a placental edge to cervical os distance of 1–20 mm and who underwent a trial of labor delivered vaginally [8,17]. Women with a placental edge that is within 1 cm of the internal os can have uncomplicated vaginal deliveries, although their risk of bleeding and requiring a cesarean is higher than in those women whose placenta is 10–20 mm distant from the os (risks of cesarean 75% and 31%, respectively) [8] (Table 28.3). As such, depending upon the distance of the placental edge from the internal os as well as other factors, these patients (i.e., those with a placenta that is not covering the internal os but that is 2 cm or less from the internal os prior to delivery) should have their circumstances discussed and be delivered in facilities with the capacity to perform emergent operative delivery if necessary [18]. In women with the placenta 2 cm or more from the internal os, a trial of labor should be encouraged.

PLACENTA ACCRETA

Diagnosis/Definition

Placenta accreta is defined as a placenta that is abnormally adherent to the uterus. The diagnosis of this condition can be quite difficult, as full ascertainment would require postpartum histologic examination of the entire uteroplacental interface with both placenta and uterus available. Since this is not typically possible, in cases of clinically suspected placenta accreta, failure to demonstrate abnormal villous invasion by pathologic examination cannot always be used to exclude this diagnosis [19]. Therefore, the diagnosis is often made clinically at delivery. The antenatal suspicion of placenta accreta can be based on history (e.g., prior cesarean delivery and an anterior placenta previa) or imaging findings (e.g., by ultrasound or magnetic resonance imaging [MRI]), but no method can provide 100% accuracy in antenatal diagnosis.

Epidemiology/Incidence

Traditionally 1/2500 deliveries, although evidence of increasing frequency (3/1000 or more), thought to be related to the increased rate of cesarean delivery, has been reported [20,21].

Etiology/Pathophysiology

Abnormal adherence of chorionic villi to myometrium, associated with total or partial lack of the decidua basalis layer. Cesarean scar pregnancy is diagnosed by ultrasound often in the first trimester, and can be an early sign of later development of placenta accreta [22].

Classification

Abnormally invasive placentas may be categorized according to the depth of their invasion [23].

- Placenta accreta: Chorionic villi are attached directly to, but do not invade, the myometrium.
- Placenta increta: Placental villi invade into the myometrium.
- Placenta percreta: Placental villi invade through the myometrium into the uterine serosa; adjacent organs (e.g., the bladder) may be involved, but this extension is not necessary for the diagnosis.

Risk Factors/Associations

Prior cesarean delivery; placenta previa (Table 28.2); prior uterine surgery involving the cavity (e.g., prior myomectomy; D&amp;E; Asherman’s syndrome; submucous leiomyomata; cesarean scar pregnancy [5,6,23].

Complications

Maternal. Blood transfusion, infection, DIC, injury to other organs, hysterectomy, venous thromboembolism, and death [5,6].

Perinatal. PTB and SGA [24].

| Table 28.3 Selected Outcomes with Low-Lying Placenta According to Placental Edge to Internal Cervical Os Distance |
|---|---|---|
| Distance between Placental Edge and Internal Cervical Os | 1–10 mm | 11–20 mm |
| Cesarean delivery | 75% | 31% |
| Antepartum hemorrhage | 29% | 3% |
| Postpartum hemorrhage | 21% | 10% |
| Postpartum hemorrhage | 8% | 10% |
| >1000 mL | |

Management
There are no trials to assess the comparative effectiveness of different interventions (e.g., planned hysterectomy versus attempted placental extraction for suspected placenta accreta; and preoperative placement versus no preoperative placement of catheters to allow the inflation for internal iliac balloons) in the management of placenta accreta.

Workup
No one imaging modality has been shown to be able to accurately diagnose placenta accreta with 100% sensitivity or specificity. Given the frequency of ultrasonographic imaging during pregnancy, this modality has been most frequently assessed with regard to attempting an antepartum diagnosis (Figure 28.3). Ultrasonographic findings that have been reported in association with placenta accreta are shown in Table 28.4 [23,25–30]. However, even the combination of all these signs is not 100% sensitive for the diagnosis of accreta, and the sensitivity, specificity, positive and negative predictive values for individual signs and combinations of signs have varied substantially in published studies, but have been commonly reported to be about >60%–80% [31,32]. Further imaging modalities to evaluate the possibility of placenta accreta include MRI, which may be informative but also is not considered to be completely diagnostic [28,29]. MRI is not recommended for routine use in suspected accreta. It may be useful as an adjunctive tool if the placenta is posterior or to assess invasion of adjacent organs in suspected percreta [33]. Cystoscopy can be considered as an adjunctive tool to assess for the possibility of placenta percreta in cases where bladder invasion is highly suspected due to radiologic studies or to signs such as frank blood in the urine.

Preparations and Plans for Delivery
If placenta accreta is suspected, appropriate counseling and preparations should be made (Table 28.5) [23]. Labor and delivery staff (nursing and anesthesia) as well as blood bank should be notified regarding delivery plans, and the delivery should occur in a location that has the capacity to provide large volumes of blood transfusion and emergent surgery (including hysterectomy). It should be considered whether the particular circumstances would suggest the need for additional surgical services (e.g., gynecologic oncology, urology, general surgery, and vascular surgery). Another potential adjunctive strategy that may be considered is the cell saver. Although multiple other strategies (e.g., ureteral stents, preoperative placement of pelvic artery catheters for potential postpartum embolization and/or balloon inflation) have been utilized, these strategies have not been shown in either prospective trials or retrospective studies to have clear benefits that outweigh risks [23,34–36].

The patient (and family members if available) should be counseled regarding risks, complications, and management. The possible need for hysterectomy as a life-saving procedure should be discussed with the patient. Plans for future reproduction should be discussed and weighted against the risk of retaining the uterus. Other preventive or therapeutic interventions as described above and below should be discussed.

Table 28.4 Ultrasoundographic signs associated with placenta accreta

<table>
<thead>
<tr>
<th>First trimester</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Gestational sac that is located in the lower uterine segment</td>
<td></td>
</tr>
<tr>
<td>Multiple irregular vascular spaces noted within the placental bed</td>
<td></td>
</tr>
<tr>
<td>Implantation of the gestational sac imbedded into the uterine window at the site of the prior cesarean delivery (cesarean scar ectopic)</td>
<td></td>
</tr>
<tr>
<td>Second trimester</td>
<td></td>
</tr>
<tr>
<td>Multiple vascular lacunae within the placenta</td>
<td></td>
</tr>
<tr>
<td>Third trimester</td>
<td></td>
</tr>
<tr>
<td>Loss of the normal hypoechoic retroplacental zone</td>
<td></td>
</tr>
<tr>
<td>The presence of multiple vascular lacunae within placenta (Swiss cheese appearance)</td>
<td></td>
</tr>
<tr>
<td>Abnormalities of the uterine serosa–bladder interface (interruption of the line, thickening of the line, irregularity of the line, and increased vascularity)</td>
<td></td>
</tr>
<tr>
<td>Extension of the villi into the myometrium, serosa, or bladder</td>
<td></td>
</tr>
<tr>
<td>Retroplacental myometrial thickness of less than 1 mm</td>
<td></td>
</tr>
<tr>
<td>Turbulent blood flow through the lacunae on Doppler ultrasonography</td>
<td></td>
</tr>
<tr>
<td>Increased subplacental vascularity</td>
<td></td>
</tr>
<tr>
<td>Vessels bridging from the placenta to the uterine margin</td>
<td></td>
</tr>
<tr>
<td>Gaps in myometrial blood flow</td>
<td></td>
</tr>
</tbody>
</table>

Table 28.5 Preparations in Cases of Suspected Placenta Accreta

<table>
<thead>
<tr>
<th>Outpatient procedures</th>
<th>Inpatient procedures</th>
<th>Postoperative</th>
</tr>
</thead>
<tbody>
<tr>
<td>• MFM and Gyn Oncology consultations as outpatient</td>
<td>• Admit the day before</td>
<td>• SICU—If necessary, decide intraoperatively which level of care is appropriate</td>
</tr>
<tr>
<td>Interventional radiology (IR), urology, and anesthesia consultations as outpatient</td>
<td>Type and cross—10 U pRBC, 7 FFP, in room</td>
<td>Possible post op embolization with IR</td>
</tr>
</tbody>
</table>
| Prenatal care and ultrasounds with MFM. Specify: | Blood bank to be notified | |]
| • Placenta location | • Order the blood products before the case, not same day | | |
| • Relevant ultrasound findings | • Clear liquid diet/consider bowel preparation if concern for bowel involvement | | |
| • Obstetric history | • Placement of large bore IV | | |
| • No. of prior cesarean deliveries | • If IR involved: | | |
| • Other prior uterine surgery | • Fetal heart monitoring before IR | | |
| • Latest hemoglobin (within 4 weeks of surgery; aim for >10 mg/dL) | • Heparin 5000 U SQ | | |
| • Betamethasone at 33.5 weeks | • Preop for IR by 7 AM first case | | |
| • Third trimester MRI if posterior placenta or concern for percreta | • Heparin 5000 U SQ | | |
| • Cesarean hysterectomy in main OR | • Transfer directly from IR suite to main OR | | |
| • Regional anesthesia preferred before delivery of baby; then possibly general anesthesia as per gyn oncology and anesthesia preference | • Unasyn 3 g IV before ureteral stents | | |
| • Positioning: lithotomy in Allen stirrups | • Cell saver in room | | |
| • Sequential compression devices (SCDs) placed | • Surgical intensive care unit (SICU) bed on hold | | |
| • Urology—open-ended ureteral catheters | Intraoperative considerations | | |
| • Vertical skin incision | • Regional anesthesia preferred before delivery of baby; then possibly general anesthesia as per gyn oncology and anesthesia preference | | |
| • C-section, evaluate need for hysterectomy | • If only a small area of the placenta is adherent and a focal area of the placental bed is bleeding, sewing over this area with sutures can be considered, but usually these are in | | |

Delivery

Timing. The optimal GA for delivery of a woman with placenta accreta is uncertain. Many authors recommend a delivery prior to 39 weeks (e.g., 36–37 weeks) in order to avoid an unscheduled delivery and to optimize the capacity for preparation. Fetal maturity testing has been advocated by some and not by others, and one recent decision analysis suggests that it does not help to improve overall health outcomes. This analysis also suggests that in certain high-risk cases (e.g., intermittently bleeding placenta previa with suspected placenta percreta) even earlier delivery (e.g., 34–35 weeks) may be acceptable [37]. Both the Society for Maternal-Fetal Medicine (SMFM) and the American College of Obstetricians and Gynecologists (ACOG) recommend delivery around 34 0/7–35 6/7 weeks for placenta accreta [23,38]. The final timing for delivery will need to be individualized, and take into account the risks of prematurity to the infant and the risks of major morbidity to the mother.

The two most common approaches to management of suspected placenta accreta are hysterectomy without attempt at placental removal versus attempt at placental removal. While there are different potential approaches for managing placenta accreta after delivery of the baby by cesarean, many experts would recommend that if there is proven placenta accreta (e.g., placenta directly visualized in the serosa), highly suspected accreta by history and radiologic studies (e.g., multiple prior cesarean deliveries, placenta previa, and several ultrasonographic findings of placenta accreta), or if the mother plans no more childbearing (e.g., has requested tubal ligation), a reasonable course of action is to avoid disturbing the implantation of the placenta and proceed with a hysterectomy while the placenta remains attached [39]. In these controlled situations, maternal morbidity of gravid hysterectomy may be decreased, but fertility is lost.

During hysterectomy, the uterine serosa overlying the placenta should not be dissected, including trying to avoid bladder dissection. The blood supply to the placenta is not just from the uterine arteries but also from many collateral vessels. Care should be given to dissection of this extensive blood supply by attempting uterine artery dissection laterally, and then continuing down without incising the placenta. Total hysterectomy is generally necessary, as subtotal hysterectomy may leave behind part of the lower segment, where the placenta is abnormally attached and cause bleeding if a previa is present. If the diagnosis is possible but not certain, and the patient desires to make attempts at avoiding hysterectomy despite having been counseled regarding risks, it is not unreasonable to wait for signs of placental separation, although abnormal adherence or significant hemorrhage should be ascertained and acted upon promptly. If spontaneous placental delivery fails, the operator must decide if either manual placental removal in pieces or hysterectomy is the next intervention, based on several factors, including the degree of invasiveness and amount of bleeding.

If only a small area of the placenta is adherent and a focal area of the placental bed is bleeding, sewing over this area with sutures can be considered, but usually these are in

Abbreviations: FFP, fresh frozen plasma; IV, intravenously; IR, interventional radiology; MFM, maternal-fetal medicine; MRI, magnetic resonance imaging; OR, operating room; pRBC, packed red blood cells; SQ, subcutaneously.
the very low uterine segment and cervix and often continue to bleed despite suturing or uterotonics. Some have suggested ligation of pelvic vessels (such as the internal iliac artery) in the setting of significant hemorrhage, although this may not be beneficial, given the many collateral vessels, and may incur risks as well. Pelvic packing has been used in some cases as a measure to temporarily lessen bleeding and allow attainment of hemodynamic stability. Hysterectomy may be necessary if uterine bleeding cannot be controlled, hopefully before massive blood loss and cardiovascular instability. When bleeding is from the lower part of the uterus in the setting of an accreta and placenta implanted low in the uterus (e.g., a previa), total hysterectomy is desirable, as subtotal hysterectomy may not control bleeding. Gravid hysterectomy has been associated with an incidence of maternal mortality of up to 7%, with a 90% incidence of transfusion, 28% incidence of postoperative complications, and placenta implanted low in the uterus in the setting of an accreta and placenta implanted low in the uterus (e.g., a previa), total hysterectomy is desirable, as subtotal hysterectomy may not control bleeding. Gravid hysterectomy has been associated with an incidence of maternal mortality of up to 7%, with a 90% incidence of transfusion, 28% incidence of postoperative complications, and a 5% incidence of ureteral injuries or fistula formation [19,39].

In some cases, a woman who has not completed childbearing may strongly want to avoid hysterectomy. There are several case reports of expectant (also called “conservative”) or medical management in the setting of placenta accreta. In these circumstances, the placenta is left in situ and the cord ligated close to its origin, either with no therapy or with an adjunctive therapy such as methotrexate and/or arterial embolization. There is no proven efficacy for methotrexate, or even for embolization. Antibiotic prophylaxis has been suggested given the risk of infection, as have short-term uterotonics for postpartum hemorrhage prevention, but there are no trials of these interventions. Follow-up is also not based on good evidence, but serial ultrasounds (to monitor involution and decrease in placental vascularity) and quantitative human chorionic gonadotrophin (HCG) levels have been proposed. If HCG levels plateau, or uterine size or placental vascularity do not decrease by 72 hours, methotrexate has been given as 1 mg/kg on alternate days for a total of four to six doses, or according to HCG levels and ultrasonographic findings [40]. Women on methotrexate should be monitored with liver function tests (LFTs), platelet counts, and creatinine levels. However, conservative management, especially for placenta percreta, has been associated with significant morbidity, including infection, delayed hemorrhage, a 42% chance of major morbidity (DIC, major postpartum hemorrhage, pulmonary embolism, sepsis, fistula, AV malformation), and a 58% risk of later hysterectomy (40% of the times emergent) at least up to 9 months postpartum [41–45] and should be considered only when the woman has been fully apprised of the risks and when no active uterine bleeding is present. In a follow-up study of 96 women with successful conservative management, about 9% had Asherman’s syndrome, and 25% had subsequent pregnancies, with 29% recurrent accretas, and 62% third trimester deliveries [46].

Postpartum
Consider reservation of intensive care unit bed for postpartum care.

VASA PREVIA
Diagnosis/Definition
Fetal blood vessels, unprotected by the umbilical cord or placental tissue that run through the membranes and over the internal os. Two types have been described: Type I when there is a velamentous cord insertion, and Type II when the placenta contains a succenturate lobe or is multilobed and fetal vessels connecting the two placental lobes course over or near the internal os [47].

Symptoms
Asymptomatic unless membranes rupture, at which time vaginal bleeding may be noted.

Epidemiology/Incidence
Approximately 1/2500–1/5000 deliveries [1].

Risk Factors/Associations
Placenta in the lower uterine segment (e.g., low-lying, or previa earlier in pregnancy), velamentous cord insertion, succenturiate or bilobed placenta, and multiple gestations [48].

Complications
Perinatal morbidity (e.g., neonatal anemia) and mortality (up to 56%) due to acute fetal hemorrhage [49].

Management
Principles
Timing of bleeding with antenatally diagnosed vasa previa is variable and impossible to predict. Since the fetal vessels are not protected by placental tissue or Wharton jelly, compression may lead to reduced fetal blood flow and bradycardia, and rupture of membranes with subsequent vessel laceration may result in rapid fetal exsanguination. It’s paramount to make the diagnosis of vasa previa antepartum, and not intrapartum.

Workup
TVU with color Doppler is the standard for diagnosis of vasa previa (Figure 28.4). Women with risk factors that are judged to increase their risk of vasa previa (see above) should be screened with TVU for this condition. Vasa previa is diagnosed if a vessel is visualized over the cervix with color Doppler demonstrating a rhythm consistent with the fetal heart rate. The American Institute of Ultrasound in Medicine (AIUM) and ACOG recommend that the placental cord insertion site be documented whenever technically possible [50]. A recent systematic review showed a high accuracy of TVU color Doppler (median detection rate of 93% and specificity of 99%), but also noted that the quality of the available studies was relatively poor [51]. Thus, not all vasa previa will be detected prenatally, even by adequate examinations and experienced operators using color Doppler. Women whose vasa previa has been diagnosed prenatally have been reported to have lower perinatal mortality than those with vasa previas that are undiagnosed (3% vs. 56%) [49]. The Apt test may be used to distinguish between fetal and maternal sources of vaginal bleeding, although this test may be of little use in many clinical situations with bleeding from a vasa previa, as bleeding can lead to rapid deterioration of the fetal status and require urgent delivery before an Apt test can be completed.

Therapy
Level 1 data to guide the management of antenatally diagnosed vasa previa are currently lacking. While some experts suggest that hospitalization at some time after viability may be reasonable, this strategy is unsupported by any adequately powered trials. Many admit women with proven vasa previa around 28–32 weeks. It is reasonable to consider administration of antenatal corticosteroids at 28–33 weeks of gestation,
if indications do not develop prior, in case of the need for urgent preterm delivery [52]. Women with vasa previa should be delivered by cesarean at a center capable of providing immediate neonatal blood transfusion if needed [53]. Timing of delivery is controversial, with suggestions for cesarean delivery anywhere between 34 0/7 and 35 6/7 weeks, although earlier when preterm premature rupture of membranes (PPROM), PTL, or significant bleeding occurs [1].

REFERENCES


Abruptio placentae

John F. Visintine

KEY POINTS

- Approximately 0.2%–1% of all pregnancies are complicated by placental abruption.
- Nearly 50% of women with placental abruption have no identifiable risk factors. Risk factors include abruption in a prior pregnancy, maternal hypertensive disorders, smoking, cocaine, polyhydramnios, multiple gestation, preterm and term premature rupture of membrane, chorioamnionitis, elevated maternal serum alpha-fetoprotein (MS-AFP), pregnancy-associated plasma protein-A (PAPP-A) ≤5th percentile, leiomyoma, subchorionic hematoma, vaginal bleeding <20 weeks, previous cesarean delivery, abdominal trauma, subclinical hypothyroidism, antithyroglobulin and antithyroxperoxidase antibodies. The association with thrombophilias has for the most part not been confirmed by prospective studies.
- Complications include antepartum and postpartum hemorrhage, disseminated intravascular coagulopathy (DIC), and acute renal failure, as well as growth restriction, preterm delivery, fetal hypoxia and/or exsanguination, and perinatal mortality.
- The diagnosis of placental abruption is primarily clinical. History, physical examination, laboratory and ultrasonographic studies guide management. Ultrasound is primarily useful in ruling out other causes of third-trimester bleeding.
- Placental pathology has been shown to confirm the presence of abruption in 25% of those with an acute clinical abruption, and 60% of those with a chronic clinical abruption.
- There are no trials to assess any intervention for prevention of abruption or its complications.
- Prompt delivery is indicated if the pregnancy is late-preterm or term. However, if less than 34 weeks, expectant management for mild (grade 1) abruptions may allow time for glucocorticoid administration. Maternal or fetal compromise may necessitate delivery. A decision-to-delivery interval of 20 minutes or less is associated with a substantial reduction of neonatal morbidity and mortality in placental abruption with non reassuring fetal heart testing.
- Mode of delivery is dependent primarily on the condition of the mother and fetus.
  - In most cases, for mild abruption (grade 1, no evidence of maternal or fetal compromise), vaginal delivery is indicated.
  - For moderate abruption (grade 2, evidence of fetal nonreassuring testing), rapid delivery typically by cesarean is indicated.
  - For severe abruption (grade 3, fetal demise, often with DIC), vaginal delivery is indicated.

DEFINITION/DIAGNOSIS

Placental abruption (also called abruptio placentae) is defined as a premature separation of a normally implanted placenta. The diagnosis is a clinical diagnosis of exclusion, based usually on vaginal bleeding in the second or third trimester unexplained by other etiologies (see also section “Etiology/Basic Pathophysiology”).

SIGNS AND SYMPTOMS

Signs and symptoms are shown in Table 29.1 [1,2]. About 10%–31% of abruptions present with only concealed (occult) bleeding. Occasionally the presenting sign is fetal death.

EPIDEMIOLOGY/INCIDENCE

- Approximately 0.2%–1% of all pregnancies are complicated by placental abruption [3–5]. The incidence in the United States has increased, especially in the African-American population, the ethnic group at highest risk, in particular for severe (or grade 3) abruption [3].
- About 60% of abruptions occur preterm, and 50% occur prior to labor.
- The incidence of abruption is possibly highest at 24–26 weeks (up to 9 per 100 births) [6].
- There is over a 5% recurrence risk in a subsequent pregnancy for women with a history of an abruption [4].

GENETICS

A genome wide association study identified several candidate genes associated with placental abruption. The strongest association was with the FLI-1 gene (a megakaryocyte-specific transcription factor). Genes involved in lipid metabolism, cell signaling, mitochondrial biosynthesis, and oxidative phosphorylation are also associated with the risk of placental abruption [7]. The association with thrombophilias has not been confirmed in prospective studies. See section “Risk Factors/Associations” and Chapter 27 in Maternal-Fetal Evidence Based Guidelines.

ETIOLOGY/BASIC PATHOPHYSIOLOGY

The etiology is not completely understood but appears to occur as the result of two mechanisms: mechanical separation or as the end result of defective deep placentation [8]. Placental separation occurring in association with mechanical trauma or rapid decompression of a distended uterus is believed to occur due to shearing forces resulting from a change in surface area of a relatively elastic uterine wall in relation to an inelastic placenta. Blunt trauma to the uterus resulting in abruption or rupture of membranes with rapid decompression of an over distended uterus resulting in abruption are examples of this mechanism.
Table 29.1 Clinical Findings in Women with Placental Abruption

<table>
<thead>
<tr>
<th>Clinical Findings</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vaginal bleeding</td>
<td>67–78</td>
</tr>
<tr>
<td>Uterine tenderness or abdominal/back pain</td>
<td>52–66</td>
</tr>
<tr>
<td>Nonreassuring fetal testing/Category II or III</td>
<td>60–78</td>
</tr>
<tr>
<td>Uterine contractions &gt;5/10 minutes</td>
<td>17</td>
</tr>
</tbody>
</table>


Evidence in support of a defective deep placentation mechanism comes from placental bed biopsies from cases of placental abruption, which show an absence of physiologic trophoblastic invasion, dilated vessels, and recent thrombosis of spiral arteries [9]. These changes are seen not only in abruption but also in preeclampsia, intrauterine growth restriction (IUGR), and preterm premature rupture of membranes (PPROM), suggesting that placental abruption represents one manifestation of a long-standing pregnancy disorder [8]. Moreover, abnormalities in circulating angiogenic factors have been observed in women who subsequently developed placental abruption. In a nested case–control study, serum levels of the proangiogenic factor placental growth factor (PIGF) were lower, and levels of the potent inhibitor of PIGF, soluble fms-like tyrosine kinase 1 (sFlt-1), were increased [10].

Gross pathology findings associated with placental abruption include adherent retrolenticular clot with depression or disruption of the underlying placental tissue. Microscopic changes associated with abruption resulting in perinatal mortality include thrombosed arteries and necrosis of the decidua basalis, and large recent infarcts and stromal fibrosis in the terminal villi of the placental parenchyma [11]. Severe cases of abruption can result in hemorrhagic infiltration between myometrial fibers that extends to the serosal surface giving the uterus a dark purple discoloration, also known as Couvelaire uterus in honor of the French obstetrician who first described this pathologic finding in the early twentieth century [12].

The pathologic diagnosis of a placental abruption often does not correlate with the clinical diagnosis. In a large series of placental pathologic evaluations, a placental abruption defined as an adherent clot with disruption of the underlying tissue was demonstrated in 3.8% of specimens, which is higher than the generally accepted incidence based on a clinical diagnosis (0.2%–1%) [13]. Most of those “abruptions” called by pathologists do not have clinical diagnosis of abruption, and many clinical abruptions have no pathologic evidence of abruption. We have structured the chapter around a clinical definition of abruption. A retrospective review of abruption cases diagnosed on clinical grounds or by pathologic criteria were analyzed based on risk factors, acute (cocaine use, trauma <12 hours from delivery) or chronic (hypertension, preeclampsia, acute chorioamnionitis, trauma occurring >12 hours prior to delivery). Placental pathology confirmed the presence of abruption in only 25% of those with acute risk factors for abruption and 60% of those with a chronic risk factors for abruption [14].

**CLASSIFICATION**

A uniformly accepted classification system for placental abruption does not exist. The clinical classification system originally published by Page in 1954 has been used by some authors as a means of grouping placental abruptions in those that can be potentially managed conservatively (grade 1) and those that require more aggressive management (grades 2 and 3) [15] (Table 29.2). A more recent classification defines severe abruption as at least one maternal (disseminated intravascular coagulation, hypovolemic shock, blood transfusion, hysterectomy, renal failure, or in-hospital death), fetal (nonreassuring fetal status, intrauterine growth restriction, or fetal death), or neonatal (fetal growth restriction, preterm birth) complication. Using these criteria, two-thirds of placental abruptions are defined as severe (overall abruption rate 9.6/1000, severe abruption 6.5/1000) [5]. It should be noted that the definition of placental abruption in this study does not include abruptions only identified through pathologic evaluation of the placenta (grade 0).

**RISK FACTORS/ASSOCIATIONS**

Nearly 50% of women with placental abruption have no identifiable risk factors (Table 29.3) [16].

- **History of an abruption in a prior pregnancy:** The risk of recurrence is about 5%–17% [17]. After two abruptions, the risk of recurrence is about 25%.
- **Maternal hypertensive disorders:** Associated with up to almost 50% of grade 3 abruption cases. In particular, chronic hypertension (incidence 1.5%–2.5%, odds ratio [OR] 2.8), superimposed preeclampsia (about 3%), and severe preeclampsia (OR 4.1) are associated with placental abruption [18,19].
- **Abdominal trauma** is a recognized cause of placental abruption but is responsible for only 1% of cases [20]. See also Chapter 39 in Maternal-Fetal Evidence Based Guidelines.
- **Smoking:** 90% increase in abruption in women who smoke compared with controls. Smoking is responsible for 15%–25% of episodes of abruption [18].
- **Cocaine:** 1.9% rate of abruption [21].
- **Previous cesarean delivery** is associated with an increased risk of abruption in subsequent pregnancies (OR 1.3) [22].
- **Polyhydramnios** has been associated with placental abruptions in patients >37 weeks’ gestation [23].
- **Multiple gestation:** 1.2% risk of abruption in twins, 1.5% in triplets [24].
- **PPROM:** OR 3.5 [25]. Evidence of old decidual hemorrhage can be found in nearly 40% of placentas from patients with PPROM [26].
- **Chorioamnionitis:** OR 9 [25]. Neutrophil infiltration of the fetal membranes and cervix is seen in premature rupture of membranes (PPROM), and chorioamnionitis is associated with placental abruption [27].
- **Leiomyomas** detected at second-trimester ultrasound are associated with a small increase in the risk of abruption (OR 2.1) [28].
- **Ultrasound-detected subchorionic hemorrhage** before 22 weeks of gestation results in an increased risk of placental abruption (OR 2.6) [28].
- **Vaginal bleeding <20 weeks** is associated with an increased risk of abruption (RR 1.6) [29].
- **Unexplained elevated MS-AFP** in the second trimester: OR 6–10 for placental abruption [30,31].
- **PAPP-A ≤5th percentile** in the first trimester is associated with placental abruption (OR, 1.9) [32].
- **Inherited thrombophilias** have been associated with abruption in case–control studies [33,34]. However, most prospective studies have shown no increased risk of abruption [35–37]. A retrospective cohort study
Abruption Placenta

Table 29.2  Classification of Placental Abruption

<table>
<thead>
<tr>
<th>Grade</th>
<th>Clinically Evident Bleeding</th>
<th>Uterine Tenderness</th>
<th>Maternal Hypotension</th>
<th>Maternal Coagulopathy</th>
<th>Fetal Distress</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>1</td>
<td>Yes</td>
<td>Yes or no</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>2</td>
<td>Yes or no</td>
<td>Yes</td>
<td>No</td>
<td>Rare</td>
<td>Yes</td>
</tr>
<tr>
<td>3</td>
<td>Yes or no</td>
<td>Yes</td>
<td>Yes</td>
<td>Often</td>
<td>Death</td>
</tr>
</tbody>
</table>


*Grade 0: Diagnosis based on examination of the placenta.

Table 29.3  Risk Factors for Placental Abruption

<table>
<thead>
<tr>
<th>Risk Factor</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prior abruption</td>
<td>Chronic hypertension, Severe preeclampsia, Smoking, Cocaine, Chorioamnionitis, Unexplained elevated MS-AFP, PAPP-A levels ≤5th percentile, (P)PROM, Subchorionic hemorrhage, Leiomyoma, Vaginal bleeding &lt;20 weeks, Polyhydramnios, Multiple gestation, Trauma, Subclinical hypothyroidism, Thyroperoxidase or thyroglobulin antibodies</td>
</tr>
</tbody>
</table>

**Abbreviations:** MS-AFP, maternal serum alpha-fetoprotein, PAPP-A, pregnancy-associated plasma protein-A; (P)PROM, preterm premature rupture of membrane.

Comparing thromboprophylaxis with heparin for women with an inherited thrombophilia demonstrated no difference in rates of abruption or other adverse outcomes regardless of treatment [38]. But one large prospective cohort study did demonstrate an increased risk of abruption for women with the prothrombin gene mutation 20210A (OR 12), but not for factor V Leiden mutation, MTHFR C677T or A1298C mutations, or the thrombomodulin gene mutation [39].

- **Advanced maternal age:** It is unclear if there is an association with abruption, as the data are conflicting [40,41].
- **Thyroperoxidase antibodies** have been associated with an increased risk of placental abruption in two large cohort studies (OR 1.5–3.4) [42,43]. An association with abruption was also seen with **thyroglobulin antibodies** (OR 1.4–1.7) [43]. The association with abruption was confirmed in a cohort of low risk women with **subclinical hypothyroidism** [44].

**Complications**

- **Maternal**
  - **Serious maternal complications** (pulmonary edema, acute respiratory failure, acute heart failure, acute myocardial infarction, cardiomyopathy, puerperal cerebrovascular disorder, coma, and amniotic fluid embolism) occur at a higher ratio for women with **severe abruption** (141.7/10,000) compared with those with a **mild abruption** (33.3/10,000) or nonabruption (15.4/10,000 births) [5].
  - **Antepartum hemorrhage** remains a leading cause of maternal mortality. For pregnancies ending in stillbirth, hemorrhage related to placental abruption is the leading cause of maternal mortality [45].
  - **DIC** was first reported to occur in association with placental abruption by De Lee in 1901 [46]. The development of DIC is thought to be due to a release of thromboplastins, as well as consumption of coagulation factors secondary to an enlarging hematoma. Nearly 30% of patients who present with a severe (grade 3) abruption develop DIC.
  - **Acute renal failure** is a potential maternal complication associated with abruption. Fortunately the incidence of acute renal failure appears to be decreasing, possibly due to improved medical management.
  - **Postpartum hemorrhage secondary to uterine atony** is associated with abruption, as is postpartum anemia and infection.

- **Perinatal**
  - **Perinatal mortality** (both fetal and neonatal deaths) varies from 4 to 12/1000 [47]. This high perinatal mortality with abruption is attributable, in part, to its association with **preterm delivery**.
  - Of the excess perinatal deaths, about 55% can be attributed to prematurity. The other is associated with **fetal hypoxia** and/or **exsanguination**, or **growth restriction**.
  - Among abruption cases from a large multicenter case-control study, 60% were found to be small for gestational age; the authors concluded that this association was primarily due to preterm birth [48].
  - Of children with spastic quadriplegic or dyskinetic cerebral palsy, 5.7% are associated with placental abruption [49].

**Management**

Unfortunately there are **no trials to assess any intervention for management of abruption or its complications** [50]. Recommendations for management are primarily based on expert opinion or at best retrospective case-control studies (Figure 29.1).

**Prevention**

- A reduction in the risk of placental abruption was demonstrated with **prenatal vitamin C and E supplementation in smokers** (relative risk (RR) 0.09; 95% confidence interval [CI] 0.00–0.87) but not in nonsmokers (RR 0.92; 95% CI 0.52–1.62) [51].
- The use of **magnesium sulfate for women with preeclampsia** was shown to reduce the risk of placental abruption (RR 0.64; 95% CI 0.5–0.83) [52].
- **Smoking cessation counseling, avoidance of cocaine, and, if possible, avoidance of other risk factors may prevent placental abruption.**
Preconception Counseling

Women with risk factors should be counseled regarding the risk factors (Table 29.3) and complications of placental abruption, as well as interventions for its prevention.

Workup

- The diagnosis of an abruption is made clinically (vaginal bleeding, abdominal pain/uterine tenderness, nonreassuring fetal testing, or uterine tachysystole). Pathologic evaluation of the placenta may confirm the diagnosis in some cases (25% of acute and 60% of chronic abruptions), but the absence of pathologic findings does not exclude an abruption [14]. Placental abruptions may be occult. At times presenting as preterm labor, they may go undiagnosed until after delivery with examination of the placenta. If a pathologic diagnosis (rather than clinical diagnosis) of placental abruption is used the incidence of placental abruption is 3.8% [13] compared with around 1% in epidemiologic studies that employ a clinical diagnosis.

- History and physical examination, as well as appropriate laboratory and ultrasonographic studies, guide management. Routine assessment should be conducted including vital signs, oxygenation status, and urine output. Laboratory assessment may include a hematocrit, platelet count, coagulation studies (prothrombin time [PT], partial thromboplastin time [PTT], and fibrinogen), blood type and screen, or cross-match, serum creatinine, and a drug screen. Other causes of third-trimester bleeding must be excluded. The differential diagnosis includes placenta previa, vasa previa, cervical lesions (for example, malignancy) and vaginal lesions.

- An ultrasound examination is useful primarily in the exclusion of placenta previa or vasa previa. The sensitivity and specificity of ultrasound in the diagnosis of placental abruption were 24% and 96%, respectively [53]. Of note, a placental abruption was defined in this study by clinical signs or symptoms or by placental examination. So while ultrasound is very helpful in ruling out other causes of third-trimester bleeding, it lacks the sensitivity needed to reliably detect placental abruption [53]. The echogenicity of the collection of blood of an abruption depends on the time the ultrasound was performed relative to the onset of symptoms [54]. Acute hemorrhage is hyperechoic to isoechoic compared with the placenta. Resolving hematomas become hypoechoic within 1 week and sonolucent within 2 weeks.

- Computed tomography (CT) has been used to identify abruptions in trauma patients. A retrospective study of pregnant trauma patients who were evaluated with CT found a high sensitivity (86%) and specificity (98%) in the detection of placental abruption [55]. In a retrospective review of pregnant trauma patients, as placental enhancement on CT imaging decreased the presence of clinical findings of placental abruption were found to increase. The authors found that clinical signs and/or symptoms of abruption were significantly more likely with placental enhancement on CT imaging <50%, and that the likelihood of delivery for abruption increased as the placental enhancement decreased to less than 25% [56].

- Magnetic resonance (MR) imaging has also been used to identify placental abruption. In a prospective series of 60 consecutive patients with clinical suspicion of abruption ultrasound imaging and MR imaging were compared. Placental abruption (defined as adherent clot or fibrin by placental examination at delivery) was found in 19 patients. Abruption was identified in 10 of the 19 patients (52%) with ultrasound and in all 19 (100%) with MR imaging (p = .002) [57]. This study underscores the relative ability of imaging modalities to identify retroplacental collections, but the presence of absence of adherent placental clot does not always correlate with a clinical abruption.
• A vaginal speculum examination should be performed to rule out cervical or vaginal sources of bleeding.

• Laboratory findings: A prolonged PT, prolonged PTT, hypofibrinogenemia, and thrombocytopenia can occur in women with abruption for whom DIC develops. D-dimer, a fibrin degradation product, has been evaluated as a diagnostic tool for abruption in two small studies with conflicting results [58,59]. At present the data do not support its routine use. The Kleihauer-Betke (KB) test for fetal red cells in the maternal circulation appears to be a low yield test for women with abruption. In a recent retrospective study reported no positive KB tests out of 27 cases of abruption confirmed by pathology [61].

General Principles

Once the diagnosis of placental abruption has been made, attention should be focused on ensuring maternal and fetal well-being. Maternal status should be addressed with attention paid to signs or symptoms of hemorrhage, hypovolemic shock, and DIC. The frequency of repeated evaluations is dependent primarily on the acuity and severity of the abruption. Preparations should be made in anticipation of potential maternal complications. This should include intravenous access; two large-bore peripheral lines will allow for rapid fluid or blood component replacement. The availability of blood or blood components may be life saving; therefore, close cooperation with blood banking services is essential.

Fetal status is typically assessed with continuous electronic fetal monitoring, at least in the acute setting. For women with a chronic abruption, once clinically stable, intermittent monitoring may be a consideration.

Timing of Delivery

Late-Preterm or Term Placental Abruption: ≥34 Weeks

Prompt delivery is indicated if the pregnancy is ≥34 weeks [62–64]. Fortunately rapid labor often ensues as a result of the abruption (Figure 29.1).

Preterm Abruption: <34 Weeks

Maternal or fetal compromise necessitates delivery. In selected patients with mild (grade 1) abruption, with no evidence of maternal or fetal compromise, expectant management may allow time for glucocorticoid administration and appears to be a safe option [1,15,64]. Tocolysis should in general not be used. Antepartum testing should be frequent, and at least initially, continuous fetal monitoring is indicated.

Mode of Delivery

Deciding on the method of delivery is dependent primarily on the condition of the mother and fetus.

Mild Abruption (Grade 1): No Evidence of Maternal or Fetal Compromise

With close monitoring of the mother and fetus, vaginal delivery may be accomplished. In studies of women with mild (or grade 1) abruptions, those who delivered vaginally had a similar perinatal mortality rates compared with those with a cesarean delivery [62].

Moderate Abruption (Grade 2): Evidence of Fetal Compromise

Rapid delivery: typically cesarean delivery is indicated. In a study of placental abruption complicated by fetal bradycardia, a decision-to-delivery interval of ≤20 minutes was associated with substantially reduced neonatal morbidity and mortality [65].

Severe Abruption (Grade 3): Fetal Death, Often with DIC

Vaginal delivery is preferred in this group as a cesarean delivery may exacerbate maternal hemorrhage. If present, DIC will typically resolve with evacuation of the uterus, with possible improvement in clotting parameters even prior to delivery [66,67].

Anesthesia

No specific suggestions. Anesthesia support is particularly important with DIC, hemorrhagic shock, and massive transfusion cases.

Postpartum

Attention should be paid to hemodynamic state and possible late hemorrhage from uterine atony after abruption.

REFERENCES


Postpartum care includes breast-feeding, contraception, postpartum endometritis, postpartum wound complications, and postpartum mood and anxiety disorders.

**BREAST-FEEDING KEY POINTS**

- **In normal reproductive physiology, lactation follows pregnancy.** For the infant, use of artificial breast milk substitutes is associated with increased acute and chronic disease risk. For mothers, never or curtailed (e.g., short-term) breast-feeding is associated with increased risks of malignancy and metabolic disease.

- **All major medical organizations recommend 6 months of exclusive breastfeeding, with continued breast-feeding through 1–2 years, or longer as mutually desired by mother and infant.**

- **Infant demand drives maternal milk supply.** To ensure adequate milk production, encourage frequent, on-demand feeding in response to infant cues, continuing until the infant is satisfied.

- **Pre- and postnatal lay and/or professional support and evidence-based physician training** improve breast-feeding duration and exclusivity.

- **Maternity care practices affect breastfeeding initiation and duration.** Implementation of the United Nations International Children’s Emergency Fund (UNICEF)/World Health Organization (WHO) Baby Friendly Hospital Initiative (BHI) improves breast-feeding duration, intensity, and infant health outcomes.

- **Prenatal treatment of inverted nipples does not increase breast-feeding success.**

- **Distribution of formula company marketing materials and samples adversely affects breast-feeding outcomes.** Formula company materials should not be distributed in healthcare facilities.

- **At birth, early skin-to-skin contact increases breast-feeding duration. Healthy infants should be placed skin-to-skin on the mother’s chest** and remain there, undisturbed, until the first feeding is accomplished.

- **Treatment for mastitis begins with frequent removal of milk, hydration, and analgesia. Healthy term infants can continue to feed on the affected side.** Antibiotics are used if conservative management is ineffective or the patient is acutely ill.

- **There is limited evidence that galactagogues increase milk production in placebo-controlled trials.** Optimal breast-feeding education can increase milk supply among women with low production.

- **Most medications are compatible with breast-feeding, or a safe alternative medication exists.** The physiology of the placenta differs from the breast, and providers should not extrapolate drug safety information from pregnancy to lactation.

- **Breast-feeding is contraindicated in the setting of an infant with classic galactosemia, active maternal use of illicit drugs, maternal medications that are contraindicated in breast-feeding, or maternal infection with human T-cell lymphotropic virus type I or II (e.g., HTLV).** Recommendations for breast-feeding in the setting of maternal human immunodeficiency virus (HIV) or tuberculosis vary by region. Women with HIV in the United States are advised not to breast-feed because of the risk of maternal–infant transmission and the availability of safe artificial feeding. Breast-feeding is temporarily contraindicated during perinatal maternal varicella infection or from a breast with an active herpetic lesion.

### Definition

Breast-feeding is the physiologic form of infant nutrition. Breast-feeding is defined as the infant receiving any amount of human milk. **Exclusive breast-feeding** is defined as receiving human milk alone for nutrition. **Predominant breast-feeding** is defined as receiving human milk and nonformula supplements, and **mixed feeding** is defined as receiving both human milk and infant formula.

### Breast-Feeding Physiology

Secretory differentiation occurs during pregnancy, as placental hormones stimulate development of mammary alveoli. After birth, secretory activation occurs, as progesterone levels fall and milk production increases from 50 mL/day to approximately 500 mL/day in the first 2–3 days after birth. **Positive feedback via infant stimulation** of the breast causes secretion of prolactin from the anterior pituitary and oxytocin from the posterior pituitary. **Prolactin** stimulates continuous milk synthesis, whereas **oxytocin** stimulates intermittent milk secretion, when myoepithelial cells contract to transfer milk from the alveoli to the areola. A milk is not removed regularly, negative feedback downregulates prolactin receptors and reduces production [1]. Frequent, on-demand feeding is essential to establish and maintain lactation.

### Health Effects of Infant Feeding

Infants who are artificially fed face higher risks of infectious morbidity and chronic disease than infants who are breast-fed. An Agency for Healthcare Research and Quality (AHRQ) meta-analysis of outcomes in developed countries found that **artificially fed infants** faced a 2.0-fold risk of otitis media, a 3.6-fold risk of pneumonia, and a 2.8-fold risk of gastrointestinal infection compared with infants who were exclusively breast-fed [2]. Artificial feeding is also associated with a 1.8- to 3.7-fold [3] risk of sudden infant death syndrome (SIDS),...
a 1.1- to 1.4-fold [4] risk of child obesity, and a 1.5-fold risk of type 2 diabetes [4] compared with breast-feeding. Artificial or mixed feeding is also associated with higher risks of asthma, odds ratio (OR) 1.2-1.3 [5], compared with exclusive breast-feeding. Among preterm infants, artificial feeding is associated with a 5% absolute increased risk of necrotizing enterocolitis compared with being fed mother’s milk [2].

For mothers, artificial feeding is similarly associated with increased health risks. Compared with women who have breast-fed, parous women who have never breast-fed or weaned early face higher rates of breast cancer [6,7], particularly triple-negative breast cancer [8–10], ovarian cancer [11], retained gestational weight gain [12,13], type 2 diabetes [14–17], hyperlipidemia [18], metabolic syndrome [19,20], hypertension [16,21,22], and myocardial infarction [16,23].

Optimal Duration of Exclusive Breast-Feeding
Six months of exclusive breast-feeding, compared with 3–4 months of exclusive breast-feeding with mixed feeding through 6 months, reduces infant risk of gastrointestinal and respiratory tract infections without any adverse effect on infant growth [24]. For mothers, longer exclusive breast-feeding was associated with delayed resumption of menses and, in one small study, with greater maternal weight loss [25].

Infant Feeding Recommendations
All major medical organizations [26–29] recommend exclusive breast-feeding for the first 6 months of life. The WHO recommends continued breast-feeding up to 2 years of age or beyond [29]. The American Academy of Pediatrics (AAP) recommends continued breast-feeding for at least the first year of life, continuing for as long as mutually desired by mother and child [30].

Promoting Breast-Feeding
Antenatal interventions can increase breast-feeding initiation and duration (Table 30.1) [31]. Breast-feeding education improves initiation rates among low-income U.S. populations. In subgroup analyses, one-on-one, needs-based, informal education sessions were more effective (relative risk [RR] 2.40, 95% confidence interval [CI] 1.57–3.67) than generic, formal antenatal sessions (RR 1.26, 95% CI 1.00–1.60) [32]. Both lay and professional supports for mothers after birth improve breast-feeding duration and exclusivity, with the strongest effects found with proactive, face-to-face, lay support involving four to eight contacts [33]. An USPSTF review similarly found that interventions to promote and support breast-feeding increase initiation, duration, and exclusivity of breast-feeding (grade B).

Interventions that include both prenatal and postnatal components may be most effective [34]. Proactive, integrated pre- and postnatal support for breast-feeding mothers improves breast-feeding duration and intensity.

Table 30.1 Successful Breast-Feeding Recommendations

<table>
<thead>
<tr>
<th>Recommendations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Offer additional breast-feeding support to women who have had a narcotic/general anesthetic, a caesarean, or delayed contact with their baby.</td>
</tr>
<tr>
<td>Ensure breast pumps are available for women who have been separated from their babies, and give instruction on how to use them.</td>
</tr>
<tr>
<td>Encourage unrestricted breast-feeding frequency and duration.</td>
</tr>
<tr>
<td>Reassure women about breast milk supply and help them gain confidence.</td>
</tr>
<tr>
<td>Advise women that babies will stop feeding when satisfied.</td>
</tr>
<tr>
<td>Advise women of the signs that a baby is successfully feeding:</td>
</tr>
<tr>
<td>• Swallowing is audible and visible.</td>
</tr>
<tr>
<td>• There is a sustained rhythmic suck.</td>
</tr>
<tr>
<td>• The arms and hands are relaxed.</td>
</tr>
<tr>
<td>• The mouth is moist.</td>
</tr>
<tr>
<td>• Regular soaked/heavy nipples.</td>
</tr>
</tbody>
</table>

Reassure women that they may feel the following:
• Brief discomfort at the start of feeds in the first few days; this is not uncommon but should not persist.
• Softening of their breast during the feed.
• No compression of the nipple at the end of the feed.
• Relaxed and sleepy.

Attachment and positioning
Advise women of the following signs of good attachment and positioning:
• The baby’s mouth is wide open.
• There is less areola visible underneath the chin than above the nipple.
• The baby’s chin is touching the breast, the lower lip is rolled down, and the nose is free.
• There is no pain.

If the baby is not attaching effectively, advise teasing the baby’s lips with the nipple to open the mouth.


improved breast-feeding outcomes [40]. Healthcare providers who work with mother and infants should receive comprehensive breast-feeding training.

The Baby Friendly Initiative
Maternity care affects breast-feeding outcomes. The Baby Friendly Initiative (BFI) is WHO-developed set of maternity care recommendations designed to increase initiation and duration of breast-feeding. The BFI requires maternity centers to implement The Ten Steps to Successful Breast-feeding (Table 30.2) [41]. Each country is responsible for establishing processes and procedures for maternity center designation as baby friendly [41]. In a cluster-randomized trial in Belarus, BFI implementation improved breast-feeding exclusivity at 3 months (43.3% vs. 6.4%, p < .001) and 6 months (79% vs. 0.6%, p = .01) and increased any breast-feeding rates at 12 months (19.7% vs. 11.4%). Infants born in intervention hospitals had lower rates of gastrointestinal illness and atopic eczema in the first year of life [42]. Observational studies in the United States have found a dose–response association between implementation of BFI steps and maternal achievement of breast-feeding goals [43,44]. Cohort studies suggest that the Ten Steps are particularly effective among women with lower education, suggesting that BFI maternity care can reduce socioeconomic disparities in breast-feeding [45].

The National Institute for Health and Clinical Excellence (NICE) postnatal care guideline recommends that all healthcare

Training of Providers
Breast-feeding education and knowledge among healthcare providers is inconsistent [35–37]. A meta-analysis found that training breast-feeding support personnel using the WHO/UNICEF breast-feeding training course [38] increases duration of exclusive breast-feeding. In a cluster-randomized trial, implementation of a breast-feeding training curriculum for residents improved provider knowledge, practice patterns, and confidence, and increased exclusive breast-feeding rates at 6 months (OR 4.1) compared with control sites [39]. Interventions for hospital staff are similarly associated with
Breast-Feeding in Clinical Care

Antenatal Breast Examination
Clinical guidelines recommend evaluation of breast anatomy as part of prenatal care [27, 46]. However, there are no randomized controlled trials (RCTs) regarding the effects of an antenatal breast examination on breast-feeding outcomes or maternal satisfaction [47]. There is no evidence that antenatal manipulation for inverted nipples improves breast-feeding outcomes [46, 49].

Avoiding Commercial Infant Feeding Materials
In randomized trials, provision of formula company materials and gift packs during prenatal care [50] or during the maternity hospitalization [51–53] reduces exclusive breast-feeding. In a time series design study, changing from formula marketing discharge packs to noncommercial packs was associated with higher breast-feeding rates, but more than one-third of women in the noncommercial cohort received formula samples, demonstrating that policy implementation is challenging [54]. Formula company marketing materials should not be distributed in the healthcare setting.

Skin-to-Skin Contact at Birth
Early skin-to-skin contact improves breast-feeding outcomes. Skin-to-skin contact is defined as placing the naked infant prone on the mother's bare chest, with the infant's back covered with a warm blanket, ideally beginning immediately after birth. In a Cochrane meta-analysis, early skin-to-skin contact increased breast-feeding at 1–4 months postbirth (RR 1.27), and increased duration of total breast-feeding by about 6 weeks [55]. Based on these data, the NICE [56], AAP [30], and the American College of Obstetricians and Gynecologists (ACOG) [27] recommend that all healthy infants should be placed skin-to-skin at birth. Infants should remain with their mothers for at least 1 hour, and providers should encourage mothers to recognize when the infant is ready to feed and offer help if needed [41]. Routine procedures, such as weighing, bathing, and vitamin K and eye prophylaxis, should be avoided during the first hour, or performed on the mother's chest. Skin-to-skin care during procedures, such as a heel lance, appears to reduce infant pain [57]. Skin-to-skin is also feasible following cesarean section [58].

Anticipatory Guidance to Support Normal Physiology
Human milk production is driven by infant demand. Therefore, frequent feeding, on demand in response to infant feeding cues, continuing until the infant is satisfied, is a cornerstone of breast-feeding success [59], although a recent meta-analysis did not identify trials comparing baby-led feeding with scheduled breast-feeding [60]. NICE clinical guidelines have outlined anticipatory guidance for successful breast-feeding (Table 30.1). A small pilot study found that early, limited supplementation with hydrolyzed formula for infants with ≥5% weight loss increased exclusive breast-feeding rates at 1 week and 3 months postpartum [61]; several trials to test whether these findings can be replicated are underway (NCT02313181 and NCT02221167).

Management of Breast-Feeding Complications

Engorgement
Breast engorgement typically occurs between 2 and 5 days postpartum. A meta-analysis found insufficient evidence to support any single treatment strategy for engorgement [62]. Cold packs and cabbage leaves may be helpful, and acupuncture was associated with some improvement in symptoms. NICE guidelines recommend breast massage, continued breast-feeding, and analgesia for symptom relief [31].

Mastitis
Mastitis is defined as acute inflammation of connective tissue within the breast, which may or may not be accompanied by infection [63]. Interventions to prevent mastitis have not proven effective [64]. A Cochrane meta-analysis (two RCTs, n = 125 patients) concluded that there is insufficient evidence to support treatment with antibiotics [65]. An RCT suggests that lactobacilli strains isolated from breast milk may be effective for treatment of mastitis [66].

Clinical guidelines emphasize effective milk removal, hydration, and analgesia as key elements of clinical management of mastitis. Healthy term infants may continue to breastfeed on the affected breast. If symptoms do not improve with conservative management or the woman is acutely ill, antibiotics are recommended. Penicillinase-resistant penicillins (e.g., dicloxacillin) are preferred [67].

Breast-feeding and Maternal Medications
Most medications are compatible with breast-feeding, or a safe alternative medication exists. The physiology of the placenta differs from that of the maternal breast and infant gut, so the provider should not assume that a drug’s pregnancy safety profile applies to breast-feeding. Many drug databases utilized by commercial pharmacies do not contain accurate information on drug safety in lactation [68]. The National Library of Medicine’s LactMed database (http://lactmed.nlm.nih.gov/) provides a comprehensive set
of monographs on medication safety in lactation. Decisions about medication use in lactation should consider the risks and benefits of maternal treatment or nontreatment, the risk of drug transfer to the infant, and the risks to mother and infant of discontinuing breast-feeding [69]. Maternal providers should collaborate with the pediatric provider regarding counseling about medication use in breast-feeding.

**Milk Expression**

If mothers and infants are separated, milk expression is necessary to maintain supply and provide milk to the infant. Evidence suggests that early initiation of milk expression, simultaneous pumping of both breasts, breast massage, relaxation techniques, and expressing milk after kangaroo (i.e., skin-to-skin) mother care improve milk production among neonatal intensive care unit (NICU) mothers [70–75].

**Galactogogues**

There is limited evidence to support pharmacotherapy for low milk supply [76,77]. Two small randomized, placebo-controlled, blinded studies have found that domperidone increases milk supply among mothers of preterm infants [78,79]. Domperidone is not approved by the U.S. Food and Drug Administration (FDA) for any indication. Routine use of metoclopramide in the early postpartum period does not improve milk production [80,81]. In two studies among women with low milk supply who received education on optimal breast-feeding [82,83], metoclopramide provided no additional benefit compared with placebo. The AAP notes that galactogogues have a limited role in facilitating lactation; mothers experiencing milk production difficulties should undergo assessment by a lactation specialist and be offered nonpharmacologic measures to increase supply [69].

**Maternal Diet during Lactation**

In two small trials, maternal dietary restrictions during lactation did not reduce the incidence of atopic eczema or the severity of existing disease [84]. Maternal supplementation with long-chain polyunsaturated fatty acids (LCPUFA) during lactation does not improve infant neurodevelopmental outcomes. In two studies, LCPUFA supplementation increased infant head circumference [85]. A meta-analysis found limited evidence for LCPUFA supplementation during pregnancy and/or lactation to reduce allergic diseases in children [86]. There is insufficient evidence to recommend LCPUFA supplementation during breast-feeding.

**Contraindications**

Breast-feeding is contraindicated in the setting of the following:

- Infant with classic galactosemia (galactose-1-phosphate uridyltransferase deficiency).
- Maternal medications:
  - Mothers with current, active illicit drug use, in absence of a coordinated treatment plan among maternal and infant providers [87].
  - Mothers requiring medications that are contraindicated in breast-feeding.
- Maternal infectious disease:
  - Mothers who are HTLV type I or II positive.
  - Mothers with untreated brucellosis.
  - Mothers with active herpetic lesions on the breast(s). Mothers can continue to breast-feed on the unaffected breast, and provide expressed milk from the affected breast.
- Mothers with varicella onset within 5 days before until 48 hours after delivery should be separated from their infants, but expressed milk can be provided to the infant.
- Mothers with HIV infection in settings where safe artificial feeding is available (see below).

**Tuberculosis**

The AAP lists active, untreated tuberculosis as a contraindication to breast-feeding. According to AAP guidelines, the infant may continue to receive expressed milk while the mother is treated, but mother and infant should be separated until the mother is treated for a minimum of 2 weeks and is documented to be no longer infectious [30]. The WHO recommends continued breast-feeding in the setting of maternal tuberculosis. The infant should be treated with 6 months of isoniazid preventive therapy, followed by immunization with Bacillus Calmette–Guérin (BCG) [88].

**Human Immunodeficiency Virus**

Artificial infant feeding is efficacious in preventing maternal–child transmission of HIV, but RCTs in regions of the world where access to clean water is limited demonstrate similar mortality and malnutrition among breast-fed and artificially fed infants. If infants are breast-fed, early exclusive breast-feeding and extended antiretroviral prophylaxis reduce the risk of HIV transmission [89].

Women with HIV in the United States are advised not to breast-feed because of the risk of maternal–infant transmission and the availability of safe artificial feeding.

The WHO recommends that national or subnational health authorities determine recommendations regarding infant feeding for HIV-positive mothers, balancing HIV prevention with protection from other causes of child mortality [90]. In countries where breast-feeding is recommended, antiretroviral prophylaxis for mothers and infants reduces HIV transmission. Exclusive breast-feeding is recommended for the first 6 months, continuing through 1 year. When mothers decide to stop breast-feeding, they are advised to wean gradually, over 1 month, and mothers or infants receiving antiretroviral prophylaxis should continue for 1 week after breast-feeding is fully stopped.

**CONTRACEPTION**

**Key Points**

- Postpartum educational interventions can increase contraceptive use and delay repeat pregnancy.
- Intrauterine device (IUD) placement <48 hours postpartum is not associated with infectious morbidity, but expulsion rates are higher than with placement >4 weeks. Since many women do not present for postpartum follow-up, overall IUD use is highest when inserted in the immediate postpartum period.
- Postpartum tubal ligation (PPTL) is highly effective. However, risk of regret is higher for PPTL than for interval TL.
- Postpartum placement of the contraceptive implant shows promise as an additional long-acting option for women early in the postpartum period, with potentially less lactogenic effects.
- Lactation amenorrhea is an effective but not perfect method of contraception, until the infant is 6 months old, begins complementary feeding, or the mother’s menses resume.
• Barrier methods are preferred during lactation compared with hormonal contraception because of theoretical effects of hormonal methods on milk supply and hormone exposure for the infant.

Background
Sexual activity is resumed by about 53% of women within 6 weeks postpartum, with 41% attempting vaginal sex. By 8 weeks, about 65% attempt vaginal sex. Women with uncomplicated vaginal births resume sex earlier, compared with women who had perineal lacerations, assisted vaginal births, or cesarean deliveries [91]. Two-thirds of women have unmet contraceptive needs after childbirth [92]. Educational interventions during the postpartum hospitalization increase contraceptive use, compared with no intervention. Home visit programs reduce repeat pregnancies among teenagers [93].

Contraceptive Guidelines
A detailed review of contraception is behind the scope of this book. The WHO has published detailed guidelines regarding timing of initiation of postpartum contraception and other relative and absolute contraindications. Postpartum recommendations for healthy women are summarized in Tables 30.3 and 30.4. The full guidelines are available online [94].

Intrauterine Devices
IUDs are highly effective methods of contraception. Unfortunately less than 50% of women who express interest in an IUD postpartum actually receive one [95]. Immediate postpartum placement has been shown to be safe, and allows women to access contraception during the maternity hospitalization though is associated with an increased risk of expulsion compared with delayed insertion [96,97]. In an RCT of postplacental versus delayed insertion, women randomized to postplacental insertion were more likely to have a device inserted (98% vs. 90.2%, p = .20). There were no differences between groups in IUD use at 6 months postpartum (84.3% vs. 76.5%). However, among women who were ineligible for the study and were advised to follow-up for IUD placement as part of routine postpartum care, only 26.8% were using an IUD at 6 months postpartum [98]. These results were confirmed in a more recent Cochrane review, with IUC use at 6 months twice as likely, though expulsion four times more likely [99]. These results suggest that women undergoing postplacental placement are more likely to use an IUD than those advised to follow-up for placement during routine postpartum care.

Postpartum Sterilization
Sterilization is the most commonly used form of contraception worldwide. Postpartum partial salpingectomy has a 1 year failure rate of 0.6/1000 and a 10-year failure rate of 7.5/1000, which compares favorably with other sterilization methods [100,101]. In a small RCT, operative times were shorter with postpartum Filshie clip placement compared with the Pomeroy technique, but failure rates were not evaluated [102]. Another larger RCT with 1400 women compared postpartum titanium clips to partial salpingectomy (Pomeroy technique);

Table 30.4 WHO Guidelines for Intrauterine Devices

<table>
<thead>
<tr>
<th>Breast-feeding</th>
<th>Not Breast-feeding</th>
</tr>
</thead>
<tbody>
<tr>
<td>LNG-IUD, &lt;48 hours postpartum</td>
<td>2</td>
</tr>
<tr>
<td>Cu-IUD, &lt;48 hours postpartum</td>
<td>1</td>
</tr>
<tr>
<td>LNG-IUD or Cu-IUD&gt;48 hours to &lt;4 weeks</td>
<td>3</td>
</tr>
<tr>
<td>&gt;4 weeks</td>
<td>1</td>
</tr>
<tr>
<td>Puerperal sepsis</td>
<td>4</td>
</tr>
</tbody>
</table>


Abbreviations: CU-IND, copper-bearing intrauterine device; LNG-IUD, levonorgestrel-releasing intrauterine device.

Notes: 1: A condition for which there is no restriction for the use of the contraceptive method; 2: A condition where the advantages of using the method generally outweigh the theoretical or proven risks; 3: A condition where the theoretical or proven risks usually outweigh the advantages of using the method generally; 4: A condition that represents an unacceptable health risk if the contraceptive method is used.

Table 30.3 WHO Guidelines for Postpartum Contraception: Hormonal Contraception

<table>
<thead>
<tr>
<th>Progestin-Only Methods</th>
<th>Combined Hormonal Contraceptives</th>
</tr>
</thead>
<tbody>
<tr>
<td>Breast-feeding</td>
<td></td>
</tr>
<tr>
<td>&lt;6 weeks postpartum</td>
<td>3/2c</td>
</tr>
<tr>
<td>&gt;6 weeks to &lt;6 months postpartum (primarily breast-feeding)</td>
<td>1</td>
</tr>
<tr>
<td>&gt;6 months postpartum</td>
<td>1</td>
</tr>
<tr>
<td>Not breast-feeding</td>
<td></td>
</tr>
<tr>
<td>&lt;21 days postpartum</td>
<td>1</td>
</tr>
<tr>
<td>&gt;21 days to 42 days</td>
<td>1</td>
</tr>
<tr>
<td>&gt;42 days</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
</tbody>
</table>


Abbreviations: VTE, venous thromboembolism; WHO, World Health Organization.

Notes: 1: A condition for which there is no restriction for the use of the contraceptive method; 2: A condition where the advantages of using the method generally outweigh the theoretical or proven risks; 3: A condition where the theoretical or proven risks usually outweigh the advantages of using the method; 4: A condition that represents an unacceptable health risk if the contraceptive method is used.

a Progestogen-only pills, levonorgestrel and etonogestrel implants, depot medroxyprogesterone acetate, and norethisterone enanthate.
b Combined oral contraceptives, combined contraceptive patch, combined contraceptive vaginal ring, and combined injectable contraceptives.
c 3: Depot medroxyprogesterone or norethisterone enanthate, 2: all other Progestogen-only methods.
though this study did not assess operative time, it did find that pregnancy probability at 2 years was .017 and .004 for clips and salpingectomy, respectively (p = .04) [103]. Maternal age <30 increases risk of regret and request for tubal reversal [104]. Risk of regret is higher among women undergoing PPTL compared with interval TL. Other risk factors for regret include ligation at the time of C-section, abrupt decision to undergo PPTL, and sterilization performed for obstetrical indications [105].

In a cohort study of women planning postpartum sterilization, one-third did not receive the procedure prior to hospital discharge. Of these women, 47% were pregnant within 1 year, compared with 22% of women who had not planned a postpartum sterilization [106]. Women who desire but do not undergo postpartum sterilization are at high risk for unplanned pregnancy.

Contraception during Breast-Feeding

Breast-feeding reduces fertility and prolongs amenorrhea after childbirth. In the first 6 months after birth, cohort studies suggest that 0%–7.5% of women who are fully breast-feeding and amenorrheic become pregnant [107,108]. To maintain effective protection against pregnancy, another method of contraception must be used as soon as menstruation resumes, the frequency or duration of breast-feeds is reduced, bottle feeds are introduced, or the baby reaches 6 months of age [94].

Nonhormonal methods, such as the copper IUD, and if these are not available, barrier methods, including diaphragm, cervical cap, male condom, and female condom, are the preferred method of contraception during breast-feeding [109]. The fall in progesterone after birth triggers lactogenesis, raising theoretical concerns that progesterone-containing contraceptives may interfere with lactation, particularly in the early postpartum period. Existing RCTs of hormonal contraception during lactation are of mixed quality, and evidence is insufficient to determine effects on milk production and quality [110]. The WHO medical eligibility criteria for contraceptive use guidelines note that although adverse health effects of exogenous estrogen in infants exposed to combined hormonal contraception have not been reported, existing studies have not been designed to quantify long-term effects. In addition, animal data suggest that progesterone may affect the developing brain, raising questions about the theoretical risks of progestogen-only injectables in the early postpartum period [111]. WHO recommendations regarding hormonal contraception during breast-feeding are listed in Table 30.3. In summary, given the mixed data on hormonal contraception and breast-feeding, the copper IUD or barrier methods are preferred.

WHO guidelines indicate that breast-feeding women can generally receive the levonorgestrel intrauterine device (LNG-IUD) at <48 hours postpartum [111]. In a small RCT [112], immediate postpartum placement of an LNG-IUD decreased exclusive breast-feeding rates at 3 and 6 months postpartum compared with insertion at 6–8 weeks. A study comparing insertion at 6–8 weeks of the LNG-IUD with the Cu T380A IUD found no differences in breast-feeding outcomes, infant growth, or development [113]. WHO recommendations regarding timing of IUD insertion are listed in Table 30.4.

The contraceptive implant is a progestosterone-only option that can be placed immediately postpartum. However, theoretical concerns regarding the effect on lactogenesis and lactation exist. In a recent RCT comparing early insertion (1–3 days postpartum) to standard insertion (n = 69), there was no difference in hours to lactogenesis (mean difference, −1.4 hours, 95% CI −10.6 to 7.7 hours) or lactation failure (risk difference 0.03, 95% CI −0.02 to 0.08) between the two groups [114]. There was no difference in the percentage of women partially or completely breast-feeding at 3 or 6 months. Women with early insertion were more likely to be contracepting at 3 months. A second small RCT (n = 24) used deuterium to index milk ingestion among healthy, term newborns of mothers randomized to immediate postpartum etonogestrel (ENG) implant placement or 6-week placement [115]. Participation was limited to nonobese women who had previously breast-fed for at least 3 months. No differences were found in milk intake.

POSTPARTUM ENDOMETRITIS

Key Points

- The diagnosis of postpartum endometritis is based on the presence of ≥2 of the following: fever >100.3°F, at least twice, ≥6 hours apart; fundal tenderness; tachycardia (heart rate >100 beats/minute); and foul-smelling lochia. Endometrial cultures are usually not necessary.
- Postpartum endometritis is most often associated with cesarean delivery; effective strategies include administration of preoperative antibiotics (either ampicillin or first-generation cephalosporin for just one dose), avoidance of manual placental extraction, nonclosure of both visceral and parietal peritoneum, and suture closure of the subcutaneous tissue when thickness is ≥2 cm.
- Gentamicin and clindamycin intravenously, preferably once daily dosing, are most effective for the treatment of postpartum endometritis.
- Once uncomplicated endometritis has clinically improved with intravenous therapy (usually 24–48 hours afebrile), oral therapy is not indicated.

Diagnosis/Definition

The diagnosis is based on clinical criteria. Table 30.5 describes criteria for diagnosis.

Symptoms/Signs

Those described in Table 30.5, plus abdominal pain, malaise, and elevated white blood cell count.

Epidemiology/Incidence

Endometritis complicates about 1%–3% of vaginal deliveries and 5%–27% of cesarean deliveries [116]. The lower incidence in certain cesarean delivery populations is due to infection precautions at delivery and antibiotic prophylaxis (see Chapter 13). In specific populations such as diabetic, obese, or HIV-positive patients, the risk appears to be higher [117,118].

Etiology/Basic Pathophysiology

Endometritis is an inflammatory process that involves both the endometrium and decidual tissue, secondary to infection. It is surmised that additional factors, (such as host defense, bacterial inoculum, and virulence) other than the presence of bacterial colonization play a role in pathogenesis, since 94% of postpartum patients have positive endometrial samples, but

Table 30.5  Diagnosis of Postpartum Endometritis (≥2 of the Following)

- Fever >100.3°F, at least twice, ≥6 hours apart
- Fundal tenderness
- Tachycardia (heart rate >100 beats/minute)
- Foul-smelling lochia
only a small fraction of these develop the infection. Bacteria usually ascend from the vagina and initially colonize the innermost layer of the endometrial cavity. If this colonization is not treated, infection can spread locally and through the bloodstream, leading to life-threatening complications. The use of prophylactic antibiotics has reduced, but not eliminated, these risks.

**Microbiology**

Endometrial infection is usually polymicrobial. Isolated bacteria are usually Gram-positive cocci, Gram-negative rods, or anaerobes that might be present in normal female genital tract and presumably reach the endometrium ascending from the vagina (Table 30.6) [119]. A recent systematic review identified isolates from the geneses *Bacteroides*, *Staphylococcus*, and *Streptococcus* as the most common pathogens associated with postpartum endometritis [120]. The presence of these microorganisms and the colonization of the decidua generate multiple microabscesses that trigger invasion of inflammatory cells. These cells release chemical mediators responsible for the different manifestations of endometritis. The presence of two microorganisms in the vagina as assessed by smears or cultures has been associated with postpartum development of endometritis.

**Risk Factors/Associations**

“Classic” risk factors for postpartum endometritis are shown in Table 30.7 [118,121–127]. Longer labor, longer duration of ruptured membranes, and more frequent vaginal examinations are associated with higher is the risk of infection, as is cesarean or operative vaginal delivery. HIV-positive women with CD-4 count ≤500 cells/mL have similar risks of postpartum endometritis and wound infection as HIV-negative women if they receive prophylactic antibiotics [128,129].

**Complications**

Treatment failure is uncommon. However, if fevers persist, addition of ampicillin to the primary antibiotic regimen should be considered. When fevers continue to persist, imaging to assess for infected hematoma, pelvic abscess, or septic pelvic thrombophlebitis should be considered. Sepsis is an uncommon complication when endometritis is actively managed. Though rare, endometritis secondary to Group A strep, Clostridium Sordelli, and *Staphylococci* can lead to necrotizing myometritis, toxic shock syndrome, and multiorgan system failure. Prompt recognition and treatment is imperative when these organisms are present.

**Management**

**Workup**

Endometrial cultures are usually not necessary, as the microorganisms identified are usually susceptible to the antibiotic regimens used [130]. Although not a current standard of care [130], some authors recommend the use of endometrial cultures at the time of diagnosis, advocating the need for specific antibiotic coverage [131], but no trials are available to assess their efficacy. Elevation of temperature may be the only sign found in patients with endometritis. Since one single episode of temperature ≥100.4°F (38°C) is commonly present in postpartum patients, and most of them will not develop any infection, it is recommended that two episodes of temperature elevation outside of the first 24 hours postpartum are identified in order to consider the diagnosis [132]. Physical examination is the cornerstone for assessment, with midline abdominal pain and uterine tenderness being central findings [133]. Although laboratory studies are not criteria for diagnosis, an increased neutrophil count, as well as elevated proportion of bands, may suggest the presence of an infectious disease [134]. leukocytosis alone is nonspecific and often physiologic in the postpartum period. However, urine analysis and culture should be obtained. The utility of routine blood cultures is unclear, though they are useful in immunocompromised patients, those at increased risk for bacterial endocarditis, and for women who present with overt sepsis or fail initial antibiotic therapy [135]. In selected patients, chest X-rays can be taken. Differential diagnosis for a postpartum febrile episode includes atelectasis, pneumonia, viral syndrome, engorgement, mastitis, pylonephritis, and appendicitis.

**Table 30.6** Microorganisms More Frequently Associated with Postpartum Endometritis

<table>
<thead>
<tr>
<th>Facultative Gram Positive: 51%</th>
<th>Facultative Gram Negative: 28%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group B streptococci</td>
<td>Gardnerella vaginalis</td>
</tr>
<tr>
<td>Enterococci</td>
<td>Escherichia coli</td>
</tr>
<tr>
<td><em>Staphylococcus</em> epidermidis</td>
<td>Enterobacter spp.</td>
</tr>
<tr>
<td>Lactobacillus</td>
<td><em>Proteus mirabilis</em></td>
</tr>
<tr>
<td>Diphtheroids</td>
<td>Others</td>
</tr>
<tr>
<td><em>S. aureus</em></td>
<td></td>
</tr>
<tr>
<td>Others</td>
<td></td>
</tr>
<tr>
<td>Anaerobic: 49%</td>
<td></td>
</tr>
<tr>
<td><em>Bacteroides bivius</em></td>
<td></td>
</tr>
<tr>
<td>Peptococcus</td>
<td></td>
</tr>
<tr>
<td><em>asaccharolyticus</em></td>
<td></td>
</tr>
<tr>
<td><em>Bacteroides</em> spp.</td>
<td></td>
</tr>
<tr>
<td><em>Peptostreptococcus</em> spp.</td>
<td></td>
</tr>
<tr>
<td>Others</td>
<td></td>
</tr>
<tr>
<td>Veillonella spp.</td>
<td></td>
</tr>
<tr>
<td>Others</td>
<td></td>
</tr>
</tbody>
</table>

Source: Modified from Rosene K et al., *J Infect Dis*, 153(6), 1028–1037, 1986.

**Table 30.7** Risk Factors for Postoperative Endometritis

<table>
<thead>
<tr>
<th>Cesarean delivery (directly correlated to operative time)</th>
<th>Labor (directly correlated to duration)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rupture of membranes (directly correlated to duration)</td>
<td>Socioeconomic status</td>
</tr>
<tr>
<td>Number of vaginal examinations</td>
<td></td>
</tr>
<tr>
<td>Internal fetal monitoring</td>
<td></td>
</tr>
<tr>
<td>Manual extraction of placenta</td>
<td></td>
</tr>
<tr>
<td>Epiotomoy</td>
<td></td>
</tr>
<tr>
<td>Operative vaginal delivery</td>
<td></td>
</tr>
<tr>
<td>Chorioamnionitis</td>
<td></td>
</tr>
<tr>
<td>Preterm delivery</td>
<td></td>
</tr>
<tr>
<td>Young age (&lt;17 years old)</td>
<td></td>
</tr>
<tr>
<td>Obesity (BMI &gt;30)</td>
<td></td>
</tr>
<tr>
<td>Postterm pregnancy</td>
<td></td>
</tr>
<tr>
<td>Blood loss</td>
<td></td>
</tr>
<tr>
<td>Bacterial vaginosis</td>
<td></td>
</tr>
<tr>
<td>GBS colonization</td>
<td></td>
</tr>
<tr>
<td>Diabetes</td>
<td></td>
</tr>
</tbody>
</table>


Abbreviations: BMI, body mass index; GBS, group B streptococcus.
Prophylactic antibiotics (either ampicillin or first-generation cephalosporin for just one dose about 30 minutes before skin incision), spontaneous placental removal, nonclosure of both visceral and parietal peritoneum, and suture closure of the subcutaneous tissue when thickness is ≥2 cm should routinely be performed in cesarean delivery (see Chapter 13). Vaginal cleansing with povidone-iodine prior to cesarean delivery, especially in the setting of labor or rupture of membranes should be considered especially if preincision antibiotics cannot be administered [136]. Each of these interventions decreases the incidence of postpartum endometritis and/or fever. In particular, both ampicillin and first-generation cephalosporins have similar efficacy in reducing postoperative endometritis from about 18% to 12%, with no added benefit found in using more broad-spectrum agents [137,138]. In urban, indigenous, mainly Afro-American women with incidence of postpartum endometritis over 24% despite first-generation cephalosporin prophylaxis, the addition of doxycycline 100 mg together with the cephalosporin and azithromycin 1 g orally 6–12 hours postoperatively can decrease the incidence of endometritis to 17%, possibly by targeting Ureaplasma urealyticum [139]. The number of vaginal examinations, nulliparity, early gestational age, and cefazolin use were predictors of prophylaxis failure in one study [140]. Cleansing of the vagina with povidone-iodine prior to cesarean delivery has been associated with a reduction in the risk of postpartum endometritis, particularly among women with ruptured membranes [141] (see Chapter 13). In HIV-positive women with a CD4 count less than 500 who had vaginal deliveries, intrapartum cefoxitin reduced the rate of postpartum endometritis by 53% (95% CI 0.24–0.9) [117].

Therapy
Since more than one microorganism is usually involved, a combination of antibiotics is used to assure adequate coverage and prevent resistance. Parenteral, broad-spectrum antibiotics should be initiated and continued until the patient is afebrile. The combination of gentamicin and clindamycin is appropriate for the treatment of endometritis [14]. Compared with clindamycin and an aminoglycoside, other regimens have a 1.44 relative risk (RR) of treatment failure [142]. Compared with regimens with activity against penicillin-resistant anaerobic bacteria, regimens with poor activity have a 1.94 RR of treatment failure. There was no evidence of difference in incidence of allergic reactions. Cephalosporins were associated with fewer diarrheas. There is no evidence that any one regimen is associated with fewer side effects [142].

In four studies comparing once daily with thrice daily dosing of gentamicin, there were fewer failures with once-daily dosing. Once-daily gentamicin can be given 5 mg/kg, and once-daily clindamycin phosphate 2700 mg, both intravenously [143]. Thrice-daily dosing consists of clindamycin 900 mg and gentamicin 1.5 mg/kg every 8 hours. It is recommended that levels (peak/trough) of gentamicin should be taken after the third dose, to make sure therapeutic regimens are achieved. Once uncomplicated endometritis has clinically improved, oral therapy is not needed; in four trials, there were no differences in outcome with oral therapy versus no oral therapy [142].

In low resource setting, where intravenous options are not readily available, alternative regimens should be considered, as untreated infection can lead to sepsis. In a recent systematic review, oral clindamycin with intramuscular gentamicin was suggested as the first line recommendation, with amoxicillin–clavulanic acid or amoxicillin and metronidazole as alternative regimens [120]. Each of these regimens had at least an 85% cure rate, and they are compatible with breast-feeding.

Response is usually prompt. If fever persists >48 hours (<10% of women), the addition of ampicillin can be considered. If fever still persists, pelvic abscess, wound infection, septic pelvic thrombophlebitis (SPT), inadequate antibiotic coverage, and retained placental tissue should be ruled out. SPT, though uncommon, should be suspected when fevers follow a spiking pattern and are unresponsive to antibiotics for more than 5 days [144]. Imaging (CT or MRI) is often used to help aid in the diagnosis of SPT, though no data exist to suggest that one is superior to the other. These modalities can also evaluate for pelvic abscess. Treatment for SPT has been conventionally anticoagulation, though there is no high-quality data to support this. There is one randomized trial with 14 patients that compared antibiotics alone to heparin and antibiotics for women with CT evidence of SPT; the primary outcomes of fever duration and hospital stay did not differ between the two groups [145]. Despite these limited data, expert opinion suggests anticoagulation for women with SPT [144,145]. For women with negative imaging, the likelihood of a resistant organism, a nongenital source of infection (pyelonephritis, pneumonia, and intravenous catheter phlebitis), or noninfectious fever should also be considered [131].

Because of neonatal implications, information on the mother's condition should be provided to the neonate's health-care provider [130].

Breast-feeding
Most antibiotics, including clindamycin, gentamicin, and penicillin, are considered safe in breast-feeding, although they may cause changes in infant gastrointestinal flora, potentially resulting in diarrhea or thrush [146]. The mother and the pediatric provider should be advised so that they can monitor the infant for side effects.

POSTPARTUM WOUND COMPLICATIONS

Key Points
- Risk factors for post cesarean wound infection are chorioamnionitis, maternal preoperative condition or infection, preeclampsia, higher body mass index (BMI), nulliparity, increased surgical blood loss, and diabetes.
- Prophylactic preincision antibiotics (either ampicillin or first-generation cephalosporin for just one dose), suture closure or drainage of subcutaneous fat in women with ≥2 cm thickness, and suture closure of skin reduce the risk of post cesarean wound infection.
- Penicillins are first line antibiotics for wound infection. Wound drainage and debridement of necrotic tissue may be necessary.
- Compared with healing by secondary intention, reclosure of the disrupted laparotomy wound after the infection has resolved is associated with success in >80% of women, faster healing times, and fewer office visits.

Diagnosis/Definition
The Centers for Disease Control defines and classifies surgical site infection (SSI) as either superficial, deep or organ/space, as shown in Table 30.8. These criteria should be addressed at the time of diagnosis [147]. The vast majority of significant wound infections in obstetrics are postpartum following cesarean delivery. Breakdown and infection of perineal repair is uncommon.
and staples for skin closure 

Risk factors described are 

Risk factors for perineal break down include third or fourth degree laceration, episiotomy, operative delivery, meconium, and a prolonged second stage. In one retrospective case-control, an operative delivery with a mediolateral episiotomy was associated with a 6.36-fold increased risk of perineal breakdown or infection, while third/fourth degree laceration and meconium were associated with a 3.7- and 3.22-fold increase risk, respectively [161]. Among women with obstetric anal sphincter injuries, a large prospective study noted at 19.8% infection rate and a nearly 25% breakdown rate, with operative delivery being significantly associated with both outcomes (OR 2.54, 95% CI 1.32–4.87, p = .008) [162]. In this study, intrapartum obstetric antibiotics were associated with a decreased risk of wound complications (OR 0.50, 95% CI 0.27–0.94, p = .03).

**Prevention**

Prophylactic antibiotics (either ampicillin or first-generation cephalosporin for just one dose) are associated with a 59% decrease in the incidence of wound infection compared with no antibiotics in women undergoing cesarean delivery [138]. The timing for prophylaxis should be within 30–60 minutes before the start of procedure, as while it is not associated with

**Epidemiology/Incidence**

The incidence of wound infection after cesarean delivery ranges from 2.8%–11.6% to 16.6%–30% in obese patients [148–150]. Perineal wound breakdown occurs between 0.1%–4.6% of deliveries, though actual incidence may be higher due to underreporting [151,152].

**Table 30.8 Classification of Surgical Site Infection (SSI)**

<table>
<thead>
<tr>
<th>Classification</th>
<th>Occurs within 30 days after surgical procedure, involves skin and subcutaneous tissue only, and at least one of the following:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Superficial</td>
<td>a. Purulent drainage from incision</td>
</tr>
<tr>
<td></td>
<td>b. Organism isolated from culture of fluid or tissue from SSI</td>
</tr>
<tr>
<td></td>
<td>c. At least one of the following: pain, redness, swelling or heat, and superficial incision is deliberately opened by the surgeon</td>
</tr>
<tr>
<td></td>
<td>d. Diagnosis of superficial SSI by surgeon or attending physician</td>
</tr>
<tr>
<td>Deep</td>
<td>Occurs within 30 days from surgical procedure and involves deep soft tissues (fascia, muscle) of the incision, and at least one of the following:</td>
</tr>
<tr>
<td></td>
<td>a. Purulent drainage from incision</td>
</tr>
<tr>
<td></td>
<td>b. Spontaneous dehiscence or deliberately opened by a surgeon when the patient presents at least one of the following: fever, pain, and tenderness</td>
</tr>
<tr>
<td></td>
<td>c. An abscess involving the deep incision is found on direct examination during reoperation, or by histopathologic or radiologic examination</td>
</tr>
<tr>
<td></td>
<td>d. Diagnosis of deep SSI by surgeon or attending physician</td>
</tr>
<tr>
<td>Organ/Space</td>
<td>Occurs within 30 days from surgical procedure and the infection appears to be related to the operation, and infection involves any part of the anatomy (organs, spaces) other than the incision, which was opened or manipulated during an operation, and at least one of the following:</td>
</tr>
<tr>
<td></td>
<td>a. Purulent drainage from a drain placed in the organ/spaces</td>
</tr>
<tr>
<td></td>
<td>b. Organisms isolated from a culture of fluid or tissue in the organ/spaces</td>
</tr>
<tr>
<td></td>
<td>c. An abscess involving the organ/spaces found on direct examination during reoperation or by histopathologic or radiologic examination</td>
</tr>
<tr>
<td></td>
<td>d. Diagnosis of an organ/spaces SSI by surgeon or attending physician</td>
</tr>
</tbody>
</table>

**Symptoms**

Wound complications most commonly present with pain, redness, swelling or drainage from the incision site. Fever and purulent drainage may be signs that infection is present. Perineal infection may be associated with malodorous discharge, pain with urination and/or defecation, and dyspareunia.

**Classification**

See Table 30.8.

**Risk Factors/Associations**

Risk factors described are preoperative remote infection, chorioamnionitis, maternal preoperative condition, preeclampsia, higher BMI, nulliparity, increased surgical blood loss, and staples for skin closure [122,153–156]. Women with diabetes have five times the risk of wound infection compared with women without diabetes [157]. One large case-control study identified subcutaneous hematoma as an additional risk factor for wound infection, OR 11.6, 95% CI 4.1–33.2) [158]. Risk for wound infection increase with increasing BMI [159]. Staples removed prior to postoperative day 4 are also associated with increased risk of wound complication [160]. Staple closure has been associated with a twofold higher rate of wound infection and complications [155,156].

**Etiology/Basic Pathophysiology**

The most common microorganisms identified by cultures from wound infections after cesarean delivery include *Staphylococcus epidermidis*, *Enterococcus faecalis*, *S. aureus*, *Escherichia coli*, and *Proteus mirabilis*. The pathophysiology involves seeding of bacteria either from the uterine cavity or from the skin [148].

**Management**

**Workup**

For women presenting with history concerning for a wound complication, perineal or abdominal, thorough examination of the incision is important. Areas of erythema, induration and fluctuance should be noted, and potentially marked to assess for spread over time. For an abdominal incision, a sterile swab can be used to probe any open areas to assess the dimensions of the separation and to ensure fascial integrity. Observation of any discharge (serosanguinous vs. bloody vs. purulent) may be helpful in diagnosis. Imaging is not indicated unless there is concern for fascial dehiscence or more severe infection.

In early onset wound infection (<48 hours after the procedure), the microorganisms are likely to be either group A streptococcus or *Clostridium*. A wound culture can be taken and a Gram stain will depict either Gram-positive cocci or Gram-positive rods, respectively. There are no trials to confirm the efficacy or utility of obtaining a wound culture from abdominal or perineal wounds.
significant change in neonatal outcomes it reduces risk of maternal infectious morbidity (RR 0.59, 95% CI 0.44–0.81) [163–165].

Suture closure of subcutaneous fat in women with >2 cm thickness is associated with a significant decrease in wound disruptions, including infections [166] (see also Chapter 13). For incisions closed with staples, removal on or after POD4 also decreases wound complications [155,156]. Based on a recent meta-analysis and more recent RCT, suture should be strongly encouraged over staples [155,156]. Some centers are using negative pressure wound therapy (NPWT) empirically in morbidly obese patients in an effort to decrease wound complications; early data appear promising [150].

One RCT examining the benefit of prophylactic antibiotics during repair of third and fourth degree lacerations exists. This trial of 147 women did show a difference in perineal wound complication rates between women who received the antibiotics and those who did received placebo (8.4% vs. 24.1%) [167]. This trial used a single dose of intravenous cefoxitin. Further study is needed before recommendations can be made [165,168] (see Chapter 9).

**Therapy**

For women with perineal wound breakdown, a thorough examination (potentially under anesthesia) is helpful to better characterize the defect and adequately cleanse the wound. There is insufficient evidence to suggest resuturing or allowing the wound to heal by secondary intention in the absence of infection [169]. Given the active bacterial ecosystem of the vagina, some providers will assume that a subclinical infection is the reason for the wound breakdown and treat with antibiotics. In the setting of active perineal infection, debridement with antibiotic therapy is indicated. Antibiotic regimens vary, though some experts suggest a 7-day course of 875 mg amoxicillin–clavulanate and 500 mg of metronidazole twice daily [162]. Similar to wound breakdown, insufficient evidence exists regarding repair of the defect versus conservative management in the setting of perineal infection; clinical judgment and patient preferences should be used make this decision [169].

For women with cesarean delivery, the onset of the possible abdominal wound infection defines the need for antibiotics. For infection arising <48 hours after cesarean, penicillin is the drug of choice, based largely on expert opinion. Empiric therapy for cellulitis is reasonable with dicloxacillin, cephalaxin, or clindamycin [170]. Wound drainage and debridement of necrotic tissue may be necessary. In late-onset wound infection (4–8 days postoperatively), the management consists purely of drainage. Antibiotics are not considered indicated in this setting, unless extensive cellulitis is present, or if the patient does not improve after drainage, in which case necrotizing fasciitis should be considered [131].

**Disrupted (Open) Laparotomy Wound, after Infection Has Resolved or If No Infection on Presentation**

There are different ways to manage the open wound [171]. Women who present to the clinic or emergency wound with evidence of an abdominal wound separation, hematoma, or seroma without evidence of infection need a thorough evaluation of the incision. The incision should be cleansed with sterile water or saline. When hematoma is present, the wound may need to be partially opened in order to evacuate the hematoma, thus decreasing the risk of subsequent infection.

Compared with healing by secondary intention, reclosure of the disrupted laparotomy wound is associated with success in >80% of women, faster healing times (16–23 vs. 61–72 days), and fewer office visits [172]. No serious morbidity or mortality is associated with either method. There is insufficient evidence to assess optimal timing (probably 4–6 days after disruption if noninfected) and technique (superficial vertical mattress or “en bloc” reclosure of entire wound thickness with absorbable sutures, or adhesive tape) of reclosure, as well as utility of antibiotics. After the wound is free of infection and is granulating properly, local anesthesia can be applied at bedside, and polypropylene mattress stitches can be used to close the skin.

Compared with reclosure using sutures, reclosure using permeable, adhesive tape (Cover-Roll; Biersdorf Inc., Norwalk, CT) was a faster procedure with lower pain scores and similar healing times in a small trial [173]. However, a subsequent study found shorter healing times (16.1 vs. 23.0 days) with suture closure compared with surgical tape [174].

Closure with secondary intention using NPWT, also called vacuum-assisted closure, has not been studied in any trials after cesarean section [175]. A recent meta-analysis in nonpregnant adults found some evidence of improved healing with NPWT but concluded that existing data are insufficient to establish clinical benefit [176].

**POSTPARTUM MOOD AND ANXIETY DISORDERS**

**Key Points**

- See Chapter 21 in *Maternal-Fetal Evidence Based Guidelines* for details of management of Mood disorders in pregnancy.
- One in 7 women experience perinatal depression (PND), making it the most common complication of pregnancy.
- Risk factors include history of maternal anxiety or depression, lack of social support traumatic birth experience, infant admission to neonatal intensive care, and breastfeeding problems.
- Routine screening is recommended at least once during perinatal care.
- Positive screens require additional evaluation. If depression is diagnosed, referral and follow-up to ensure treatment occurs is essential.

**Definition**

PND is an episode of moderate or severe major depressive disorder (MDD) beginning either during pregnancy or within 4–6 weeks after delivery [177,178].

**Symptoms/Signs**

In addition to typical depression symptoms of sadness, despair, disrupted sleep and appetite, women with postpartum depression often experience prominent anxiety symptoms.

**Epidemiology/Incidence**

Its prevalence is about 15%, affecting one in seven women [179,180].

**Risk Factors and Associations**

Risk factors for prenatal depression include past history of depression or anxiety, life stressors, lack of social support, unintended pregnancy, lower income or education, domestic violence, and smoking. Additional risk factors for postpartum depression include a traumatic birth experience, infant admission to neonatal intensive care, and difficulties breast-feeding [181].
Complications
During the postpartum period, women with depression are at increased risk of maternal suicide, infanticide, and impaired maternal sensitivity and attachment with the infant [182-185]. Women with depression are also less likely to engage in enriching interactions with the child, such as reading or singing [186]. Anxiety symptoms have been associated with increased maternal healthcare utilization and reduced breast-feeding duration [187,188].

Prevention
Psychosocial and psychological interventions are effective to reduce postpartum depression. In a large meta-analysis, both lay and professional prevention interventions reduced postpartum depression symptoms (average RR 0.78, 95% CI 0.66–0.93). Effective strategies included postpartum home visits, telephone support, and interpersonal psychotherapy.

Screening
Screening for PND should be performed using a validated instrument, such as the Edinburgh Postnatal Depression Scale (EPDS). Despite the high prevalence and substantial morbidity associated with PND, this condition is under-recognized. Major obstetric and pediatric medical organizations recommend routine screening [3,181]. Although clinicians are generally supportive of screening for PND [189,190], these attitudes do not consistently translate into practice. In the United States, less than half of women are formally screened for PND [191]. Consistent with such efforts, less than 50% of PND cases are detected in routine clinical practice, with antenatal recognition rates reported at 41% [192] and postnatal rates ranging from 29% [193] to 43% [194]. Screening alone does not ensure treatment: in a recent review, one in five women who screened positive had at least one mental health visit [195]. Patient engagement strategies increased the change that women with depression symptoms received mental health care.

Treatment
Both psychotherapy and medications are effective for treating PND. For further details on treatment of postpartum depression, see Chapter 21 in Maternal-Fetal Evidence Based Guidelines. As with any patient diagnosed with a health condition, women with postpartum depression should be followed-up to ensure adequate engagement with treatment and response to therapy. Monitoring patient symptoms is essential, just as measurement of blood pressure is necessary to guide treatment of hypertension. A stepped care approach with serial measurement of symptoms with a tool such as the Patient Health Questionnaire 9 (PHQ-9) to inform depression management can increase the likelihood that patients receive adequate therapy [196].

OTHER POSTPARTUM ISSUES
Women report receiving insufficient guidance from their providers about recovery from birth, and feeling unprepared for postpartum concerns is correlated with postpartum mood symptoms [197]. In a randomized trial, routine provision of guidance about expected symptoms and coping strategies measures reduced depression symptoms [198] and increased breast-feeding duration [199] among black and Latina women.

Table 30.9 Postpartum (Including Postcesarean) Advice Regarding Common Daily Life Activities

<table>
<thead>
<tr>
<th>Advice</th>
<th>Evidence</th>
<th>Suggested Recommendations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lifting</td>
<td>Lifting increases intra-abdominal pressure much less than Valsalva, forceful coughing, or rising from supine to erect position [201].</td>
<td>(1) Patients should continue lifting patterns as before pregancy.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(2) Patients need an adequate postoperative analgesic regimen as necessary.</td>
</tr>
<tr>
<td>Climbing stairs</td>
<td>Climbing stairs increases intra-abdominal pressure much less than Valsalva, forceful coughing, or rising from supine to erect position [201].</td>
<td>(1) Patients should continue climbing stairs as before pregnancy.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(2) Patients need an adequate postoperative analgesic regimen as necessary.</td>
</tr>
<tr>
<td>Driving</td>
<td>No consistent prospective or retrospective evidence.</td>
<td>(1) Patients need an appropriate postoperative analgesic regimen that does not cause a clouded sensorium when driving.</td>
</tr>
<tr>
<td>Exercise</td>
<td>Limited retrospective and prospective evidence.</td>
<td>(2) Patients may resume driving when comfortable with hand and foot movements required for driving.</td>
</tr>
<tr>
<td></td>
<td>Forceful coughing increases intra-abdominal pressure as much as jumping jacks [201].</td>
<td>(3) Exercise program may need to be tailored for postpartum women.</td>
</tr>
<tr>
<td>Vaginal intercourse</td>
<td>No retrospective or prospective evidence.</td>
<td>(4) Preprocedure and postprocedure recommendations should be consistent.</td>
</tr>
<tr>
<td>Returning to work</td>
<td>No consistent retrospective evidence; no prospective evidence,</td>
<td>(1) Women and their partners should make the decision to resume intercourse mutually.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(2) Women should use appropriate contraception after childbirth.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(3) There is insufficient evidence as to when to resume vaginal intercourse.</td>
</tr>
</tbody>
</table>

There is insufficient evidence for general postpartum advice on common issues such as lifting, climbing stairs, driving, exercise, vaginal intercourse, and return to work. Suggestions based on this limited evidence and mostly expert opinions are shown in Table 30.9 [200,201]. Women are routinely counseled to defer intercourse until 6 weeks postpartum, but there are no data supporting this recommendation. The risk of bleeding or infection is minimal after 2 weeks, and guidelines recommend that women resume intercourse when it is comfortable and desired, once the perineum has healed and bleeding has decreased [202]. More than half of women resume sexual intercourse by 6 weeks postpartum, and 90% resume by 12 weeks postpartum [203]. Multiple prospective cohort studies have attempted to identify risk factors for persistent dyspareunia after delivery. Though data are mixed, Connolly, Serati, and Barrett each conducted prospective cohort studies, which failed to implicate mode of delivery, episiotomy, and perineal lacerations as risk factors for postpartum dyspareunia [204–206]. Barrett et al. [204] did note that a history of dyspareunia was a major risk factor for postpartum dyspareunia (OR 4.97).

For other postpartum issues, see also Chapter 1, Chapter 9 (third stage of labor and its complications—including repair of vaginal lacerations, etc.), Chapter 13 (cesarean delivery—especially for prevention of postpartum endometritis and postpartum wound infection), and chapters pertinent to specific conditions in both this book and its companion Maternal-Fetal Evidence Based Guidelines.

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The neonate
Gary A. Emmett and Swati Murthy

KEY POINTS

- **Necessary equipment** for neonatal stabilization must be available at every delivery.
- **Personnel trained in neonatal resuscitation** should be present at every delivery.
- An infant who is 37+ weeks of gestation, is crying or breathing, and has good muscle tone does not require resuscitation. Dry the baby, place the baby skin-to-skin with the mother, and cover the baby with dry linen to maintain temperature.
- Provide routine care, provide warmth, assure open airway, dry, and continuously evaluate color, activity and breathing.
- Neonatal resuscitation begins if child does not proceed through successful transition, not based on a specific Apgar number at any time. Stimulate the baby and open the airway. Routine suctioning is no longer suggested. If further resuscitation is needed, it is frequently due to poor respiratory transition and can usually be resolved with bag-and-mask support of airway and breathing.
- **Initial bag-and-mask resuscitation should be with room air.** Oxygen should be titrated with continuous pulse oximetry to maintain the baby within the target saturations (Table 31.6).
- **Low-risk infants** are 37 weeks of gestation or older and have birth weight between 2500 and 4200 g, Apgar scores of 7 or above at 5 minutes, normal vital signs, and no signs of congenital anomalies or respiratory distress.
- A difficult transition may be anticipated by infants who are not low risk and may be due to hypothermia, hypoglycemia, or congenital anomalies.
- The **preterm infant** (especially 35 weeks of gestation or less) is at greater risk for complications in the delivery room and in the well nursery. With high-risk infants, there must be immediate access to means of establishing intravenous (IV) access and airway pressure.
- **Late-preterm infants** are those born at 34 0/7–36 6/7 weeks. While these infants are often the size and weight of some term infants, they are physiologically more immature and at higher risk for complications.
- Every nursery should have preexisting protocols to institute best stabilization practices and should have clear statements of what conditions and degree of illness are appropriate to go to the well-baby nursery.
- The majority of cases of cerebral palsy do not result from isolated intrapartum asphyxia with resultant hypoxemia and organ damage.
- The health benefits to circumcision include prevention of urinary tract infection, penile cancer, and transmission of sexually transmitted infections. In the United States, the benefits are not considered great enough for universal circumcision of males.

DELIVERY ROOM MANAGEMENT

For management of the normal neonate, see also Chapter 9 (third stage). For management specific to meconium, see Chapter 23. The perinatal period is the riskiest time in a child’s life [1–3]. Every newborn has the right to a resuscitation performed at a high level of competence. A competent resuscitation means that the proper equipment and well-trained personnel must be at every delivery. **Necessary equipment** is shown in Table 31.1.

**Initiation of Resuscitation**

Due to the complex physiologic changes that occur at birth, many newborns will experience apnea or bradycardia that will require opening the airway, stimulation, and ventilation. The need for medication is rare.

The first steps include the following:

- **Thermal management**: Placing the infant under a preheated radiant warmer, drying and stimulating the newborn, and frequently replacing wet blankets with dry ones.
- **Opening the airway**: Because of evidence that suctioning of the nasopharynx can lead to bradycardia during resuscitation, the first step is simply opening the airway. Clearing the airway should be reserved for babies with obvious obstruction of the airway or who require positive pressure ventilation. Most historic methods of removing meconium from the airway (suctioning of the oral pharynx before delivery of the shoulders and elective endotracheal intubation and direct suctioning of the trachea) have not been shown to improve outcomes. There is insufficient evidence to recommend a change in the current practice of endotracheal suctioning of nonvigoruous babies with meconium stained amniotic fluid, but if this cannot be done easily and quickly, start bag-mask ventilation, especially if child is bradycardic.
- **Tactile stimulation**: Rubbing the baby’s back, trunk, or extremities may occasionally be necessary to stimulate the baby to normal breathing and heart rate. For persistent apnea after tactile stimulation, positive pressure ventilation should be initiated immediately.

Chest compressions may be necessary if there is continued bradycardia (heart rate less than 60 beats/minute) after breathing is adequately supported. On rare occasions, medications are necessary and should be available (refer to Table 31.2 for the full list.). The most commonly used are as follows:

1. **Epinephrine 1:10,000**: 0.1–0.3 mL/kg IV or via endotracheal tube (ETT) given rapidly
2. **Volume expanders**: normal saline, Ringer’s lactate, or whole blood. Dose is 10 mL/kg IV over 5–10 minutes
3. Sodium bicarbonate 0.5 mEq/mL: 2 mEq/kg IV given over at least 2 minutes
4. Naloxone hydrochloride 1.0 mg/mL: 0.1 mg/kg IV or intra-muscularly (IM) given rapidly

### Delivery Room Resuscitation

The need to treat the newborn is based on frequent evaluation of respirations, heart rate, and oxygenation. The algorithm in Figure 31.1 can be used as a guideline for the resuscitation of a term or late preterm infant. The Apgar score (outlined in Table 31.3) is a useful tool for communicating an infant’s status, but should not be used to guide resuscitation.

### The Difficult Transition

Successful transition to extrauterine life generally occurs over the first hours after birth. Delay in transition can occur for many reasons. Common causes of delayed transition are listed in Table 31.4.

In infants with signs of poor transition (tachypnea, cyanosis, mottled skin or pallor, tremors, and/or jitteriness), additional measurements of the child’s vital signs should be considered. These include temperature, blood glucose, and arterial blood saturation (by pulse oximetry or blood gas). Some of the more common reasons for delayed transition are discussed below.

#### Hypoglycemia

Low blood glucose in the newborn is a common problem often associated with a diabetic mother or with abnormal in utero growth—either too small or too large. A less common cause is congenital abnormalities of the pancreas. The exact definition of hypoglycemia is not agreed upon but all symptomatic infants should be treated. Treatment should be initiated immediately. Specific guidelines vary with the institution.

According to the 2011 report by the Committee of Fetus and Newborn of the American Academy of Pediatrics (AAP), treatment should proceed as in Table 31.5. Screening should be limited to term babies who are symptomatic (see Table 31.5), or ones that are at high-risk, which include babies that are less than 37 weeks gestation, small for gestational age (SGA), large for gestational age (LGA), and/or infant or diabetic mother (IDM). Routine screening of blood glucose is not needed in healthy, term newborns after uneventful pregnancy and delivery, except if they are symptomatic. No studies have demonstrated harm from asymptomatic hypoglycemia in the first 12 hours of life. Screening in late preterm infants and SGA infants should continue up to at least 24 hours. After 24 hours, any infant with glucose that is persistently <45 mg/dL should continue to be screened until stable. If glucose is not stable at 24 hours of age, or symptoms continue in spite of treatment, transfer to a higher level of care is necessary [4].
Meconium is the thick, green bowel movement of a newborn infant. All newborns should have a bowel movement within 24 hours of birth. Intrauterine stress or stress during delivery may cause the meconium to pass early. Meconium-stained fluid may be present in 8%–20% of deliveries and is almost exclusively found in term or postterm infants [1]. The fluid may be lightly stained and thin or very dark and thick, but all meconium-stained fluid presents a risk to the baby. The treatment is based on the condition of

**Table 31.3** Apgar Score

<table>
<thead>
<tr>
<th></th>
<th>0</th>
<th>1</th>
<th>2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Color</td>
<td>Blue/pale</td>
<td>Acrocyanosis</td>
<td>Pink</td>
</tr>
<tr>
<td>Heart rate</td>
<td>Absent</td>
<td>&lt;100/minute</td>
<td>&gt;100/minute</td>
</tr>
<tr>
<td>Reflex/irritability</td>
<td>No response</td>
<td>Grimace</td>
<td>Cry/active withdrawal</td>
</tr>
<tr>
<td>Tone</td>
<td>Limp</td>
<td>Some flexion</td>
<td>Active motion</td>
</tr>
<tr>
<td>Respiration</td>
<td>Absent</td>
<td>Weak cry/hypoventilation</td>
<td>Good/crying</td>
</tr>
</tbody>
</table>

**Figure 31.1** Neonatal Resuscitation Program (NRP) guidelines for resuscitation. CPAP, continuous positive airway pressure; HR, heart rate; PPV, positive pressure ventilation.
the infant at birth. If the baby is vigorous and cries immediately, resuscitation should proceed normally. If the infant is limp or lethargic, he or she should be delivered immediately and placed on the warmer. Meconium suctioning, by intubating the baby and directly suctioning the trachea with a meconium aspirator attached to the ETT, may be helpful if it can be done quickly and easily. If not, simply bag the baby with room air. Current recommendation is to not suction the baby before delivery of the shoulders and to apply mask-and-bag resuscitation with room air as soon as possible to a baby who is not vigorous.

### Respiratory Distress

Respiratory distress and/or tachypnea may occur and can be the manifestation of underlying illnesses. Etiologies include abnormalities of the lung (respiratory distress or aspiration syndromes, infections, pneumothorax, and congenital lung abnormalities), structural airway problems (choanal atresia, tracheal-esophageal fistula, and micrognathia), abnormalities of the cardiovascular system (primarily congenital heart disease—cyanotic or not, congestive heart failure, and pulmonary hypertension), abnormalities of the neurologic system (central nervous system [CNS] infections, hypoxic injury, and hydrocephalus), blood dyscrasia (anemia or polycythemia), infections, metabolic problems, or exposure to maternal drugs. Specific treatment should be aimed at the underlying etiology. Screening laboratory tests and imaging studies such as complete blood count (CBC), C-reactive protein (CRP), blood gas, and chest X-ray should be initiated if indicated.

Clear evidence exists that either insufficient or excessive oxygen can be harmful to the newborn infant. Hypoxia/ischemia can result in injury to the brain and other organs, but even brief exposure to excessive oxygen can result in vision or other neurological problems. Newborn babies do not have high levels of blood oxygen until at least 10 minutes after birth, so pulse oximetry below 90 is completely normal just after birth. Also, because of circulation changes at birth, cyanosis does not always mean lack of oxygen and a pink color of the skin does not always mean presence of oxygen in the newborn. When using pulse oximetry on the newborn in the first hour of life, it should be applied preductally (the right, upper extremity).

### Table 31.4 Factors Associated with Delayed Transition

<table>
<thead>
<tr>
<th>Infant</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Hypothermia</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hypoglycemia</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Retained pulmonary fluid</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Preterm birth (&lt;37 weeks)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Multiple births</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Small or large for gestational age (&lt;2500 g or &gt;4200 g)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Postterm birth (≥42-week gestation)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Meconium</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low Apgar scores (i.e., &lt;7 at 5 minutes)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Congenital problems (i.e., chromosomal abnormalities and structural problems)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Infection</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mother</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diabetes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hypertension or preeclampsia</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tocolysis</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Licit or illicit drug</td>
<td></td>
<td></td>
</tr>
<tr>
<td>No prenatal care</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age greater than 35 years</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### Table 31.5 Hypoglycemia Management in the Child after 34 Weeks of Gestation at Risk (34–36 Weeks' Gestation, Small for Gestation Age, Infants of Diabetic Mothers, and/or Large for Gestational Age)

<table>
<thead>
<tr>
<th>Symptomatic* with Serum Glucose &lt;40 mg/dL</th>
<th>Asymptomatic (4–24 Hours of Age)</th>
<th>Asymptomatic (Birth to 4 Hours of Age)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Use IV glucose</td>
<td>Initial feed within 1 hour. screen glucose 30 minutes after first feed. If initial screen is &lt;25 mg/dL, feed and recheck in 1 hour. If repeat glucose is still &lt;25 mg/dL, start IV glucose. If 25–40 mg/dL, refeed and use IV glucose as needed. If repeat glucose, use IV glucose as needed.</td>
<td>Continue feeds every 2–3 hours. Screen glucose prior to each feed. If screen &lt;35 mg/dL, feed and recheck in 1 hour. If repeat glucose is still &lt;35 mg/dL, start IV glucose. If repeat glucose 35–45 mg/dL, refeed and glucose as needed.</td>
</tr>
</tbody>
</table>

*Symptomatic hypoglycemia: irritability, tremors, jitteriness, exaggerated reflexes, high-pitched cry, seizures, lethargy, floppiness, cyanosis, apnea, and poor feeding.

**Note:** Target glucose >45 mg/dL before routine feedings. IV infusion, if needed, at 5–8 mg/kg/minute (80–100 mL/kg/24 hours).
clear posting of the new protocol in every venue where neonates are resuscitated is of the utmost importance.

**Transient Tachypnea of the Newborn**

In the newborn, tachypnea without respiratory distress may be a benign condition called transient tachypnea of the newborn (TTN). These infants are usually term or near term and will develop tachypnea following birth. They may have respiratory rates up to 120 breaths/minute, nasal flaring, grunt ing, and retractions, but upon physical examination, they do not appear to be in serious distress. Risk factors include cesarean delivery, precipitous delivery, fetal polycythemia, and delivery to a diabetic mother. There is no clear etiology to this disorder, but it is thought to be due to a combination of delayed resorption of fetal lung fluid from the lymph system, pulmonary immaturity, and a mild surfactant deficiency. Unfortunately, TTN is not easily distinguishable from other causes of tachypnea, such as pneumonia or sepsis. If the condition does not quickly resolve, these infants should have screening laboratory tests, including a CBC, CRP, and blood culture, along with a chest X-ray and a pediatric or neonatal consult. In TTN, the lab work will be normal and the chest X-ray will have increased pulmonary vascular markings and fluid in the fissures. More seriously affected infants may be treated with broad-spectrum antibiotics until infection can be excluded. In addition, those infants with respiratory rates above 60 breaths/minute should not be fed by mouth, and some will require either nasogastric (NG) tube feeds or IV fluids if the tachypnea lasts more than 12 hours.

**Common Malformations Affecting Transition**

Some congenital abnormalities of the newborn can present as problems in the delivery room. Below is a list of common malformations and best stabilization practices before transfer to a tertiary care nursery.

2. Pierre Robin sequence (small mandible with relatively large tongue): Secure an oral airway, but consider intubation if inadequate.
3. Congenital diaphragmatic hernia: At birth, immediate insertion of an NG tube to decompress the stomach and intubation of the airway are required.
4. Abdominal wall defects (omphalocele and gastroschisis): Cover the defect with a silo as soon as possible after birth. Gastroschisis may require prompt surgical correction; therefore, surgery should be called as soon as the baby is born.
5. Neural tube defects: Best emergent management is controversial, but the goal is to keep the area moist and free of contamination. The area may be covered with sterile plastic wrap, or the baby may be placed in a sterile plastic bag to just above the lesion.

**Withholding or Discontinuing Resuscitation**

There are conditions associated with high mortality and poor outcomes in which withholding resuscitation may be appropriate, particularly when there has been an opportunity for parental agreement. Cases should be addressed on an individual basis with a consistent and coordinated approach by both the obstetric and neonatal teams and in consultation with the parents. Cases where noninitiation of therapy could be considered may vary by region and include:

- Gestational age less than 23 weeks
- Birth weight less than 400 g
- Anencephaly
- Some chromosomal abnormalities, such as Trisomy 13

Resuscitation should always be considered in cases with high rates of survival and acceptable rates of morbidity. This generally includes infants who are 25 weeks gestation and above and most infants with congenital abnormalities. In cases of uncertain prognosis, parental desire should be weighed heavily.

It is reasonable to consider stopping resuscitation in infants who are born without a detectable heart rate, if the heart rate remains undetectable at 10 minutes of life.

**TYPE OF NURSERY: LOW- VERSUS ELEVATED-RISK NEONATES**

Minimal intervention is needed for the infant with no risks and a normal delivery. Therefore, it is important to distinguish which infants are at low risk and can go to the well baby nursery, and which infants are at high risk and require further evaluation.

**Low Risk**

These infants can be admitted without pediatric specialist consultation. Low-risk infants are 37 weeks of gestation or older and have birth weight between 2200 and 4200 g, Apgar scores of 7 or above at 5 minutes, normal vital signs, and no signs of congenital anomalies or respiratory distress.

**Elevated Risk**

Elevated-risk infants require pediatric specialist consultation prior to admission to the well baby nursery. Criteria include infants at risk for neonatal abstinence syndrome, sepsis, rupture of membranes greater than 16 hours, exposure to group B streptococcus (GBS) that was inadequately treated, maternal temperature of greater than 100.4°F (38°C) within 24 hours of delivery, infants of diabetic mothers, infants born outside of the hospital, infants of uncertain gestation, infants 36 weeks gestation or less, infants less than 2200 g, and infants greater than 4200 g.

**Late-preterm infants** at 34–36 weeks gestation are of often the size and weight of term infants, but are physiologically immature. They are at increased risk for hypothermia, hypoglycemia, dehydration, and poor feeding. The AAP recommends that late-preterm infants receive special attention prior to discharge and close outpatient follow-up in order to prevent rehospitalization [5].
CARE OF THE WELL NEWBORN AFTER LABOR AND DELIVERY (L&D) CARE

All infants must be observed until they complete transition. A skilled staff member must assess the vital signs every 30 minutes for at least 2 hours and observe that the newborn is completely stable on its own. Completion of transition includes thermostability, respiratory rate less than 60 breaths/minute, heart rate greater than 100 beats/minute, and good muscle tone and sucking reflex.

NEWBORN SCREENING

Neonatal screening has arisen from the politics of medicine and of government, and has not been subject to consistent cost-benefit analysis or evidence-based medicine [6-8]. The required tests vary from state to state, but each state requires screening after 24 hours of life. Both the American College of Medical Genetics and Centers for Disease Control (CDC) have made model screening lists, but there is no single list of tests universally required in 2015. The tests suggested are referenced in Table 31.7.

CIRCUMCISION

The health benefits to circumcision include prevention of urinary tract infection (from about 7-14/1000 to 1-2/1000), penile cancer (about 1000 circumcisions needed to prevent one penile cancer), and transmission of sexually transmitted infections (about 15% decrease in lifetime human immunodeficiency virus [HIV] risk). In the United States, the benefits are not considered great enough for universal circumcision of males. The AAP recommends that circumcision should be parental choice and only agreed to after a discussion with a health care provider about the risks and benefits [9]. The World Health Organization encourages circumcision as a method to reduce the risk of acquiring HIV infection in populations at high risk for HIV [10]. Complications of circumcision, which have been reported to range from 0.3% to 20%, include excessive bleeding (1/100-1/1,000), infection (1-6/10,000), penile injury (4/10,000). Circumcision is often a social decision rather than a medical decision. The pediatric practitioner should delay or prevent circumcision in the following cases:

- Suspected sepsis or bacteremia
- Family history of bleeding problems (until the child is fully assessed by hematologist)
- Genital abnormalities, including ambiguous genitalia, micropenis (<2.5 cm from pubic bone to penile tip), hypospadias, epispadias, or chordae.

Performing a Circumcision

Prior to circumcision, infants should have completed transition, voided at least once, and been fasting for 1 hour prior to the procedure. Pain control should be utilized and can include a dorsal penile nerve block, topical analgesic, or subcutaneous ring block. Sucrose water or sweet wine also provides some pain control. Techniques for circumcision include the Mogen™, Plastibell™, and Gomco™. After the procedure, breast- or bottle-feeding is an excellent pain control method. If needed, the dose of acetaminophen is 12–15 mg/kg/dose. A gauze wrap is no longer recommended after the procedure. The area should be kept clean with plain water and covered with lubricant for up to 1 week until completely healed. The

<table>
<thead>
<tr>
<th>Table 31.7</th>
<th>Disorders Recommended by the American College of Medical Genetics Task Force for Inclusion in Newborn Screening</th>
</tr>
</thead>
<tbody>
<tr>
<td>Disorders of organic acid metabolism</td>
<td>Isovaleric acidemia</td>
</tr>
<tr>
<td>Glutaric aciduria type 1</td>
<td>3-Hydroxy-3-methylglutaric aciduria</td>
</tr>
<tr>
<td>Multiple carboxylase deficiency</td>
<td>Methylmalonic acidemia, mutase deficiency form</td>
</tr>
<tr>
<td>3-Methylcrotonyl-CoA carboxylase deficiency</td>
<td>Methylmalonic acidemia, Cb1 A and Cb1 B forms</td>
</tr>
<tr>
<td>Propionic acidemia</td>
<td>Beta-ketothiolase deficiency</td>
</tr>
<tr>
<td>Disorders of fatty acid metabolism</td>
<td>Medium-chain acyl-CoA dehydrogenase deficiency</td>
</tr>
<tr>
<td>Very long chain acyl-CoA dehydrogenase deficiency</td>
<td>Long-chain L-3-hydroxy acyl-CoA dehydrogenase deficiency</td>
</tr>
<tr>
<td>Trifunctional protein deficiency</td>
<td>Carnitine uptake defect</td>
</tr>
<tr>
<td>Disorders of amino acid metabolism</td>
<td>Phenylketonuria</td>
</tr>
<tr>
<td>Maple syrup urine disease</td>
<td>Homocystinuria</td>
</tr>
<tr>
<td>Citrullinemia</td>
<td>Argininosuccinic acidemia</td>
</tr>
<tr>
<td>Tyrosinemia type 1</td>
<td>Hemoglobinopathies</td>
</tr>
<tr>
<td>Sickle cell anemia</td>
<td>Hemoglobin S-b-thalassemia</td>
</tr>
<tr>
<td>Hemoglobin SC disease</td>
<td>Other disorders</td>
</tr>
<tr>
<td>Congenital hypothyroidism</td>
<td>Biotinidase deficiency</td>
</tr>
<tr>
<td>Congenital adrenal hyperplasia</td>
<td>Galactosemia</td>
</tr>
<tr>
<td>Hearing deficiency</td>
<td>Cystic fibrosis</td>
</tr>
</tbody>
</table>


Abbreviations: CoA, coenzyme A; Cb1 A, cobalamin A; Cb1 B, cobalamin B; S-b-thalassemia, sickle beta-thalassemia; SC, sickle cell.

The American College of Medical Genetics Task Force also recommended reporting an additional 25 disorders (secondary targets) that can be detected through screening but that do not meet the criteria for primary disorders (6). At this time, there is a state-to-state variation in newborn screening; a list of the disorders that are screened for by each state is available at http://genes-r-us.uthscsa.edu.

Parents should be told to expect a white yellow discharge for 5–7 days.

CARE OF THE PRETERM INFANT

The delivery of a preterm infant follows many of the guidelines generated for the term infant [1–3]. These infants born at 36 weeks’ gestation or less are at high risk for problems immediately following birth and in the subsequent newborn period. If a preterm delivery is anticipated, the delivery should be attended by a team of neonatal intensive care unit (NICU) personnel (physician, nurse, and respiratory therapist). Management of the infant includes accentuated attention to drying and maintaining thermal stability and close observation of blood glucose, fluid requirements, and respiratory function. Babies less than 29 weeks gestation should be wrapped in polyethylene in order to prevent evaporative heat loss.
In high-risk infants, the prophylactic placement of an umbilical venous line should be considered. Also, early institution of positive airway pressure may be necessary to prevent respiratory failure. Preparation for and anticipation of the delivery with the availability of the proper equipment and personnel for stabilization and transport will optimize survival.

A detailed review of the topic of neonatal care of the preterm infant is beyond the scope of this book. Table 31.8 has recent data on survival at very early (22–28 weeks) gestational ages, helpful for obstetric and neonatal counseling [11].

### NEONATAL ENCEPHALOPATHY AND CEREBRAL PALSY

#### Neonatal Encephalopathy

Neonatal encephalopathy (NE) is a clinically defined syndrome of disturbed neurologic function in the earliest days of life in the term infant, manifested by difficulty initiating and maintaining respiration, depression of tone and reflexes, subnormal level of consciousness, and, often, seizures. It can result from a myriad of conditions and may or may not result in permanent neurologic impairment [12,13].

#### Hypoxic–Ischemic Encephalopathy

Hypoxic–ischemic encephalopathy (HIE) is a subset of neonatal encephalopathy for which the etiology is considered to be a limitation of oxygen and blood flow near the time of birth. A comprehensive assessment of neonatal status, maternal history, obstetric antecedents, intrapartum factors, and placental pathology is required in order to determine if hypoxic–ischemia occurred in close temporal proximity to L&D. The following conditions increase the likelihood that hypoxia–ischemia played a role in causing NE [14]:

- Apgar score of 6 or lower at 5 minutes
- Fetal umbilical artery pH <7.0 or base deficit ≥12 mmol/L
- Magnetic resonance (MR) imaging or MR spectroscopy showing hypoxic–ischemic pattern of cerebral imaging (deep nuclear gray matter or watershed cortical injury)
- Multisystem organ failure
- Sentinel hypoxic/ischemic event immediately before or during labor/delivery (uterine rupture, umbilical cord prolapse, maternal cardiovascular collapse, fetal exsanguination)

Therapeutic Hypothermia

Induced therapeutic hypothermia is recommended for infants 36 weeks and greater with evidence of moderate-to-severe HIE. Several randomized controlled trials have demonstrated lower mortality and less neurodevelopmental disability at 18 months in babies who were not cooled. Therapeutic hypothermia should be administered at institutions with multidisciplinary care and capability for long-term follow-up [3].

### Cerebral Palsy

Cerebral palsy is a chronic static neuromuscular disability characterized by aberrant control of movement or posture, appearing early in life and not the result of recognized progressive disease. Cerebral palsy affects 2/1000 live births.
Prevention is elusive. The majority of cases of cerebral palsy do not result from isolated intrapartum asphyxia with resultant hypoxemia and organ damage [13]. Other factors associated with cerebral palsy and/or neonatal encephalopathy are as follows:

- Increasing maternal age
- Family history of seizures or neurologic disease
- Maternal history of infertility treatment or previous neonatal death
- Severe preeclampsia
- Intrauterine growth restriction
- Congenital malformations or genetic abnormalities
- Autoimmune and coagulation disorders
- Infections

Timing of Neonatal Encephalopathy

Estimating the time when neonatal encephalopathy occurred is extremely difficult. Some [12] have reported risk factors related to this condition:

- 69% Only antepartum risk factors
- 25% Both antepartum risk factors and evidence for intrapartum hypoxia
- 4% Intrapartum hypoxia without antepartum risk factors
- 2% No recognized risk factors

Therefore, the incidence of neonatal encephalopathy attributed to intrapartum hypoxia in absence of other preconception or antepartum abnormalities is about 1.6/10,000 infants [12,13].

Seventy-five percent of children with cerebral palsy had normal Apgar scores at birth [12] (see Table 31.9).

Criticism of the management of labor should not be confused with cerebral palsy causation because the two often may not be linked.

### Table 31.9 Apgar Scores and Cerebral Palsy

<table>
<thead>
<tr>
<th>Apgar Score</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>0–3 at 5 minutes</td>
<td>0.3%–1.0% of babies with cerebral palsy</td>
</tr>
<tr>
<td>0–3 at 10 minutes</td>
<td>10% of babies with cerebral palsy (but rates drop to 5% if scores improve at 15 and 20 minutes)</td>
</tr>
<tr>
<td>&lt;3 at 15 minutes</td>
<td>53% mortality, 36% of survivors with cerebral palsy</td>
</tr>
<tr>
<td>&lt;3 at 20 minutes</td>
<td>60% mortality, 57% of survivors with cerebral palsy</td>
</tr>
</tbody>
</table>

REFERENCES

The adnexal mass

Lauren Cooper Hand, George Patounakis, and Norman G. Rosenblum

KEY POINTS
- There are no randomized trials on interventions for adnexal masses in pregnancy.
- Ultrasound, with transvaginal and Doppler capabilities, is the mainstay of diagnosis and prognosis.
- Most adnexal masses are simple cysts and will resolve.
- The most common persistent adnexal mass in pregnancy is a mature teratoma.
- The risk of an adnexal mass representing malignancy is 5%. Half of all ovarian cancers in pregnancy are epithelial and half of those are of low malignant potential.
- For a persistent adnexal mass in pregnancy, early preoperative consultation with a gynecologic oncologist, anesthesiologist, and neonatologist is recommended.
- Complications related to the persistent adnexal mass in gravid patients may include severe pain (5%–26%), ovarian torsion (7%–12%), cyst rupture (9%), pelvic impaction and obstruction of labor (17%), and ovarian cancer (5%).
- CA-125 is nonspecific and can be elevated simply due to pregnancy; between 15 weeks and delivery, levels between 1,000 and 10,000 units/mL cannot be attributed to pregnancy alone.
- Management in the first trimester is almost uniformly expectant unless the clinical presentation is acute or malignancy is suspected.
- Although there are no guidelines, surgery in pregnancy can be reserved for adnexal masses that persist in the second trimester, and are either ≥10 cm, symptomatic, or have solid or mixed solid and cystic findings concerning for malignancy.
- When necessary and feasible, surgery should be scheduled in the early second trimester when organogenesis is complete, most spontaneous abortions have occurred, and the risk for premature delivery is low.
- Intervention should be considered at any point in gestation if a mass is complex or suspicious for malignancy and increases in size.
- If a suspicious adnexal mass is identified incidentally at the time of cesarean section, it should be removed and not simply aspirated.
- If a malignant neoplasm of the ovary is found at the time of exploration, the surgeon should consult a gynecologic oncologist to properly stage the disease.
- For more advanced ovarian cancer, the degree of cytoreductive surgery and the timing of initiation of chemotherapy will depend on fetal viability and maternal choice.

DEFINITION/DIAGNOSIS
An adnexal mass is any mass in the ovary or tube or attached to them (adnexa). There is an increase in the detection of asymptomatic adnexal masses in pregnancy due to the increase in prenatal ultrasounds. The vast majority (>90%) of adnexal masses in pregnancy are ovarian. Most are benign simple cysts under 5 cm. Table 32.1 outlines the differential diagnosis of an adnexal mass during pregnancy [1]. The diagnosis is most accurately made by ultrasound, even if it is possible to diagnose an adnexal mass by bimanual physical examination. A persistent adnexal mass is one that does not resolve by the second trimester.

EPIDEMIOLOGY/INCIDENCE
In earlier studies, approximately 1%–4% of women were diagnosed with an adnexal mass in pregnancy with persistence of only 1 in 200 to 1 in 600 into the second trimester [2,3]. A more recent study of adnexal masses that performed ultrasound of asymptomatic women during nuchal translucency evaluation (11–14 weeks) found an incidence of almost 25%, with 85% of those resolving during the pregnancy without intervention [4]. This dramatic increase in incidence is likely secondary to the early prevalence of functional cysts that spontaneously resolve during the pregnancy. Meanwhile, a recent study looking at over 7.7 million deliveries from 2003 to 2011, showed the incidence of ovarian masses to be 0.25% at the time of delivery [5]. The majority of the adnexal masses are simple cysts such as corpus luteal or other functional cysts. Most of them (up to 90%) when identified during pregnancy will spontaneously regress prior to the second trimester. The likelihood of regression is inversely related to size. Only 6% of cysts ≤6 cm compared with 39% of cysts >6 cm persist into the second trimester of pregnancy [6–8]. Two adnexal conditions are specifically associated with pregnancy and spontaneously regress in the postpartum period requiring no further treatment—luteomas of pregnancy and theca lutein cysts. Only 5% of adnexal masses found in the beginning of pregnancy are malignant ovarian tumors [3]. The incidence of ovarian cancer in pregnancy is rare, 1 in 18,000 to 1 in 47,000 deliveries [9,10]. It is the fifth most common cancer diagnosed in pregnancy [11].

CLASSIFICATION
Among persistent adnexal masses diagnosed during pregnancy (age group 18–35 years), mature teratomas are the most common followed by benign serous and mucinous cystadenomas (Table 32.2) [12]. Most of the literature on ovarian cancer in pregnancy is based on case reports and series. In one of the largest case series, the most common ovarian cancers found in pregnancy are serous and mucinous tumors of low malignant potential [1].

RISK FACTORS/ASSOCIATIONS
Maternal age is a risk factor for malignancy, but ovarian malignancies can occur at any reproductive age. A study assessing the accuracy of combining patient demographics, serum CA-125,
Table 32.1  Differential Diagnosis of Most Common Adnexal Masses in Pregnancy

<table>
<thead>
<tr>
<th>Benign</th>
<th>Malignant</th>
</tr>
</thead>
<tbody>
<tr>
<td>Corpus luteum</td>
<td>Germ cell tumor</td>
</tr>
<tr>
<td>Simple cyst</td>
<td>Cystadenocarcinoma</td>
</tr>
<tr>
<td>Hemorrhagic cyst</td>
<td>Borderline tumors</td>
</tr>
<tr>
<td>Dermoid</td>
<td>Sex cord stromal tumor</td>
</tr>
<tr>
<td>Endometrioma</td>
<td>Metastatic cancer to ovary</td>
</tr>
<tr>
<td>Myoma</td>
<td></td>
</tr>
<tr>
<td>Luteoma</td>
<td></td>
</tr>
<tr>
<td>Hyperstimulated ovary</td>
<td></td>
</tr>
<tr>
<td>Hydroalpinx</td>
<td></td>
</tr>
<tr>
<td>Theca lutein cyst</td>
<td></td>
</tr>
<tr>
<td>Ectopic/hetertopic pregnancy</td>
<td></td>
</tr>
<tr>
<td>Cystadenoma</td>
<td></td>
</tr>
<tr>
<td>Peritoneal cyst</td>
<td></td>
</tr>
<tr>
<td>Paraovarian/paratubal cyst</td>
<td></td>
</tr>
</tbody>
</table>


Table 32.2  Relative Frequency of Adnexal Masses Diagnosed in Pregnancy

<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>Percent</th>
</tr>
</thead>
<tbody>
<tr>
<td>Benign (95%)</td>
<td></td>
</tr>
<tr>
<td>Dermoid</td>
<td>37</td>
</tr>
<tr>
<td>Serous or mucinous cystadenoma</td>
<td>24</td>
</tr>
<tr>
<td>Corpus luteum</td>
<td>17</td>
</tr>
<tr>
<td>Endometrioma</td>
<td>5</td>
</tr>
<tr>
<td>Leiomyoma</td>
<td>5</td>
</tr>
<tr>
<td>Other (paraovarian, luteoma, theca-lutein)</td>
<td>12</td>
</tr>
</tbody>
</table>

Malignant (5%)                   |         |
| Epithelial                     | 50      |
| Low malignant potential        | 66      |
| Invasive                       | 33      |
| Germ cell tumor                | 30      |
| Sex cord stromal tumor         | 20      |

Source: Adapted from Liu JR et al., Cancer Obstetrics and Gynecology, Lippincott Williams & Wilkins, Philadelphia, PA, 1999.

and ultrasonographic features in predicting the risk of malignancy of an ultrasonographically confirmed adnexal mass in nonpregnant women found age, menopausal status, weight, tumor morphology, presence of ascites, tumor laterality, tumor diameter, and CA-125 level to be associated with risk of malignancy [13]. Specifically, age ≤55, premenopausal weight >200 lb, cystic morphology, negative ascites, unilaterality, diameter <10 cm, and CA-125 <35 were associated with benign masses. None of the cystic tumors in that study were malignant. Furthermore, a large study in postmenopausal women with unilocular cystic masses <10 cm found none to have malignancy [14]. Assisted reproductive technologies increase the incidence of enlarged ovaries, cysts, and therefore adnexal masses.

**COMPLICATIONS**

Many studies show that planned abdominal surgery in pregnancy is safe [11,15–17]. Complications related to the persistent adnexal mass in gravid patients include severe pain (26%), ovarian torsion (1%–22%), cyst rupture (0%–9%), and pelvic impaction and obstruction of labor (2%–17%) [11,15]. Ovarian torsion, the most common significant complication in pregnancy, occurs usually in less than 15% of the cases and is most common in adnexal masses with sizes between 6 and 8 cm [16]. Approximately 60% of torsion happens between 10 and 17 weeks’ gestation and less than 6% happens after 20 weeks’ gestation [16]. Ovarian cancer is estimated to occur in about 5% of adnexal masses found in the beginning of pregnancy [3]. Most malignancies are either germ cell, stromal, or epithelial tumors of low malignant potential [17].

**MANAGEMENT Principles**

There are no trials available to assess any intervention in the management of adnexal masses in pregnancy. As about 90% of these masses resolve and the risk of malignancy is 5%, in general, expectant management with serial ultrasounds is usually appropriate in the vast majority of, but not all, clinical situations. If malignancy is suspected, the best gestational age for surgery is about 16–18 weeks, as the risk of loss is lowest.

**Workup**

**Ultrasound**

Highly skilled ultrasonographers may be able to accurately diagnose complications of adnexal masses in pregnancy (e.g., cancer and torsion) without surgical intervention. Suspicious characteristics of an adnexal mass include complex masses consisting of both solid and cystic components with nodularity, thick septations, irregular borders, solid masses containing irregular echoes, and papillary projections [18]. While cancer can be present in a cyst of any size, adnexal masses of ≤5 cm have an incidence of malignancy less than 1% [11]. Even unilocular cysts ≤10 cm in postmenopausal women can be safely followed by serial ultrasounds until they resolve or develop solid and/or wall abnormalities prompting surgical intervention secondary to a high suspicion of malignancy [14]. In addition to routine ultrasonography, color Doppler studies may be used to distinguish between malignant and benign adnexal masses [19]. Low pulsatility index of <1.0 and low impedance are associated with ovarian neoplasms. Transvaginal ultrasound may also help to better visualize the adnexal mass. The overall sensitivity of high-resolution ultrasound in distinguishing malignant from benign adnexal masses is 96.6%, specificity of 77%, and negative predictive value of 99% [19]. After 20 weeks, adnexal masses are more difficult to see by ultrasound given the larger uterine size. However, definitive diagnosis requires pathologic confirmation.

**Magnetic Resonance Imaging**

Although ultrasound is usually the extent of preoperative imaging, magnetic resonance imaging (MRI) has been used to characterize adnexal masses. There are insufficient data to assess the effect of MRI on management of adnexal masses in pregnancy, but a review of the topic with respect to each type of adnexal mass and the usefulness of MRI has been published [20]. The conclusions are that MRI can assist ultrasound in distinguishing an exophytic leiomyoma, degenerating leiomyoma, an endometrioma, a dermoid cyst, and a decidualized endometrioma, from other masses. Furthermore, it supports the value in MRI of detecting massive ovarian edema and distant findings such as widespread ascites, peritoneal implants, and lymphadenopathy that can help distinguish benign from malignant masses.
Laboratory
Most tumor markers may be elevated in normal pregnancy and are generally not helpful in distinguishing between a benign or malignant ovarian mass in pregnancy. For example, up to 16% of pregnant patients may have an elevated CA-125 [21]. Levels of CA-125 peak in the first trimester (range, 7–251 units/mL), and decrease consistently thereafter. Low-level elevations in pregnancy are typically not associated with malignancy [17]. Levels from 1,000–10,000 units/mL after 15 weeks of gestation are likely to be cancer [3].

Therapy
Treatment planning is dependent upon the timeliness of detection of an adnexal mass in pregnancy. When an adnexal mass is diagnosed in the first trimester, the likelihood of a functional etiology is high, as is the probability of spontaneous resolution. In pregnant women, most simple cysts ≤6 cm have been shown to spontaneously resolve [22–25]. Given the high obstetrical risk during this period, the management in the first trimester is almost uniformly expectant when the clinical presentation is not acute [11]. Similarly, intervention in the third trimester is typically deferred until delivery or the postpartum period as the risk of delaying surgery rarely outweighs the risk of surgery to the mother and the fetus. When necessary and feasible, surgery should be scheduled in the early second trimester after most functional cysts have resolved, organogenesis is complete, most spontaneous losses have occurred, and the risk for premature delivery is low. Progesterone supplementation is necessary if the corpus luteum is removed prior to 8 weeks [3]. Intervention should be considered at any point in gestation if a mass is complex or highly suspicious for malignancy and increases in size (Figure 32.1). In addition to suspected malignancy, surgical intervention may be indicated if torsion, rupture, or hemorrhage is identified.

For persistent adnexal masses in pregnancy, early preoperative consultation with a gynecologic oncologist, anesthesiologist, and neonatologist is recommended [26]. Consultation at <15 weeks is recommended for better operative planning. Masses present after the first trimester that should be resected include those masses with the following characteristics: greater than 10 cm, symptomatic, or are solid or solid and cystic on ultrasound [27].

Surgery
Removal of adnexal masses can be accomplished via laparoscopy, laparotomy, or at time of cesarean section. When exploration is necessary, all efforts should be made to avoid unnecessary manipulation of the uterus to minimize premature uterine contractions. Other intraoperative and postoperative considerations should be kept in mind when operating on a pregnant patient (Table 32.3). Pre- and postoperative fetal doppulons are recommended in the first trimester or early second trimester. Continuous fetal heart monitoring can be considered when viability is reached with the ability for an emergent cesarean section if needed [28–30].

While traditionally a laparotomy was the standard recommendation used to explore the abdomen of a pregnant patient with an adnexal mass, laparoscopy is currently not only an acceptable alternative (in experienced hands) but also is preferred in cases with low suspicion of malignancy, as it offers the benefit of a more expeditious recovery [31–33]. In a Cochrane review on laparoscopy for adnexal tumors in pregnancy, the available case series were too limited for conclusions to be made on risks and benefits of laparoscopy. The need for a randomized controlled study was concluded [34]. However, laparoscopic surgery can be safely performed for ovarian torsion during pregnancy [35,36]. Abdominal surgery during pregnancy, in particular laparotomy, has been associated with higher rates of miscarriage and preterm birth compared with no surgery [11].

If a suspicious adnexal mass is identified incidentally at the time of cesarean section, it should be removed and not simply aspirated [2]. With aspiration and cytologic evaluation, malignancy could be missed [37].

If a malignant neoplasm of the ovary is suspected prior to surgery, a vertical incision is preferred and a frozen section should be sent. The surgeon’s obligation is to consult a gynecologic oncologist in order to properly stage the disease (Table 32.4).

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Stage I—Growth limited to the ovaries or fallopian tubes
  - Stage IA—Growth limited to one ovary or fallopian tube, no ascites, no tumor on external surface, and capsule intact
  - Stage IB—Growth limited to both ovaries or fallopian tubes, no ascites, no tumor on external surface, and capsule intact
  - Stage IC—Tumor limited to one or both ovaries or fallopian tubes with any of the following:
    - Stage IC1—Surgical spill intraoperatively
    - Stage IC2—Capsule ruptured before surgery or tumor on ovarian or fallopian tube surface
    - Stage IC3—malignant cells in the ascites or peritoneal washings
Stage II—Growth involving one or both ovaries, with pelvic extension (below pelvic brim) or peritoneal cancer
  - Stage IIA—Extension and/or implants to the uterus and/or fallopian tubes and/or ovaries
  - Stage IIB—Extension to other pelvic intraperitoneal tissues
Stage III—Tumor involving one or both ovaries, fallopian tubes, or peritoneal cancer, with cytologically or histologically confirmed peritoneal implants outside the pelvis and/or metastasis to the retroperitoneal lymph nodes
  - Stage IIIA—Metastasis to the retroperitoneal lymph nodes with or without microscopic peritoneal involvement beyond the pelvis
  - Stage IIIA1—Positive retroperitoneal lymph nodes only (cytologically or histologically proven)
    - Stage IIIA1(i)—Metastasis ≤10 mm in greatest dimension
    - Stage IIIA1(ii)—Metastasis >10 mm in greatest dimension
  - Stage IIIB—Macroscopic peritoneal metastases beyond the pelvic brim ≤2 cm in greater dimension, with or without metastasis to the retroperitoneal lymph nodes
  - Stage IIIC—Macroscopic peritoneal metastases beyond the pelvic brim >2 cm in greater dimension, with or without metastasis to the retroperitoneal lymph nodes
Stage IV—Distant metastases excluding peritoneal metastases
  - Stage IVA—Plural effusion with positive cytology
  - Stage IVB—Metastases to extra-abdominal organs (including inguinal lymph nodes and lymph nodes outside the abdominal cavity)


Since most present as stage I disease, a unilateral salpingo-ophorectomy, omentectomy, and limited pelvic and paraaortic lymph node dissection is the procedure of choice. All suspicious lesions should be biopsied, with surface excrescences and ascites being suggestive of malignancy. If benign disease is suspected, cystectomy is preferred over a salpingo-oophorectomy.

If the disease appears to be confined to the pelvis, comprehensive surgical staging is indicated. The staging procedure includes peritoneal cytology, multiple peritoneal biopsies, omentectomy, and pelvic and paraaortic lymph node sampling. Rarely is a hysterectomy indicated. For more advanced disease, cyto-reductive surgery should be attempted. The timing of initiation of chemotherapy will depend on fetal viability and maternal choice, and should be managed by a gynecologic oncologist. Returning for staging after completion of pregnancy is not thought to adversely impact survival, though late stage of disease has poor overall survival.

REFERENCES


Cervical cancer screening and management in pregnancy

Talia M. Maas and Christine H. Kim

KEY POINTS
- Cervical cancer screening guidelines for nonpregnant women can be followed in pregnant women. These suggest initiating Pap smear screening at age 21, regardless of onset of sexual activity. Routine screening intervals have also been extended to every 3 years for women in their twenties without human papillomavirus (HPV) cotesting and every 5 years in women over 30 with the addition of HPV cotesting.
- The only cervical diagnosis that is considered to alter management in pregnancy is invasive cancer.
- A more conservative approach should be used to manage cervical intraepithelial neoplasia (CIN).
- Due to risks of bleeding and premature rupture of membranes, endocervical curettage (ECC) is never recommended in pregnancy.
- Diagnostic conization during pregnancy should only be considered when either the biopsy, or cytology is suggestive of invasive cancer and the diagnosis of invasion would result in a modification of treatment recommendations, timing, or mode of delivery.
- In the diagnosis of invasive cancer with a grossly visible lesion, cesarean delivery is indicated and results in better survival outcomes as well as a decreased risk of obstetrical bleeding in the intrapartum and postpartum periods.
- If a nonvisible microinvasive lesion (stage IA1 or IA2) is identified, either the abdominal or vaginal route of delivery is acceptable, depending on obstetrical and gynecologic circumstances.
- Once the diagnosis of cervical cancer is established, individualized recommendations for the management of the malignancy as well as the pregnancy are formulated with consideration for the stage of disease, gestational age at the time of diagnosis, and maternal desires regarding the continuation of her pregnancy.
- Fertility-sparing surgery and neoadjuvant chemotherapy for earlier stage cervical cancer found in pregnancy may be considered without worsening survival.

DIAGNOSIS/DEFINITION
Microinvasive cervical cancer is defined as cancer spread to no more than 5 mm into the tissue of the cervix. Invasive cervical cancer is defined as cancer spread from the surface of the cervix to tissue deeper in the cervix, possible spread to part of the vagina, to the lymph nodes, to the other tissues surrounding the cervix, within the pelvis, or beyond the pelvic areas into nearby organs.

EPIDEMIOLOGY/INCIDENCE
The overall rate of an abnormal Pap smear with atypical squamous cells—undetermined significance (ASC-US) or higher in the United States is estimated around 4% from data collected between 2003 and 2010 [4]. The peak age incidence of cervical cancer is in the mid-forties [5]. In low- and middle-income countries, cervical cancer is the second most common cancer among women, the third most common cause of cancer-related death, and the most common cause of mortality from gynecologic malignancy. In contrast, in high-income countries, the success of Pap smear screening has greatly reduced the incidence of disease by accurately detecting preinvasive and early-invasive cervical disease. In the United States, the incidence of cervical cancer ranges from 1.5 to 12 cases in 100,000 pregnancies [6]. About 1% of the women who have cervical cancer are pregnant at the time of diagnosis. The likelihood that a pregnant woman with ASC pathology has a detectable high-risk human papillomavirus (HR HPV) is 84% [7].

CLASSIFICATION
The Pap smear report consists of the following [2]:

Specimen Type
- Conventional Pap smear: Cells from the transformation zone are sampled and transferred directly onto a slide [8].
- Liquid-based cytology: Cells from the transformation zone are placed in a prepared liquid preservative and processed by a laboratory. This is the method most commonly used in clinical practice in the United States because of the added benefit of using a single sample for HPV cotesting. The use of liquid-based media can also control for obscuring factors including blood, inflammation, and other processes [8].
- There is no difference in unsatisfactory rates, cytology classifications, and accuracy between conventional and liquid-based cervical cytology [9].

Specimen Adequacy
Satisfactory for Evaluation
- Defined as a minimum of 5,000 well-visualized squamous cells on a liquid-based preparation or 8,000 to 12,000 well-visualized squamous cells on a conventional smear.
The presence of endocervical cells indicates that the area at risk for neoplasia, the transformation zone, has been adequately sampled [10]. Samples reported as negative but with absent or insufficient endocervical/transition zone components have raised concern about missed pathology. Prior guidelines have recommended repeat cytology. Recent meta-analysis studies reported that negative cytology (with or without sufficient sampling of the endocervical/transition zone) has both good negative predicative value as well as good specificity. Current guidelines do not recommend repeat follow-up Pap smears. Women ages 21–29 are to continue with routine screenings. Women older than 30 years of age are recommended to undergo HPV cotesting for an added margin of safety. Alternatively if cotesting is unavailable, patients are to undergo repeat testing in 2–4 months. If the subsequent Pap smear is unsatisfactory, evaluation with colposcopy and/or biopsies is appropriate [11].

Unsatisfactory for Evaluation
- Defined as more than 75% of the cells being uninterpretable or an unlabeled specimen.
- Since women with this result are more likely to have intraepithelial lesions or cancer on follow-up than women with satisfactory Pap smears [10], a repeated Pap smear in 2–4 months is recommended. If the subsequent Pap smear is unsatisfactory, evaluation with colposcopy and/or biopsies is appropriate [11].

Interpretation/Result
Squamous Epithelial Cell Abnormalities
- ASC of either of the following:
  - Undetermined significance (ASC-US)
  - Suspicious for high-grade squamous intraepithelial lesions (HSILs or ASC-H)
- Low-grade squamous intraepithelial lesions (LSIL)
- Changes consistent with HPV, mild dysplasia, or grade 1 CIN I
- HSIL
  - HSIL includes moderate or severe dysplasia, CIN II, CIN III, carcinoma in situ (CIS), and should indicate if there are features suspicious for invasive disease. When glandular cell abnormalities are present, it should be noted whether there are changes favoring neoplasia
  - Carcinoma

Glandular Cell Abnormalities
- Atypical glandular cells (AGCs) may be of endocervical, endometrial, or other glandular origin
- Endocervical adenocarcinoma in situ (AIS)
- Adenocarcinoma

Human Papillomavirus Testing
HPV testing is now an integral part of cervical cancer screening. Updated ASCCP Consensus guidelines that were most recently validated in 2015 state HPV cotesting is recommended for all female patients over the age of 30 and for all patients with ASC-US cytology.
- HPV infection is the leading etiologic agent in the development of premalignant and malignant lower genital tract disease [12].
- Cotesting should only detect for the presence of high-risk HPV species. There is no role in testing for low-risk genotypes. Detecting low-risk genotypes has been proven to cause unnecessary procedures and testing, therefore decreasing clinical specificity. It is recommended to use only Food and Drug Administration (FDA)-approved HPV DNA detection kits [8].
- The most well-studied HPV test is the Hybrid Capture 2 HPV DNA Assay (Digene Corporation, Beltsville, MD), which uses a probe mix for high-risk (HR) HPV types 16, 18, 31, 33, 35, 39, 45, 51, 52, 56, 58, 59, and 68 [13].
- A newer assay for detecting HR HPV was approved for FDA use in 2009. Cervista™ HR HPV (Hologic Corporation, Waltham, MA) includes the 13 HPV types above and adds type 66 [8].
- Cervista HPV 16/18 is another diagnostic test that specifies type 16 and 18. These two represent the most common types found in cervical squamous cell carcinomas and adenocarcinomas, respectively.
- Ongoing research looks promising in improving specificity and positive predictive value of screening for CIN2+ by looking at messenger RNA (mRNA) instead of DNA for HR HPV [14].

MANAGEMENT
Pregnancy Considerations
Pregnancy-induced changes in the cervix include hyperemia, eversion of columnar epithelium, more prominent glands, and increased production and volume of mucus. The decidual changes may exaggerate the colposcopic appearance of CIN. A biopsy during pregnancy may cause substantial bleeding [3]. In pregnancy, the general philosophy for the treatment of intraepithelial neoplasia of the cervix has become expectant management after careful diagnosis.

Screening in Pregnancy
The Pap smear is used to screen for cellular abnormalities that are associated with an increased risk for the development of cervical cancer. It selects those women who should have further evaluation, such as HPV DNA testing, colposcopy, and/or biopsy, which then are used for treatment decisions. The National Comprehensive Cancer Network (NCCN) panel has adopted recommendation set forth by the American Cancer Society on initiation and frequency of Pap smear [15]. New recommendations for 2012 include initiating Pap smear screening at age 21, regardless of onset of sexual activity. Routine screening intervals have also been extended to every 3 years for women in their twenties and should not include routine HPV cotesting unless abnormal cytology is detected. Women greater than 30 years of age should undergo screening every 5 years with both cytology and HPV cotesting. If HPV cotesting is unavailable, cytology alone every 3 years is recommended. The rationale is that invasive cancer is rare in women under 21 years. The evidence for increasing the interval is based on studies that suggested no statistical difference between annual surveillance and 2- to 3-year intervals.

Annual surveillance is recommended for patients with known immunosuppression from HIV or organ transplantation, women exposed to diethylstilbestrol in utero, or those who have been previously treated for CIN 2, CIN 3, or cervical cancer.

Pap smears are often obtained at the first prenatal visit by many providers, but the guidelines for nonpregnant women can be followed in pregnant women [8].
Consultations Required
Consultation with gynecologic oncologists is recommended in cases of cervical cancer [16]. Consultation should occur as early as possible after the diagnosis of cervical cancer for better therapeutic planning. Collaboration with a Maternal-Fetal Medicine specialist is also recommended for management of obstetrical complications and delivery planning.

Indications for Colposcopy and Cervical Biopsy during Pregnancy
As in nonpregnant women, women with abnormal Pap smears during pregnancy should undergo colposcopy with directed biopsies of suspicious areas to rule out invasive disease (Figure 33.1) [3].

ASC-US, HPV negative
• No indication for colposcopy at any age. Pregnant patients have to follow age-appropriate routine screening.

LSIL and ASC-US, HR HPV positive
• Ages 21–24 years: women are to undergo repeat Pap smear and cotesting in 12 months. Colposcopy is not recommended.
• Ages >24 years: In pregnant patients, colposcopy is recommended but it is acceptable to defer colposcopy to 6 weeks postpartum.

HSIL or more invasive cytology
• In patients with higher grade lesions noted on cervical screening examinations necessary colposcopically directed cervical biopsies can be safely performed at any time during pregnancy. Many defer biopsy until the second trimester when the risk of incidental pregnancy loss is minimal. Due to risks of bleeding and premature rupture of membranes, ECC is never recommended in pregnancy. Cervical biopsies should only be performed if CIN II or above is suspected on colposcopic examination.

CIN1:
• If adequate samples are obtained during colposcopy and invasive disease is not suspected at any point during the pregnancy, it is recommended to defer repeat colposcopy to 6 weeks postpartum.

CIN2/3 and carcinoma in situ (CIS):
• In women with histological diagnosis of CIN2/3 or (CIS) on initial colposcopy, it is recommended to perform repeat colposcopy and cytology every 12 weeks for the duration of the pregnancy. Repeat biopsy is recommended only if the appearance of the lesions worsens or if cytology is concerning for invasive cancer. It is acceptable to defer tissue biopsy to 6 weeks postpartum if no progression of the lesion is noted on colposcopy and cytology [3].

Diagnosis of Invasive Cancer
Once the diagnosis of cervical cancer is established, individualized recommendations for the management of the malignancy as well as the pregnancy are formulated with consideration for the stage of disease, gestational age at the time of diagnosis, and maternal desires regarding the continuation of her pregnancy (see Chapter 42 in Maternal-Fetal Evidence Based Guidelines). The patient should seek consultation by specialists in both maternal-fetal medicine as well as gynecology/oncology.

Management of the Abnormal Pap Smear
There are no randomized control trials assessing any aspect of management of abnormal Pap smear in pregnancy. Most of the recommendations are based on expert opinion and anecdotal experience [16–18]. The only diagnosis that is considered to alter management in pregnancy is invasive cancer.

Management of abnormal Pap smears in pregnancy should follow the recommendations delineated in Figure 33.1. Recommendations for colposcopy in pregnant patients are similar to those in the nongravid state, with certain exceptions.

Figure 33.1 Algorithm for the workup of abnormal cytology in pregnancy. AGC, atypical glandular cell; AIS, adenocarcinoma in situ; ASC-H, atypical squamous cells higher; CIN, cervical intraepithelial neoplasia; HSIL, high-grade squamous intraepithelial lesion.
as listed in Table 33.1. Newer guidelines for colposcopy in pregnancy state that colposcopy is preferred for ASC-US associated with positive HR HPV type or LSIL Pap smears, but also that colposcopy can be deferred by the clinician until after pregnancy for these two categories. Colposcopy during pregnancy has as its primary goal to assess for invasive cancer. Biopsies should only be performed for women with colposcopic impression of CIN 3/CIS, AIS, or cancer, with the purpose to exclude invasive cancer [19].

Repeat Pap and colposcopy is recommended if invasive cancer is suspected but not yet proven, specifically with evidence of CIN2/3 or CIS on cytology or biopsy. Repeat biopsy during pregnancy should only be performed if progression of lesion is suspected. ECC is not performed in pregnancy [14,19].

Conization performed during pregnancy is associated with increased morbidity. The most common complications include hemorrhage, abortion, premature delivery, and infection [20]. Therefore, cervical conization has limited indications in pregnancy (Table 33.2). In general, diagnostic conization during pregnancy should only be considered when either the biopsy or cytology is suggestive of invasive cancer or AIS and the diagnosis of invasion would result in a modification of treatment recommendations, timing, or mode of delivery [19].

Unlike standard recommendations for cervical conization in nonobstetrical patients with inadequate colposcopic biopsies or discordance between Pap smears and colposcopic biopsies [12], pregnant women with these findings can defer further examination until after pregnancy if invasive cancer has been ruled out. If a cervical conization must be performed during pregnancy, this procedure should ideally be performed in the early second trimester.

Management of Cervical Cancer
For staging and management please refer to Table 33.3 and Chapter 42 in Maternal-Fetal Evidence Based Guidelines [10] (Figure 33.1). Microinvasive cervical cancer is defined as cancer spread to no more than 5 mm deep into the cervical stroma. Microinvasive disease includes stages IA1 and IA2. Invasive cervical cancer is defined as cancer that has invaded deeper than 5 mm into the cervical stroma, is grossly visible, or involves additional structures. Invasive disease includes Stage IB1 or greater.

Stage IA1:
- Conization is recommended for both treatment and diagnosis in patients with evidence of Stage IA1 cancer [21]. To minimize the risk of spontaneous abortion it is optimal to perform conization in the early second trimester [20].
- If conization margins are negative, patients can be managed expectantly for the duration of the pregnancy [20].
- Either abdominal or vaginal delivery routes are expectable for these patients. Mode of delivery should be discussed with a maternal-fetal medicine specialist with consideration of other gynecological and obstetrical circumstances. This should be decided on an individual basis [21].
- If continued fertility is desired, patients can be followed up every 3 months for 2 years and every 6 months for the next 3 years [20].

Stage IA2
- Conization with the addition of pelvic lymphadenectomy is recommended to rule out high-risk disease. Radical tracheectomy is not recommended given the risk significant blood loss and obstetrical complications resulting from prolonged surgery [21].
- Given the technical complications performing lymphadenectomy after 25 weeks of gestation in patients diagnosed after 25 weeks, it is recommended to delay treatment and full surgical staging until after delivery. If disease progression is suspected, it is recommended to obtain an abdominal and pelvic magnetic resonance imaging (MRI) [20].
- As stated above, patients can undergo either abdominal or vaginal delivery with consideration of other obstetrical circumstances. Delivery can be delayed until fetal maturity at 37 weeks [20].
- Given that IA2 carries a higher risk of nodal metastasis, patients can undergo either classic cesarean delivery with modified radical hysterectomy or surgery can be delayed 6–8 weeks postpartum in the case of vaginal delivery [20].

Stage IB1
- Neoadjuvant chemotherapy is recommended until fetal maturity. The goal of therapy is to stabilize the tumor during pregnancy [21]. Postponement of treatment is acceptable if there is no evidence of progression of disease. As stated above, MRI should be obtained if there is any suspicion. Patients should be examined every 2–4 weeks [20].
- Delivery can be delayed until documented fetal lung maturity with amniocentesis. Testing for fetal lung maturity should begin at 32 weeks [20].
- Patient should undergo classic cesarean delivery with modified radical hysterectomy and pelvic lymphadenectomy [20].

Stage IB2 and higher stage tumors
- Termination of pregnancy is recommended given high risk of disease progression during pregnancy [21].
- If pregnancy is desired, initiation of neoadjuvant chemotherapy is recommended. Chemotherapy allows the pregnancy to be continued to an average of 33 weeks. Early delivery is recommended to expedite the initiation of standard of care treatment. The practitioner should discuss the initiation of betamethasone therapy and preterm delivery [21].
- The patient should undergo a cesarean delivery followed by initiation of therapy 6 weeks postpartum [20].
Recurrence and Follow-Up Treatment
Seventy-five percent of patients with CIN diagnosed during pregnancy have persistent or progressive disease at postpartum evaluation [21]. A regression rate of only 12% is found in pregnant women with CIN III, emphasizing the importance of reevaluation 6 weeks postpartum [12,22].

Routine surveillance can be resumed if there is no recurrence after the first 2 years of follow-up, which includes performing two Pap smears at 6-month intervals or HPV testing at 1-year intervals. Postpregnancy surveillance intervals are modified if the Pap smear or colposcopic biopsies reveal CIN 2, CIN 3, or CIS, and are the same as the nonpregnant guidelines [12].

Prevention
HPV-16 vaccine reduces the incidence of both HPV-16 infection and HPV-16-related CIN [23]. Currently, the three licensed HPV vaccines available in the United States include Cervarix (two-valent, types 16,18), Gardasil (four-valent, types 6,11,16,18), and most recently, Gardasil-9 (nine-valent, types 6,11,18,19,31,33,45,52,58). The nine-valent vaccine was recently found to be equivalent to the prior recommended vaccine [24,25].

No HPV vaccine should not be administered during pregnancy (see Chapter 38 in Maternal-Fetal Evidence Based Guidelines).

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