INTRODUCTION

The American Institute of Ultrasound in Medicine (AIUM) strives to continuously improve the quality of diagnostic ultrasound services by offering a peer review process for providers to demonstrate that they meet nationally recognized standards and guidelines. Diagnostic medical ultrasound makes important contributions to patient care and may be used in a variety of settings. Ultrasound practice accreditation is designed to set a standard of quality for the performance of standard and detailed diagnostic assessments as well as the performance of interventional procedures.

Applying for accreditation displays your commitment to the highest quality patient care and exhibits your dedication to clinical excellence. Regardless of the specialty area in which your practice decides to apply, the same policies and procedures are uniform for all ultrasound practices.

The decision to accredit a practice is based on information submitted in the application. To be eligible for ultrasound practice accreditation, a practice must be an organization, group, or business that performs, interprets and reports ultrasound examinations. This includes a mobile only business that owns or leases machines. The practice must demonstrate compliance with AIUM accreditation requirements including but not limited to those described herein. If applying with more than one site, the same policies, protocols, and diagnostic criteria must be followed at all sites. Performance, interpretation, and archiving of medically indicated ultrasound examinations constitutes practice that is auditable and eligible for accreditation. Limited use of ultrasound for other purposes is not.

Interim Changes

In case of any changes in personnel, practice name or practice location, the AIUM’s Ultrasound Practice Accreditation Department (accreditation@aium.org) must be notified within 5 business days. An update form (linked here) must be completed and signed by the Director of Ultrasound and sent to accreditation@aium.org.

Reaccreditation

Reaccreditation is required every 3 years. For reaccreditation, licensed medical providers listed on the previous accreditation application will need to demonstrate compliance with all AIUM accreditation requirements, and each interpreting provider must provide evidence of meeting maintenance of competence (as evidenced by PP specific yearly ultrasound experiential volume) and Continuing Medical Education (i.e. AMA PRA Category 1 Credit™) requirements found at the end of each relevant training guideline. All provisional providers and other provisional accreditation stipulations must be resolved prior to re-accreditation.

ACCREDITATION PROGRAMS

- Abdominal/General
  - (with or without Adjunct Contrast Enhanced US – Coming soon)
- Breast (Diagnostic only, or Diagnostic with Interventional)
- Female Pelvic Floor
- Fetal Echocardiography
- Gynecologic (with or without Adjunct Competence in 3D)
- Musculoskeletal (Diagnostic)
- Musculoskeletal (Ultrasound-Guided Interventional Procedures)
- Standard Obstetric (or Trimester-Specific Standard OB)
ULTRASOUND PRACTICE PERSONNEL, ROLES, & REPORTING POLICIES

Director of Ultrasound
The practice must designate a Director of Ultrasound who is responsible for all correspondence with the accreditation department and must sign all required accreditation documentation. The Director of Ultrasound must be a licensed medical provider who meets the training guidelines in the specific area of sonography. The Director of Ultrasound is responsible for overseeing the quality and appropriateness of ultrasound operations of the practice. This includes ensuring that appropriate clinical services are provided and support services are sufficient, and for attesting that the practice continues to meet The Guideline for Ultrasound Practice Accreditation. The Director of Ultrasound may supervise the entire operation of the facility or may delegate specific operations to associates and sonographers.

Providers who Supervise and Interpret Ultrasound Examinations
All ultrasound studies must be supervised and interpreted by a licensed medical provider who meets the relevant training guideline(s) in a specific area of sonography. If interpreting in multiple specialties, providers must meet each relevant training guideline. Physicians, including advanced clinical providers, who perform and/or interpret ultrasound exams (even occasional coverage) are required to be included in the accreditation application. They must create a profile in the application describing where and how they received ultrasound training and list each CME activity (ama pra category 1) dedicated to ultrasound within the past 36 months. Since providers not only perform but also interpret and report on examinations, signature cards will be required for all such providers listed on the Practice Accreditation Application.

General Documentation of Training and Competence Requirements
➢ Different areas may have special requirements for documentation of training and competence. Please refer to the appropriate Practice Parameter(link) and contact the AIUM office if you are unsure
➢ From each provider during initial application, and when new interpreting providers join the practice, the following must be submitted within 5 business days:

1. Copy of medical or other professional license, as appropriate.

2. Copy of board certification(s) or other certification types as required to meet applicable ultrasound training guidelines (required for all providers applying for the first time or if applying to interpret in a new specialty area).

3. Evidence of involvement in interpreting and reporting the minimum annual number of examinations recommended to maintain continuing competence in the applicable specialties.
Sonographers Who Perform Ultrasound Examinations

Qualified sonographers who perform ultrasound examinations will be responsible for those tasks specified by the Director of Ultrasound. Although a sonographer may play a critical role in extracting the information essential to deriving a diagnosis, the rendering of a final diagnosis from ultrasound studies represents the practice of medicine and, therefore, is the responsibility of the supervising licensed medical provider.

Sonographers must have appropriate training for the ultrasound examinations they perform. All sonographers must be certified in the specialty or specialties in which they perform ultrasound evaluations or must become certified before the next reaccreditation application. If the sonographer is unsuccessful at obtaining the relevant credential, he/she must not perform this type of ultrasound for the practice except for the purposes of training and in these cases only with supervision of a physician or sonographer who meets the training guideline. If a sonographer is required to become certified in multiple specialties, the sonographer must obtain a minimum of 1 additional specialty certification per accreditation cycle.

The following certifications are acceptable:

- Certification (active status) by the American Registry for Diagnostic Medical Sonography (ARDMS) in abdomen, breast, fetal echocardiography, musculoskeletal, obstetrics and gynecology, and others as applicable;
- Certification in general sonography for abdomen and obstetrics and gynecology by the American Registry of Radiologic Technologists (ARRT) or the ARRT certification in breast sonography.
- International applicants may provide equivalent credentials for review by UPAC.
Other Providers Who Perform Ultrasound Examinations
Non-physicians who perform ultrasound examinations must have appropriate training in their scope of practice. The following are specialty areas that have training guidelines approved to include other providers on the accreditation application.

- **Musculoskeletal Ultrasound**
- **Limited OB Ultrasound for ACP** – This training guideline is NOT for those performing and/or interpreting a more detailed OB examination.
- **Point of care Ultrasound**

**Documentation Requirements**

- **For each sonographer:**
  Copy of current ARDMS or ARRT registry card (can be electronic proof if applicable)

- **For each non-physician provider:**
  Documentation received documenting appropriate training as described in relevant training guidelines. (Linked above)

**Final Reports**

Final reports must meet the requirements specified in the [AIUM Practice Parameter for Documentation of an Ultrasound Examination](https://www.aium.org). If your practice interprets in interventional procedures, carefully review section V, "Reporting of Ultrasound-Guided Procedures”.

**Preliminary Reports**

A preliminary report is a written or verbal report released before being signed by the physician responsible for giving the final interpretation. Refer to "Non-routine Reporting” in the [AIUM Practice Parameter for Documentation of an Ultrasound Examination](https://www.aium.org):  

If a licensed medical provider responsible for the interpretation is not immediately available at the time an ultrasound examination is being performed:

- The sonographer performing the examination must be appropriately credentialed in the specialty area(s) of the examination.

- A mechanism must be in place to address unexpected or emergency findings.

- If the patient is released prior to the interpreting provider review of the images, a patient callback mechanism must be in place.

- A final report of the examination must be available in a timely manner not to exceed that recommended in the [AIUM Practice Parameter for Documentation of an Ultrasound Examination](https://www.aium.org).

### POLICIES AND PROCEDURES SAFEGUARDING PATIENTS AND PERSONNEL

**ALARA Principle**

AIUM advocates for the responsible use of diagnostic ultrasound for all ultrasound imaging. Application of ultrasound to humans should be limited to that which is medically necessary. Personnel must be
familiar with and show evidence of practicing the ALARA (as low as reasonably achievable) principle and imaging displays as well as permanent recorded images must include Mechanical and Thermal index values. Practices and practitioners and sonographers must follow all AIUM official statements on safety including but not limited to the following:

- **Medical Ultrasound Safety**
- **Recommended Maximum Scanning Times for Displayed Thermal Index Values**
- **Statement on Measurement of Fetal Heart Rate**
- **Statement on the Safe Use of Doppler Ultrasound During 11–14 week scans (or earlier in pregnancy)**
- **Prudent Use in Pregnancy**
- **Keepsake Fetal Imaging**
- **Conclusions Regarding Epidemiology for Obstetric Ultrasound**
- **Statement on Mammalian Biological Effects in Tissues with Gas Body Contrast Agents**

**Prevention of Infectious Diseases**

- The practice must have procedures and policies on the protection of patients and practice personnel from the transmission of infectious disease.

- The practice must follow the *Guidelines for Cleaning and Preparing External- and Internal-Use Ultrasound Probes Between Patients, Safe Handling, and Use of Ultrasound Coupling Gel*.

**Documentation Requirements**

- Most recent proof of purchase for high level disinfection agent.
- Cleaning logs

**Patient Confidentiality**

All practice personnel must adhere to Health Insurance Portability and Accountability Act regulations and professional ethics and behavior to ensure patient confidentiality.

**Patient Identification**

Patients must be identified by 2 separate identifiers before a study is performed: for example: name, date of birth, address, or medical record number.

**Precautions for Invasive Procedures**

*Time-outs*

- Before beginning an ultrasound-guided invasive procedure, the practice must take steps to verify patient identification, the type of procedure planned, and the appropriate procedure site(s).

*Specimens*
• The practice must take steps to ensure that specimens are correctly labeled.
• The practice must clearly define the personnel responsible and the steps required to hand off each specimen.

**Documentation Requirements**

- Written policy on precautions for invasive procedures.

**Incident Reporting**

A policy/procedure must exist for responding to and reporting any accidents or complications that occur in the facility.

**Documentation Requirements**

- Written policy on incident reporting

**Personnel Performance Quality Assurance**

The practice must show ongoing evaluation and improvement of the clinical practice's ultrasound personnel performance, including all interpreting providers and sonographers through regular, retrospective peer review. This includes systematic review of performance, documentation, and reporting as well as diagnostic comparison across modalities:

• To assess diagnostic accuracy, the practice must obtain correlation of ultrasound diagnoses of normal and abnormal studies with clinical, radiographic, laboratory, surgical, and pathologic findings.

• The review must include normal and abnormal case studies to evaluate the following:
  - Content, completeness, and technical quality of the images; and
  - Accuracy and timeliness of final reports.

• When deficiencies are identified, the causes should be investigated, and corrective action should be planned and implemented.

• Information obtained should be disseminated to both physician and sonographer personnel of the ultrasound practice in a timely fashion.

• A record of QA activities must be maintained and kept current. Records may be requested at the time of initial or reaccreditation.

**Documentation Requirements**

- A summary of personnel performance quality assurance activities (complete records are not required).
  - Samples:
    - Performance Evaluation Form – OB
    - Sample - Quarterly QA – Fetal Echo
    - More samples will be added for other specialty areas as soon as available
Record-keeping Guidelines and Requirements

- There must be provisions for the retrieval of examination records (ultrasound images and cine clips) of all studies performed.

- Appropriate documentation of every study must include permanent ultrasound images stored on suitable recording media, as well as a report that indicates the findings obtained by the examination and adheres to the **AIUM Practice Parameter for Documentation of an Ultrasound Examination**.

- Ultrasound images and a final report from the interpreting physician or qualified healthcare provider must be maintained in a readily accessible fashion for comparison and consultation.

- Recording media must have a shelf life compatible with the minimum number of years, required by law, for the maintenance of patient records. In most states, this will be for at least 7 years after the patient’s last examination was performed; however, these requirements vary from state to state and may be considerably longer for pediatric patients.

Policies Protecting Practice Personnel

The practice must be in compliance with all US Occupational Safety and Health Administration regulations. Medical staff should observe universal precautions.

The practice should be mindful of ergonomics as related to the use of ultrasound, and adhere to the **AIUM Official Statement on Preventing Work-related Musculoskeletal Disorders** to promote a safe work environment.

Ultrasound Equipment and Quality Assurance

The ultrasound equipment must meet all state and federal guidelines.

- Studies must be conducted with real-time equipment, and transducers must be available with a frequency range that will optimize beam penetration and resolution.

- The equipment used for diagnostic testing must be maintained in good operating condition and undergo routine quality assurance at least once a year or more frequently if problems arise. Regular assessment of transducer operation and quality control of repaired transducers are critical for optimal patient care. Practices must follow recommendations regarding **transducer testing and repair**. Routine quality assurance on ultrasound equipment must meet or exceed the recommendations specified in **Routine Quality Assurance for Diagnostic Ultrasound Equipment**.
  - User friendly guide to ultrasound QA: [https://www.aium.org/accreditation/qualityAssurance.pdf](https://www.aium.org/accreditation/qualityAssurance.pdf)

- Records of routine quality assurance must be kept for 3 years or longer depending on state and federal guidelines.

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<tr>
<th>Documentation Requirements</th>
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<tr>
<td>➢ Most recent ultrasound equipment logs</td>
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<tr>
<td>➢ Documentation of staff and physician participation in Quality Assurance Activities (such as sign-in sheets documentation of attendance or certificates of completion—general statements are NOT acceptable)</td>
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CASE STUDY SUBMISSIONS

For the purpose of accreditation, a “study” consists of images performed on a single date only with a corresponding final report. The accreditation requirements are based on the relevant AIUM Practice Parameter for the Performance of an Ultrasound Examination for each relevant specialty. Accreditation requirements may be different in order to demonstrate certain competencies.

In certain situations, for detailed obstetric ultrasounds (76811) and fetal echocardiography, fetal position can make documenting abnormalities within one visit difficult. If this occurs, a follow-up study within a week is acceptable provided all documentation requirements are met. Both the original and the follow up images and reports must be submitted for full review, along with a written explanation regarding the patient’s circumstances. These submissions must be cases that are indicated and not simply added anatomic components to a limited ob examination.

Case studies must be performed and interpreted by a representative sample of the practice’s interpreting providers and sonographers. A single interpreting provider or sonographer should not submit more than one case unless all other listed providers have each submitted a case. Original reports must be submitted.

Documentation Requirements
> See links below for requirements specific to each area of accreditation.

Case studies submitted for review must comply with the following:

1. AIUM Practice Parameter for the Documentation of an Ultrasound Exam.

2. Adhere to the Case Submission Requirements for each specialty:
   - Abdominal/General Case Requirements
     - (with or without Adjunct Contrast Enhanced US – Coming soon)
   - Breast Case Requirements
   - Fetal Echocardiography Case Requirements
   - Female Pelvic Floor Case Requirements
   - Gynecologic (with or without 3D) Case Requirements
   - Musculoskeletal (Diagnostic) Case Requirements
   - Musculoskeletal (US-Guided Interventional Procedure) Case Requirements
   - Point of Care Ultrasound
   - Standard Obstetric (or Trimester-Specific OB) Case Requirements
   - Limited Obstetric Ultrasound for ACP Case Requirements
   - Standard OB with Adjunct Detailed Fetal Anatomic US (76811) Case Requirements
   - Thyroid, Parathyroid, and Neck Case Requirements
• Urologic Case Requirements

• Ultrasound-Guided Regional Anesthesia Case Requirements

• Cases must adhere to the General File Requirements and How to Prepare Your Documents for Upload.

• Case studies must have been performed within 12 months of the date they are uploaded.

• In order to assess timeliness of the final report, interpreting providers’ signatures must be dated. If the report does not accurately display the date of signature, this should be noted and an explanation provided. A printed name must be next to the signature if the signature is illegible.

• While the Ultrasound Practice Accreditation Council (UPAC) understands that providers reading the scans are aware of what each image demonstrates, the protocol for practice evaluation is uniform and must cover a broad spectrum of practices. In order to ensure that the appropriate structure is being identified, labeling is required for cases that are submitted. Post-process labeling is acceptable.

• Video clips should not be submitted unless the case requirements specifically request them. Most ultrasound specialties can adequately display the required anatomy in still images. If video clips are allowed, clips must be labeled in the clip or labeled as the file name. If cine clips are submitted, representative thumbnail still frames from the clip should be submitted with the appropriate labels for the structures intended to be demonstrated on the cine clip.

• Only complete cases documenting all required components should be submitted for evaluation.

• Although in practice there are cases when the full anatomy is not seen, these are not acceptable for submission. Reporting a structure as “suboptimal” may clarify appropriately why anatomy was not clearly seen; however, this case would not be considered suitable for submission.