

# MQSA Facility Certification Extension Requirements for Digital Breast Tomosynthesis (DBT) System

**NOTE 1** Under MQSA, 8 hours of new modality training obtained on any DBT system, or general DBT training, is considered sufficient to meet the MQSA new modality requirement for DBT.

**NOTE 2:** In order to use the tomosynthesis portion of the unit, the facility must apply to FDA to have its certificate extended to include that portion of the unit. **The certification extension only applies to the DBT portion of the unit.** The facility must have the 2D portion of the unit accredited by one of the accreditation bodies already approved to accredit the 2D portion.

## Requirements:

**FFDM-DBT system Manufacturer Name:**

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## 2. Facility Status Information

a. Facility Name and FDA Facility ID Number

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b. FDA Certificate Expiration Date

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c. Current Accreditation Body for the 2D unit

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d. Accreditation Expiration Date

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e. Facility Contact Person for DBT unit

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f. Contact Person's Title

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g. Contact Person's Telephone, Fax, E-mail

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h. Facility Address

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i. Facility Owner

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## 3. DBT Unit Identification

a. Manufacturer

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b. Model

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c. Year of Manufacture

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d. Serial Number

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e. Accreditation Body Unit ID #

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## 4. DBT Digital Image Receptor Identification (if interchangeable)

a. Receptor Manufacturer

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b. Receptor Model

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c. Year of Manufacture

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d. Serial Number (if Applicable)

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**5. Final Interpretation Review Monitor Identification (if soft copy display is available)**

- a. Monitor Manufacturer
- b. Monitor Model
- c. Year of Manufacture
- d. Serial Number


**6. Phantom Identification**

- a. Phantom Manufacturer
- b. Phantom Model


**7. Submit either a hardcopy or softcopy 3D phantom image.** It's preferred that softcopy CD or DVD be in DICOM format and verified that the image opens properly before forwarding the 3D phantom image to the FDA. **(Failure to include a 3D phantom image will delay review of the application).**

**8. Personnel Qualifications**

- a. Interpreting Physicians who are qualified to interpret DBT mammograms (see Qualified Personnel)
- b. Radiological Technologists who are qualified to perform DBT mammography examinations and the manufacturer recommended quality assurance tests (see Qualified Personnel)
- c. Medical Physicists who are qualified to perform equipment evaluations and/or surveys of DBT mammography units (Qualified Personnel)

**9. Complete Detailed report of Mammography Equipment Evaluation (MEE)** (must have been conducted in accordance with 900.12(e)(10) within the 6 months prior to the request for use approval) **must be included when submitting application.**

- a. Statement that equipment performance, as required under the following sections of the MQSA final regulation 21 CFR 900.12(b), is met:
  - (1) Prohibited Equipment
  - (2) Specifically Designed for Mammography
  - (3) Motion of Tube-Image Receptor Assembly
  - (4)(iii) Removable Grid (if applicable to the DBT system used)
  - (5) Beam Limitation and Light Fields
  - (6) Magnification
  - (7) Focal Spot Selection
  - (8) Compression
  - (9) Technique Factor Selection and Display
  - (10) Automatic Exposure Control
- b. The results of quality control tests as required under the following sections of the MQSA final regulations 21CFR 900.12(e):

- (4)(iii) Compression Device Performance
  - (5)(i) Automatic Exposure Control Performance (if applicable to the DBT system used)
  - (5)(ii) Kilovoltage Peak Accuracy and Reproducibility
  - (5)(iii) Focal Spot Condition (Resolution)
  - (5)(iv) Beam Quality and Half-Value Layer
  - (5)(v) Breast Entrance Air Kerma and AEC Reproducibility (if applicable to the DBT system used)
  - (5)(vi) Dosimetry
  - (5)(vii) X-Ray Field/Light field/Image receptor/Compression paddle alignment
  - (5)(ix) System Artifacts
  - (5)(x) Radiation Output
  - (5)(xi) Decompression (or alternative standards allowed for these requirements)
  - (6) Quality Control Tests – Other Modalities (Facilities must perform all DBT manufacturer recommended quality control tests including the medical physicist’s tests for Soft Copy Display system)
- c. The results of the phantom image quality tests, including a sample image
  - d. If any of the requirements in 8 a, b, or c are not met, submit documentation of successful corrective action
  - e. If any of the requirements in 8 a or b are not performed, explain why the requirement is not applicable
  - f. Date of the MEE
  - g. Name and Address of the physicist(s) who performed the MEE

**10. DBT Manufacturer’s Quality Control Program**

a. Name of the Quality Control Manual	
b. Year Published	
c. Revision Number (if not original)	
d. Printing Number (if not original)	

**11. Signature of facility contact person for the DBT unit**

Signature

**Qualified Personnel**

**Interpreting Physicians**

PERSONNEL QUALIFICATIONS: INTERPRETING PHYSICIANS WHO ARE QUALIFIED TO INTERPRET DBT MAMMOGRAMS

List the current interpreting physicians who:

- (1) Meet all the requirements of 21 CFR 900.12(a)(1) "Mammography Quality Standards; Final Rule" that became effective on April 28, 1999; and
- (2) have 8 hours of initial new-modality training in DBT; training on any DBT system, or general DBT training, is sufficient to meet the MQSA new modality training requirement.\*

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\* Supporting documentation for these requirements will be checked during annual MQSA inspections.

**Radiologic Technologists**

**PERSONNEL QUALIFICATIONS: RADIOLOGIC TECHNOLOGISTS WHO ARE QUALIFIED TO PERFORM DBT MAMMOGRAMS**

List the current radiologic technologists who:

- (1) Meet all the requirements of 21 CFR 900.12(a)(3) "Mammography Quality Standards; Final Rule" that became effective on April 28, 1999; and
- (2) have 8 hours of initial new-modality training in DBT; training on any DBT system, or general DBT training, is sufficient to meet the MQSA new modality training requirement.\*

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\*Supporting documentation for these requirements will be checked during annual MQSA inspections.

**Medical Physicists**

**PERSONNEL QUALIFICATIONS: MEDICAL PHYSICISTS WHO ARE QUALIFIED TO PERFORM DBT SURVEYS**

List the current medical physicists who:

- (1) Meet all the requirements of 21 CFR 900.12(a)(3) "Mammography Quality Standards; Final Rule" that became effective on April 28, 1999; and
- (2) have 8 hours of initial new-modality training in DBT; training on any DBT system, or general DBT training, is sufficient to meet the MQSA new modality training requirement.\*

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\*Supporting documentation for these requirements will be checked during annual MQSA inspections.

**Lead Interpreting Physician Attestation to Staff Personnel Qualifications**

To the best of my knowledge and my belief, the information provided in this document is true and correct. I understand that FDA may request additional information to substantiate the statements made in the document. I understand that knowingly providing false information in a matter within the jurisdiction of an agency of the United States could result in criminal liability, punishable by up to \$10,000 fine and imprisonment of up to five years, or civil liability under MQSA, or both.

Signature  
(Lead  
Interpreting  
Physician)

Print Name:

Date: