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.

Requiremen	ts:													
FFDM-DBT system Name:	m Manufactı	ırer												
2. Facility Status I	nformation													
a. Facility Name and F	DA Facility ID	Number												
b. FDA Certificate Exp	iration Date													
c. Current Accreditation	n Body for the 2	D unit												
d. Accreditation Expira	tion Date													
e. Facility Contact Person for DBT unit														
f. Contact Person's Title														
g. Contact Person's Telephone, Fax, E-mail														
h. Facility Address														
i. Facility Owner														
3. DBT Unit Ident	ification													
a. Manufactu	urer													
b. Model														
c. Year of Manufacture														
d. Serial Nu	mber													
e. Accreditat Unit ID#	tion Body													
4. DBT Digital Ima	age Receptor	Ident	ifi	catio	on (it	inte	erch	ange	eable)				
a. Receptor	Manufacturer													
b. Receptor Model												1		
c. Year of Manufacture												1		
d. Serial Number (if														

Applicable)

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	
CD or DVD be in DICOM forma	softcopy 3D phantom image. It's preferred that softcopy t and verified that the image opens properly before to the FDA. (Failure to include a 3D phantom image tion).
8. Personnel Qualifications	
 a. Interpreting Physicians who Qualified Personnel) 	are qualified to interpret DBT mammograms (see
	who are qualified to perform DBT mammography facturer recommended quality assurance tests (see
c. Medical Physicists who are of DBT mammography unit	qualified to perform equipment evaluations and/or surveys s (Qualified Personnel)
have been conducted in accordan	Mammography Equipment Evaluation (MEE) (must ce with 900.12(e)(10) within the 6 months prior to the included when submitting application.
a. Statement that equipment po MQSA final regulation 21 C	erformance, as required under the following sections of the CFR 900.12(b), is met:
(1) Prohibited Equipment	
(2) Specifically Designed for	or Mammography
(3) Motion of Tube-Image	Receptor Assembly
(4)(iii) Removable Grid (if and Light Fields	applicable to the DBT system used) (5) Beam Limitation
(6) Magnification	
(7) Focal Spot Selection	

b. The results of quality control tests as required under the following sections of the MQSA final regulations 21CFR 900.12(e):

(8) Compression

(9) Technique Factor Selection and Display

(10) Automatic Exposure Control

- (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.*

Please enter text here:	
* Supporting docume inspections.	entation for these requirements will be checked during annual MQSA
	Radiologic Technologists
	IFICATIONS: RADIOLOGIC TECHNOLOGISTS WHO ARE RFORM DBT MAMMOGRAMS
List the current radiol	ogic technologists who:
	rements of 21 CFR 900.12(a)(3) "Mammography Quality Standards; Final Pective on April 28, 1999; and
	itial new-modality training in DBT; training on any DBT system, or general cient to meet the MQSA new modality training requirement.*
Please enter text here:	
*Supporting document inspections.	ntation for these requirements will be checked during annual MQSA
	Medical Physicists
PERSONNEL QUAL PERFORM DBT SUF	IFICATIONS: MEDICAL PHYSICISTS WHO ARE QUALIFIED TO RVEYS
List the current medic	al physicists who:
	rements of 21 CFR 900.12(a)(3) "Mammography Quality Standards; Final Pective on April 28, 1999; and
	itial new-modality training in DBT; training on any DBT system, or general cient to meet the MQSA new modality training requirement.*
	stent to meet the MQSA new modarity training requirement.
Please enter text here:	

*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:		