The American College of Radiology, with more than 30,000 members, is the principal organization of radiologists, radiation oncologists, and clinical medical physicists in the United States. The College is a nonprofit professional society whose primary purposes are to advance the science of radiology, improve radiologic services to the patient, study the socioeconomic aspects of the practice of radiology, and encourage continuing education for radiologists, radiation oncologists, medical physicists, and persons practicing in allied professional fields.

The American College of Radiology will periodically define new practice guidelines and technical standards for radiologic practice to help advance the science of radiology and to improve the quality of service to patients throughout the United States. Existing practice guidelines and technical standards will be reviewed for revision or renewal, as appropriate, on their fifth anniversary or sooner, if indicated.

Each practice guideline and technical standard, representing a policy statement by the College, has undergone a thorough consensus process in which it has been subjected to extensive review, requiring the approval of the Commission on Quality and Safety as well as the ACR Board of Chancellors, the ACR Council Steering Committee, and the ACR Council. The practice guidelines and technical standards recognize that the safe and effective use of diagnostic and therapeutic radiology requires specific training, skills, and techniques, as described in each document. Reproduction or modification of the published practice guideline and technical standard by those entities not providing these services is not authorized.

Revised 2007 (Res. 25)*

ACR–ACOG–AIUM PRACTICE GUIDELINE FOR THE PERFORMANCE OF OBSTETRICAL ULTRASOUND

PREAMBLE

These guidelines are an educational tool designed to assist practitioners in providing appropriate radiologic care for patients. They are not inflexible rules or requirements of practice and are not intended, nor should they be used, to establish a legal standard of care. For these reasons and those set forth below, the American College of Radiology cautions against the use of these guidelines in litigation in which the clinical decisions of a practitioner are called into question.

The ultimate judgment regarding the propriety of any specific procedure or course of action must be made by the physician or medical physicist in light of all the circumstances presented. Thus, an approach that differs from the guidelines, standing alone, does not necessarily imply that the approach was below the standard of care. To the contrary, a conscientious practitioner may responsibly adopt a course of action different from that set forth in the guidelines when, in the reasonable judgment of the practitioner, such course of action is indicated by the condition of the patient, limitations of available resources, or advances in knowledge or technology subsequent to publication of the guidelines. However, a practitioner who employs an approach substantially different from these guidelines is advised to document in the patient record information sufficient to explain the approach taken.

The practice of medicine involves not only the science, but also the art of dealing with the prevention, diagnosis, alleviation, and treatment of disease. The variety and complexity of human conditions make it impossible to always reach the most appropriate diagnosis or to predict with certainty a particular response to treatment. Therefore, it should be recognized that adherence to these guidelines will not assure an accurate diagnosis or a successful outcome. All that should be expected is that the practitioner will follow a reasonable course of action based on current knowledge, available resources, and the needs of the patient to deliver effective and safe medical care. The sole purpose of these guidelines is to assist practitioners in achieving this objective.

I. INTRODUCTION

The clinical aspects contained in specific sections of this guideline (Introduction, Classification of Fetal Sonographic Examinations, Specifications of the Examination, Equipment Specifications, and Fetal Safety) were revised collaboratively by the American College of Radiology (ACR), the American Institute of Ultrasound in Medicine (AIUM), and the American College of Obstetricians and Gynecologists (ACOG). Recommendations for physician qualifications, written request for the examination, procedure documentation, and quality control vary among the three organizations and are addressed by each separately.

This guideline has been developed for use by practitioners performing obstetrical sonographic studies. Fetal ultrasound\(^1\) should be performed only when there is a valid medical reason, and the lowest possible ultrasonic exposure settings should be used to gain the necessary diagnostic information. A limited examination may be performed in clinical emergencies or for a limited purpose such as evaluation of fetal or embryonic cardiac activity, or for other limited purpose.

\(^{1}\)The consensus of the committee was that the use of the terms “ultrasound” or “sonography” is at the discretion of each organization.
fetal position, or amniotic fluid volume. A limited follow-up examination may be appropriate for re-evaluation of fetal size or interval growth or to re-evaluate abnormalities previously noted if a complete prior examination is on record.

While this guideline describes the key elements of standard sonographic examinations in the first trimester and second and third trimesters, a more detailed anatomic examination of the fetus may be necessary in some cases, such as when an abnormality is found or suspected on the standard examination or in pregnancies at high risk for fetal anomalies. In some cases, other specialized examinations may be necessary as well.

While it is not possible to detect all structural congenital anomalies with diagnostic ultrasound, adherence to the following guidelines will maximize the possibility of detecting many fetal abnormalities.

II. CLASSIFICATION OF FETAL SONOGRAPHIC EXAMINATIONS

A. First Trimester Ultrasound Examination

B. Standard Second or Third Trimester Examination

A standard obstetrical sonogram in the second or third trimester includes an evaluation of fetal presentation, amniotic fluid volume, cardiac activity, placental position, fetal biometry, and fetal number, plus an anatomic survey. The maternal cervix and adnexa should be examined as clinically appropriate when technically feasible.

C. Limited Examination

A limited examination is performed when a specific question requires investigation. For example, a limited examination could be performed to confirm fetal heart activity in a bleeding patient or to verify fetal presentation in a laboring patient. In most cases limited sonographic examinations are appropriate only when a prior complete examination is on record.

D. Specialized Examinations

A detailed anatomic examination is performed when an anomaly is suspected on the basis of history, biochemical abnormalities, or the results of either the limited or standard scan. Other specialized examinations might include fetal Doppler, biophysical profile, fetal echocardiogram, or additional biometric measurements.

III. QUALIFICATIONS AND RESPONSIBILITIES OF PERSONNEL

See the ACR Practice Guideline for Performing and Interpreting Diagnostic Ultrasound Examinations.

IV. WRITTEN REQUEST FOR THE EXAMINATION

The written or electronic request for an obstetrical ultrasound examination should provide sufficient information to demonstrate the medical necessity of the examination and allow for its proper performance and interpretation.

Documentation that satisfies medical necessity includes 1) signs and symptoms and/or 2) relevant history (including known diagnoses). Additional information regarding the specific reason for the examination or a provisional diagnosis would be helpful and may at times be needed to allow for the proper performance and interpretation of the examination.

The request for the examination must be originated by a physician or other appropriately licensed health care provider. The accompanying clinical information should be provided by a physician or other appropriately licensed health care provider familiar with the patient’s clinical problem or question and consistent with the state’s scope of practice requirements. (ACR Resolution 35, adopted in 2006)

V. SPECIFICATIONS OF THE EXAMINATION

A. First Trimester Ultrasound Examination

1. Indications

A sonographic examination can be of benefit in many circumstances in the first trimester of pregnancy, including, but not limited to, the following indications:

a. To confirm the presence of an intrauterine pregnancy.
b. To evaluate a suspected ectopic pregnancy.
c. To define the cause of vaginal bleeding.
d. To evaluate pelvic pain.
e. To estimate gestational (menstrual) age.
f. To diagnose or evaluate multiple gestations.
g. To confirm cardiac activity.
h. As an adjunct to chorionic villus sampling, embryo transfer, and localization and removal of an intrauterine device (IUD).
i. To assess for certain fetal anomalies, such as anencephaly, in high risk patients.
j. To evaluate maternal pelvic masses and/or uterine abnormalities.

2 For the purpose of this document, first trimester represents 1w-13w + 6d.

3 For the purpose of this document, the terms “gestational” and “menstrual” age are considered equivalent.
k. To measure the nuchal translucency (NT) when part of a screening program for fetal aneuploidy.
l. To evaluate suspected hydatidiform mole.

Comment
Limited examination may be performed to evaluate interval growth, estimate amniotic fluid volume, evaluate the cervix, and assess the presence of cardiac activity.

2. Imaging parameters

Comment
Scanning in the first trimester may be performed either transabdominally or transvaginally. If a transabdominal examination is not definitive, a transvaginal scan or transperineal scan should be performed whenever possible.

a. The uterus, including the cervix, and adnexa should be evaluated for the presence of a gestational sac. If a gestational sac is seen, its location should be documented. The gestational sac should be evaluated for the presence or absence of a yolk sac or embryo, and the crown-rump length should be recorded, when possible.

Comment
The crown-rump length is a more accurate indicator of gestational (menstrual) age than is mean gestational sac diameter. However, the mean gestational sac diameter may be recorded when an embryo is not identified.

Caution should be used in making the presumptive diagnosis of a gestational sac in the absence of a definite embryo or yolk sac. Without these findings an intrauterine fluid collection could represent a pseudo-gestational sac associated with an ectopic pregnancy.

b. Presence or absence of cardiac activity should be reported.

Comment
With transvaginal scans, cardiac motion is usually observed when the embryo is 5 mm or greater in length. If an embryo less than 5 mm in length is seen without cardiac activity, a subsequent scan at a later time may be needed to assess the presence or absence of cardiac activity.

c. Fetal number should be reported.

Comment
Amnionicity and chorionicity should be documented for all multiple gestations when possible.

d. Embryonic/fetal anatomy appropriate for the first trimester should be assessed.

e. The uterus including the cervix, adnexal structures, and cul-de-sac should be evaluated.

Comment
The presence, location, and size of adnexal masses should be recorded. The presence of leiomyomata should be recorded, and measurements of the largest or any potentially clinically significant leiomyomata may be recorded. The cul-de-sac should be evaluated for the presence or absence of fluid.

f. If possible, the appearance of the nuchal region should be assessed as part of a first trimester scan where a live fetus is present.

Comment
For those patients desiring to assess their individual risk of fetal aneuploidy, a very specific measurement of the NT during a specific age interval is necessary (as determined by the laboratory used). See the guidelines for this measurement below.

NT measurements should be used (in conjunction with serum biochemistry) to determine the risk for having a child with Down syndrome, trisomy 13, trisomy 18, or other anatomical abnormalities such as heart defects. In this setting, it is important that the practitioner measure the NT according to established guidelines for measurement. A quality assessment program is recommended to ensure that false-positive and false-negative results are kept to a minimum.

Guidelines for NT measurement:

i. The margins of the NT edges must be clear enough for proper placement of the calipers.

ii. The fetus must be in the midsagittal plane.
iii. The image must be magnified so that it is filled by the fetal head, neck, and upper thorax.

iv. The fetal neck must be in a neutral position – not flexed and not hyper-extended.

v. The amnion must be seen as separate from the NT line.

vi. The (+) calipers on the ultrasound must be used to perform the NT measurement.

vii. Electronic calipers must be placed on the inner borders of the nuchal space with none of the horizontal crossbar itself protruding into the space.

viii. The calipers must be placed perpendicular to the long axis of the fetus.

ix. The measurement must be obtained at the widest space of the NT.

---

Diagram for the nuchal translucency measurement.
B. Second and Third Trimester Ultrasound Examination

1. Indications

Ultrasound can be of benefit in many situations in the second and third trimester, including, but not limited to, the following circumstances: (adapted from NIH publication 84-667, 1984)

- Estimation of gestational (menstrual) age.
- Evaluation of fetal growth.
- Vaginal bleeding.
- Abdominal or pelvic pain.
- Cervical insufficiency.
- Determination of fetal presentation.
- Suspected multiple gestation.
- Adjunct to amniocentesis or other procedure.
- Significant discrepancy between uterine size and clinical dates.
- Pelvic mass.
- Suspected hydatidiform mole.
- Adjunct to cervical cerclage placement.
- Suspected ectopic pregnancy.
- Suspected fetal death.
- Suspected uterine abnormality.
- Evaluation of fetal well-being.
- Suspected amniotic fluid abnormalities.
- Suspected placental abruption.
- Adjunct to external cephalic version.
- Premature rupture of membranes and/or premature labor.
- Abnormal biochemical markers.
- Follow-up evaluation of a fetal anomaly.
- Follow-up evaluation of placental location for suspected placenta previa.
- History of previous congenital anomaly.
- Evaluation of fetal condition in late registrants for prenatal care.
- To assess for findings that may increase the risk for aneuploidy.
- Screening for fetal anomalies.

Comment
In certain clinical circumstances, a more detailed examination of fetal anatomy may be indicated.

2. Imaging parameters for a standard fetal examination

- Fetal cardiac activity, fetal number, and presentation should be reported.

Comment
Abnormal heart rate and/or rhythm should be reported.

Multiple gestations require the documentation of additional information: chorionicity, amnionicity, comparison of fetal sizes, estimation of amniotic fluid volume (increased, decreased, or normal) on each side of the membrane, and fetal genitalia (when visualized).

b. A qualitative or semiquantitative estimate of amniotic fluid volume should be reported.

Comment
Although it is acceptable for experienced examiners to qualitatively estimate amniotic fluid volume, semi-quantitative methods have also been described for this purpose (e.g., amniotic fluid index, single deepest pocket, two-diameter pocket).

c. The placental location, appearance, and relationship to the internal cervical os should be recorded. The umbilical cord should be imaged, and the number of vessels in the cord should be evaluated when possible.

Comment
It is recognized that apparent placental position early in pregnancy may not correlate well with its location at the time of delivery.

Transabdominal, transperineal, or transvaginal views may be helpful in visualizing the internal cervical os and its relationship to the placenta.

Transvaginal or transperineal ultrasound may be considered if the cervix appears shortened or cannot be adequately visualized during the transabdominal sonogram.

d. Gestational (menstrual) age assessment

First-trimester crown-rump measurement is the most accurate means for sonographic dating of pregnancy. Beyond this period, a variety of sonographic parameters such as biparietal diameter, abdominal circumference, and femoral diaphysis length can be used to estimate gestational (menstrual) age. The variability of gestational (menstrual) age estimations, however, increases with advancing pregnancy. Significant discrepancies between gestational (menstrual) age and fetal measurements may suggest the possibility of fetal growth abnormality, intrauterine growth restriction, or macrosomia.
Comment
The pregnancy should not be redated after an accurate earlier scan has been performed and is available for comparison.

i. Biparietal diameter is measured at the level of the thalami and cavum septi pellucidi. The cerebellar hemispheres should not be visible in this scanning plane. The measurement is taken from the outer edge of the proximal skull to the inner edge of the distal skull.

Comment
The head shape may be flattened (dolichocephaly) or rounded (brachycephaly) as a normal variant. Under these circumstances, certain variants of normal fetal head development may make measurement of the head circumference more reliable than biparietal diameter for estimating gestational (menstrual) age.

ii. Head circumference is measured at the same level as the biparietal diameter, around the outer perimeter of the calvarium. This measurement is not affected by head shape.

iii. Femoral diaphysis length can be reliably used after 14 weeks gestational (menstrual) age. The long axis of the femoral shaft is most accurately measured with the beam of insonation being perpendicular to the shaft, excluding the distal femoral epiphysis.

iv. Abdominal circumference or average abdominal diameter should be determined at the skin line on a true transverse view at the level of the junction of the umbilical vein, portal sinus, and fetal stomach when visible.

Comment
Abdominal circumference or average abdominal diameter measurement is used with other biometric parameters to estimate fetal weight and may allow detection of intrauterine growth restriction or macrosomia.

e. Fetal weight estimation
Fetal weight can be estimated by obtaining measurements such as the biparietal diameter, head circumference, abdominal circumference or average abdominal diameter, and femoral diaphysis length. Results from various prediction models can be compared to fetal weight percentiles from published nomograms.

Comment
If previous studies have been performed, appropriateness of growth should also be reported. Scans for growth evaluation can typically be performed at least 2 to 4 weeks apart. A shorter scan interval may result in confusion as to whether anatomic changes are truly due to growth as opposed to variations in the measurement technique itself.

Currently, even the best fetal weight prediction methods can yield errors as high as ±15 percent. This variability can be influenced by factors such as the nature of the patient population, the number and types of anatomic parameters being measured, technical factors that affect the resolution of ultrasound images, and the weight range being studied.

f. Maternal anatomy
Evaluation of the uterus, adnexal structures, and cervix should be performed when appropriate. If the cervix cannot be visualized, a transperineal or transvaginal scan may be considered when evaluation of the cervix is needed.

Comment
This will allow recognition of incidental findings of potential clinical significance. The presence, location, and size of adnexal masses and the presence of at least the largest and potentially clinically significant leiomyomata may be recorded. It is frequently not possible to image the normal maternal ovaries during the second and third trimesters.

g. Fetal anatomic survey
Fetal anatomy, as described in this document, may be adequately assessed by ultrasound after approximately 18 weeks gestational (menstrual) age. It may be possible to document normal structures before this time, although some structures can be difficult to visualize due to fetal size, position, movement, abdominal scars, and increased maternal abdominal wall thickness. A second or third trimester scan may pose technical limitations for an anatomic evaluation due to imaging artifacts from acoustic shadowing. When this occurs, the report of the sonographic examination should document the nature of this technical limitation. A follow-up examination may be helpful.
The following areas of assessment represent the minimal elements of a standard examination of fetal anatomy. A more detailed fetal anatomic examination may be necessary if an abnormality or suspected abnormality is found on the standard examination.

i. Head, face, and neck
- Cerebellum
- Choroid plexus
- Cisterna magna
- Lateral cerebral ventricles
- Midline falx
- Cavum septum pellucidum
- Upper lip

Comment:
A measurement of the nuchal fold may be helpful during a specific age interval to suggest an increased risk of aneuploidy.

ii. Chest
The basic cardiac examination includes a four-chamber view of the fetal heart.

If technically feasible, views of the outflow tracts should be attempted as part of the cardiac screening examination.

iii. Abdomen
- Stomach (presence, size, and situs)
- Kidneys
- Bladder
- Umbilical cord insertion site into the fetal abdomen
- Umbilical cord vessel number

iv. Spine
- Cervical, thoracic, lumbar, and sacral spine

v. Extremities
- Legs and arms – presence or absence

vi. Gender
Medically indicated in low-risk pregnancies only for evaluation of multiple gestations.

VI. DOCUMENTATION

Adequate documentation of the study is essential for high-quality patient care. There should be a permanent record of the ultrasound examination and its interpretation. Comparison with prior relevant imaging studies may prove helpful. Images of all appropriate areas, both normal and abnormal, should be recorded. Variations from normal size should generally be accompanied by measurements. Images should be labeled with the patient identification, facility identification, examination date, and image orientation. An official interpretation (final report) of the ultrasound examination should be included in the patient’s medical record. Retention of the ultrasound examination images should be consistent both with clinical need and with relevant legal and local healthcare facility requirements.

Reporting should be in accordance with the ACR Practice Guideline for Communication of Diagnostic Imaging Findings.

VII. EQUIPMENT SPECIFICATIONS

These studies should be conducted with real-time scanners, using a transabdominal and/or transvaginal approach. A transducer of appropriate frequency should be used.

Comment:
Real time sonography is necessary to confirm the presence of fetal life through observation of cardiac activity and active movement.

The choice of transducer frequency is a trade-off between beam penetration and resolution. With modern equipment, 3 to 5 MHz abdominal transducers allow sufficient penetration in most patients while providing adequate resolution. A lower-frequency transducer (2 to 2.25 MHz) may be needed to provide adequate penetration for abdominal imaging in an obese patient. During early pregnancy, a 5 MHz abdominal transducer or a 5 to 10 MHz or greater vaginal transducer may provide superior resolution while still allowing adequate penetration.

VIII. FETAL SAFETY

Diagnostic ultrasound studies of the fetus are generally considered to be safe during pregnancy. This diagnostic procedure should be performed only when there is a valid medical indication, and the lowest possible ultrasonic exposure setting should be used to gain the necessary diagnostic information under the “as low as reasonably achievable” (ALARA) principle.

The promotion, selling, or leasing of ultrasound equipment for making “keepsake fetal videos” is considered by the U.S. Food and Drug Administration to be an unapproved use of a medical device [9]. Use of a diagnostic ultrasound system for these purposes, without a physician’s order, may be in violation of state laws or regulations.
IX. QUALITY CONTROL AND IMPROVEMENT, SAFETY, INFECTION CONTROL, AND PATIENT EDUCATION

All probes should be cleaned after use. Vaginal probes should be covered by a protective sheath prior to insertion. Following the examination, the sheath should be disposed of and the probe cleaned in an antimicrobial solution. The type of solution and amount of time for cleaning depend on manufacturer and infectious disease recommendations.

Policies and procedures related to quality, patient education, infection control, and safety should be developed and implemented in accordance with the ACR Policy on Quality Control and Improvement, Safety, Infection Control, and Patient Education appearing under the heading Position Statement on QC & Improvement, Safety, Infection Control, and Patient Education on the ACR web page (http://www.acr.org/guidelines).

Equipment performance monitoring should be in accordance with the ACR Technical Standard for Diagnostic Medical Physics Performance Monitoring of Real Time Ultrasound Equipment.

ACKNOWLEDGEMENTS

This guideline was revised according to the process described under the heading The Process for Developing ACR Practice Guidelines and Technical Standards on the ACR web page (http://www.acr.org/guidelines) by the ACR Guidelines and Standards Committee of the Commission on Ultrasound in collaboration with the AIUM and the ACOG.

Principal Reviewer: Beryl R. Benacerraf, MD

Collaborative Subcommittee

ACR
Beryl R. Benacerraf, MD, Chair
Ruth B. Goldstein, MD

AIUM
Harris Finburg, MD
Wesley Lee, MD
Larry Platt, MD

ACOG
Fredric Frigoletto, Jr., MD
William N.P. Herbert, MD
Carolyn M. Zelop, MD

ACR Guidelines and Standards Committee
Gretchen A. Gooding, MD, Chair
Raymond E. Bertino, MD
Mary C. Frates, MD

Comments Reconciliation Committee
Marcela Bohm-Velez, MD, Co-Chair
Bill H. Warren, MD, Co-Chair
Beryl R. Benacerraf, MD
Carol B. Benson, MD
Douglas L. Brown, MD
Harris J. Finberg, MD
Mary C. Frates, MD
Ruth B. Goldstein, MD
Gretchen A. Gooding, MD
Gail C. Hansen, MD
Paul A. Larson, MD
Lawrence A. Liebscher, MD
Carol M. Rumack, MD
Julie K. Timins, MD
William G. Way, Jr., MD

Suggested Reading (Additional articles that are not cited in the document but that the committee recommends for further reading on this topic)


*Guidelines and standards are published annually with an effective date of October 1 in the year in which amended, revised or approved by the ACR Council. For guidelines and standards published before 1999, the effective date was January 1 following the year in which the guideline or standard was amended, revised, or approved by the ACR Council.

**Development Chronology for this Guideline**

1990 (Resolution 5)
Revised 1995 (Resolution 35)
Revised 1999 (Resolution 37)
Revised 2003 (Resolution 19)
Amended 2006 (Resolution 35)
Revised 2007 (Resolution 25)