

User Guide MAN-09291-002 Revision 001



3Dimensions[™]

Digital Mammography System

Digital Tomosynthesis System

User Guide For Software Version 2.3

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Chapter 1 Introduction

Read all this information carefully before operating the system. Follow all warnings and precautions as stated in this manual. Keep this manual available during procedures. Physicians should tell patients about all potential risks and adverse events described in this manual with respect to the operation of the system.



Note

Hologic configures some systems to meet specific requirements. Your system configuration may not have all the options and accessories included in this manual.



Note

The features shown in this manual may not be available in all regions. Contact your Hologic representative for information.

1.1 Intended Uses

 $R_{\!X}^{}$ Only

Caution: United States federal law restricts this device to sale by or on the order of a physician.

1.1.1 3Dimensions System Intended Use

The Hologic[®] 3DimensionsTM system generates digital mammographic images that can be used for screening and diagnosis of breast cancer. The 3Dimensions (2D or 3D) system is intended for use in the same clinical applications as a 2D mammography system for screening mammograms. Specifically, the 3Dimensions system can be used to generate 2D digital mammograms and 3D mammograms. Each screening examination may consist of:

- a 2D FFDM image set; or
- a 2D and 3D image set, where the 2D image can be either an FFDM or a 2D image generated from the 3D image set and the 3D image set can be viewed as either 1-mm 3D slices or 6-mm 3D SmartSlices.

The 3Dimensions system may also be used for additional diagnostic workup of the breast.



Note

In Canada and Singapore, Tomosynthesis is not approved for screening, and must be used in conjunction with a 2D image (either a FFDM image or 2D image generated from the 3D image set).

Contrast Enhanced Digital Mammography

Contrast Enhanced Digital Mammography (CEDM) is an extension of the existing indication for diagnostic mammography with the 3Dimensions system. The CEDM application shall enable contrast enhanced breast imaging using a dual energy technique. This imaging technique can be used as an adjunct following mammography and/or ultrasound exams to localize a known or suspected lesion.

1.1.2 Genius AI Detection Intended Use

Genius AI® Detection is a computer-aided detection and diagnosis (CADe/CADx) software device intended to be used with compatible digital breast tomosynthesis (DBT) systems to identify and mark regions of interest including soft tissue densities (masses, architectural distortions, and asymmetries) and calcifications while reading DBT exams from compatible DBT systems and provide confidence scores that offer assessment for Certainty of Findings and a Case Score. The device intends to aid in the interpretation of digital breast tomosynthesis exams in a concurrent fashion, where the interpreting physician confirms or dismisses the findings during the reading of the exam.

1.1.3 Contraindications

There are no known contraindications.

1.2 Intended User

Qualified technically trained personnel position the patient and operate the device (such as a mammography technologist), to calibrate the device (such as a medical physicist) and to interpret the image produced by the device (such as a radiologist).

1.3 Patient Target Group

Asymptomatic patients who are at average risk or at higher than average risk for breast cancer require breast cancer screening imaging. Those patients who may have symptoms related to breast cancer or abnormal findings on a screening exam also need diagnostic imaging using the same or similar modalities. Qualified medically-trained personnel will decide based on shared decision making with the patient and, in some cases, the provider whether a radiographic examination is indicated based on patient age, patient health, and physical condition.

1.4 Clinical Benefits

Population based mammography screening is the best method for detecting breast cancer before symptoms occur and is the only modality that has been proven to reduce mortality.

1.5 Potential Adverse Effects of Mammography Systems on Health

Following is a list of the potential adverse effects (such as complications) associated with the use of the device (these risks are the same as for other screen-film or digital mammography systems):

- Excessive breast compression
- Excessive x-ray exposure
- Electric shock
- Infection
- Skin irritation, abrasions, or puncture wounds

1.6 System Capabilities

The system provides the user interfaces for the performance of screening and diagnostic mammograms:

- Conventional mammography with a digital image receptor equivalent in size to large mammography film.
- Tomosynthesis scan with a digital image receptor equivalent in size to large mammography film (Tomosynthesis option).
- Conventional digital mammogram and tomosynthesis scan during one compression (Tomosynthesis option).

1.7 About Genius 3D Mammography Exam

The Genius[®] 3D MammographyTM exam (also known as Genius[®] exam) is acquired on a Hologic[®] 3D MammographyTM system and has a 2D and a 3DTM image set. The 2D image can be either an acquired 2D image or a 2D image generated from the 3DTM image set. The Genius[®] exam is only available on a Hologic[®] 3D MammographyTM system.

Genius[®] 3D MammographyTM is the brand name of a Hologic[®] 3D MammographyTM exam and may not be available in all markets.

1.8 More Information About Tomosynthesis

The Hologic Selenia Dimensions system received FDA approval for the Hologic Tomosynthesis option on 11 Feb 2011 (refer to PMA number P080003). This FDA approval applies to screening and diagnostic imaging. More information is available from the FDA website at

http://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfpma/pma.cfm?id=P080003.

The Generated 2D option (C-View) in conjunction with Tomosynthesis received FDA approval on 16 May 2013 (refer to PMA number P080003 S001). More information is available from the FDA website at

http://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfpma/pma.cfm?id=P080003S001.

A list of scientific publications about Breast Tomosynthesis is available from the Hologic website. The majority of the studies were done with the commercially released Hologic Selenia Dimensions Tomosynthesis system. See the publication at <u>http://www.hologic.com/sites/default/files/Tomo-Bibliography-Rev-13.pdf</u>.

The Hologic website has several white papers and summary cards about breast imaging publications. See the documents at <u>http://www.hologic.com/en/learning-center/white-papers/breastimaging/</u>.

Independent publications have examined Hologic Selenia Dimensions Tomosynthesis systems in population-based screening sequences in Europe. Results consistently show a significant increase in invasive cancer detection rate at the same time as a reduction in false positives. The following publications are recommended.

Integration of 3D digital mammography with tomosynthesis for population breastcancer screening (STORM): a prospective comparison study.

Ciatto S, Houssami N, Bernardi D, Caumo F, Pellegrini M, Brunelli S, Tuttobene P, Bricolo P, Fantò C, Valentini M, Montemezzi S, Macaskill P. Lancet Oncol. 2013 Jun;14(7):583-9. doi: 10.1016/S1470-2045(13)70134-7. Epub 2013 Apr 25. <u>http://www.ncbi.nlm.nih.gov/pubmed/23623721</u>

Prospective trial comparing full-field digital mammography (FFDM) versus combined FFDM and tomosynthesis in a population-based screening programme using independent double reading with arbitration.

Skaane P, Bandos AI, Gullien R, Eben EB, Ekseth U, Haakenaasen U, Izadi M, Jebsen IN, Jahr G, Krager M, Hofvind S.

Eur Radiol. 2013 Aug;23(8):2061-71. doi: 10.1007/s00330-013-2820-3. Epub 2013 Apr 4.

http://www.ncbi.nlm.nih.gov/pubmed/23553585

Comparison of digital mammography alone and digital mammography plus tomosynthesis in a population-based screening program.

Skaane P, Bandos AI, Gullien R, Eben EB, Ekseth U, Haakenaasen U, Izadi M, Jebsen IN, Jahr G, Krager M, Niklason LT, Hofvind S, Gur D. Radiology. 2013 Apr;267(1):47-56. doi: 10.1148/radiol.12121373. Epub 2013 Jan 7.

http://www.ncbi.nlm.nih.gov/pubmed/23297332

Two-view digital breast tomosynthesis screening with synthetically reconstructed projection images: comparison with digital breast tomosynthesis with full-field digital mammographic images.

Skaane P, Bandos AI, Eben EB, Jebsen IN, Krager M, Haakenaasen U, Ekseth U, Izadi M, Hofvind S, Gullien R. Radiology. 2014 Jun;271(3):655-63. doi: 10.1148/radiol.13131391. Epub 2014 Jan 24

http://www.ncbi.nlm.nih.gov/pubmed/24484063

Breast Cancer Screening Using Tomosynthesis in Combination With Digital Mammography

Sarah M. Friedewald, MD1; Elizabeth A. Rafferty, MD2; Stephen L. Rose, MD3,4; Melissa A. Durand, MD5; Donna M. Plecha, MD6; Julianne S. Greenberg, MD7; Mary K. Hayes, MD8; Debra S. Copit, MD9; Kara L. Carlson, MD10; Thomas M. Cink, MD11; Lora D. Barke, DO12; Linda N. Greer, MD13; Dave P. Miller, MS14; Emily F. Conant, MD15

JAMA. 2014;311(24):2499-2507. doi:10.1001/jama.2014.6095

http://jama.jamanetwork.com/article.aspx?articleid=1883018



IMPORTANT:

Hologic strongly recommends that users make themselves familiar with local or regional regulations. These regulations could impose restrictions on the different types of clinical use. Because the regulations could evolve and change over time, periodic review is recommended.



About C-View and Intelligent 2D Imaging Technologies



Note

Intelligent 2DTM imaging technology may not be available in all regions. Contact your sales representative for information.

The C-View[™] and Intelligent 2D[™] software uses image data available from a breast tomosynthesis acquisition to generate one digital mammogram (2D) per breast tomosynthesis acquisition. The synthesized 2D image is created without the need for an additional digital mammography exposure. The synthesized 2D image is designed to appear similar to, and serve the same purpose as, a digital mammogram (2D) when used as part of a screening study employing tomosynthesis. The C-View or Intelligent 2D image is interpreted in combination with a breast tomosynthesis image set and is not intended to be used without the accompanying breast tomosynthesis images to make a clinical decision or diagnosis.

1.9.1 Warnings and Precautions for C-View and Intelligent 2D Imaging Technologies



Warning:

Do not make a clinical decision or diagnosis from the C-View or Intelligent 2D images without reviewing the accompanying tomosynthesis image set.

Use the C-View or Intelligent 2D synthesized 2D images in the same way you would use conventional digital mammography (2D) when performing a screening study employing tomosynthesis.

- While reviewing the C-View or Intelligent 2D images for items or areas of interest, compare to a prior digital mammogram (2D) if priors exist and then review the related tomosynthesis images carefully.
- Carefully examine the entire tomosynthesis image set before making a clinical decision.



Warning:

The appearance of a C-View or Intelligent 2D synthesized 2D image may differ from that of a conventional digital mammography (2D) image, just as 2D film and digital mammography (2D) images from different vendors may look different.

Users should ensure they are adequately trained and are familiar with the appearance of C-View and Intelligent 2D synthesized 2D images before using them in conjunction with tomosynthesis image sets.

1.9.2 Theory of Operation for C-View and Intelligent 2D Imaging Technologies

Overview

The C-View and Intelligent 2D software is an image processing application for postprocessing the pixel data from tomosynthesis data, captured on a Hologic tomosynthesis imaging system, into a digital mammography (2D) image. The C-View or Intelligent 2D synthesized 2D image can be used in place of a digital mammogram (2D) as part of a screening study employing tomosynthesis.

The C-View and Intelligent 2D software processes tomosynthesis data to generate 2D images which are designed to appear similar to and serve the same purpose as a digital mammogram (2D) when used as part of a screening study employing tomosynthesis.

Configuration

The C-View and Intelligent 2D software has no user-configurable settings that influence the appearance of the resulting synthesized 2D images. C-View and Intelligent 2D images are produced in either DICOM Breast Tomosynthesis Image format, as a single thick slice, or DICOM Digital Mammography X-Ray Image format. The site PACS administrator, in consultation with Hologic connectivity engineers, can select the output format most appropriate for the site's IT infrastructure and workstations. Each C-View or Intelligent 2D image DICOM header contains the information needed to differentiate it from any accompanying conventional 2D image(s) or tomosynthesis image sets in the same view. An annotation ("C-View" or "Intelligent 2D") is also burned to the synthesized 2D image pixel data.

Workflow

As with any imaging study, the technologist selects the patient and identifies the type of imaging procedure that will be done. For an exam with C-View or Intelligent 2D images, the process of imaging the subject and completing the study is all that is necessary. The C-View and Intelligent 2D software itself operates with no direct human intervention.

1.10 About 3DQuorum Software



Note

3DQuorum[®] imaging technology may not be available in all regions. Contact your sales representative for information.

3DQuorum[®] software uses image data available from a high-resolution breast tomosynthesis acquisition to generate SmartSlices, which are "slabbed" 3D images. The SmartSlices are created without the need for an additional digital tomosynthesis exposure. A SmartSlice can be perceived as a "thick" version of a tomosynthesis slice and is equivalent to six slices (6 mm) from the original tomosynthesis reconstruction. The SmartSlice volume consists of sequentially synthesized slices. Each SmartSlice shares half of its contents with the SmartSlice before and the other half with the SmartSlice after. Therefore, when reviewing SmartSlices, the images are advanced three 1-mm tomosynthesis slices at a time. The resulting number of images to be reviewed by a radiologist in a study is one-third the number of a tomosynthesis 1-mm reconstruction image set.

SmartSlices are designed to reduce the total tomosynthesis screening study interpretation time by decreasing the total number of images a radiologist is required to review. SmartSlices images are not intended to be used without the synthesized 2D images to make a clinical decision or diagnosis.

1.10.1 Warnings and Precautions for 3DQuorum Software



Warning:

Do not make a clinical decision or diagnosis from the SmartSlice images without reviewing the accompanying synthesized 2D image set.



Warning:

Refer to the imaging system *User Guide* **for a complete list of warnings and cautions.**

Use SmartSlices in the same way you would use the 3D 1-mm slices (or tomosynthesis reconstructed slices) when reviewing a tomosynthesis screening study. Users should ensure they are adequately trained and are familiar with the appearance of SmartSlices before using them in place of tomosynthesis reconstructed slices.

1.10.2 Theory of Operation for 3DQuorum Software

Overview

3DQuorum software is an image processing application for post-processing a limited number of slices of tomosynthesis data, captured on a Hologic tomosynthesis imaging system, into a combined 3D image. The SmartSlices image set reduces the number of images to one-third compared to tomosynthesis reconstructed slices data set while maintaining clinical information. As a result, read times for the radiologist can be reduced by about 13%. And for archival and transmission purposes, file sizes can be reduced by over 50%. SmartSlices can be used in place of the tomosynthesis reconstructed slices when reviewing a screening study employing tomosynthesis. SmartSlices achieve the same sensitivity as reading 1-mm 3DTM slices.

Configuration

3DQuorum software has no user-configurable settings that influence the appearance of the resulting SmartSlices. SmartSlices are stored and transmitted as a series of frames using the DICOM Breast Tomosynthesis Image object format. The DICOM header for SmartSlices contains the information needed to differentiate it from any accompanying tomosynthesis reconstructed slices in the same view. An annotation ("3DQuorum") is also burned into the SmartSlices pixel data.

During the initial installation of 3DQuorum software, the system is configured to allow sending of tomosynthesis reconstructed slices and SmartSlices to PACS and diagnostic workstations. The default output groups can be configured to send only SmartSlices, only tomosynthesis reconstructed slices, or both to the preferred devices automatically. The site PACS administrator, in consultation with Hologic connectivity engineers, can select the output formats most appropriate for the site's IT infrastructure and workstations.

As with the introduction of any new imaging technology, Hologic advises testing SmartSlices with your PACS, workstations, and other Imaging Enterprise devices to ensure compatibility.

Workflow

As with any imaging study, the technologist selects the patient and identifies the type of imaging procedure that will be done. For tomosynthesis screening exams with 3DQuorum software, the process of imaging the subject and completing the study is all that is necessary. The 3DQuorum software itself operates with no direct human intervention.

1.11 About Genius Al Detection Software

Genius AI Detection[®] software is a software module intended to identify suspicious breast lesions appearing as soft-tissue densities (masses, architectural distortions, and asymmetries) or calcification clusters in breast tomosynthesis images. Genius AI Detection software analyzes the tomosynthesis data set for each view from a Hologic 3D Mammography[™] exam to identify such lesions using deep learning neural networks. For each identified lesion, Genius AI Detection software produces the following CAD findings:

- Slice where the lesion is best represented.
- Location of the lesion.
- Outline of the lesion.
- Confidence score of the lesion.
- Correlated findings for the lesion (if applicable).

In addition, Genius AI Detection software produces a case score for the entire breast tomosynthesis exam representing the confidence that the exam contains a malignant lesion.

Genius AI Detection software stores all properties of these CAD findings into a DICOM Mammography CAD SR object for storage and display on DICOM-compliant review workstations and archive systems. An option to create a DICOM Secondary Capture Image is available when the review workstations cannot interpret DICOM Mammography CAD SR objects.

Workflow

As with any imaging study, the technologist selects the patient and identifies the type of imaging procedure that will be done. For a Genius 3D[™] Mammography exam, the process of imaging the subject and completing the study is all that is necessary. The Genius AI Detection software itself operates with no direct human intervention.

1.11.1 Warnings and Precautions for Genius AI Detection Software



Important

The radiologist should base interpretation only on diagnostic-quality images and should not solely depend on Genius AI Detection software marks for determining clinical disposition.



Important

Genius AI Detection software is an aid to be used by radiologists concurrently while interpreting Hologic 3D MammographyTM image sets.



Important

Genius AI Detection software does not enhance what the user sees; rather, it helps to identify regions on 3DTM mammograms that should be examined.

1.12 User Profiles

1.12.1 Mammography Technologist

- Meets all requirements that apply to the location in which the Mammography Technologist operates.
- Completed training on the mammography system.
- Has training in mammography positions.
- Knows how to operate a computer and its peripherals.

1.12.2 Radiologist

- Meets all requirements that apply to the location in which the Radiologist operates.
- Knows how to operate a computer and its peripherals.

1.12.3 Medical Physicist

- Meets all requirements that apply to the location in which the Medical Physicist operates.
- Knows about mammography.
- Has experience with digital imaging.
- Knows how to operate a computer and its peripherals.

1.13 Training Requirements

In the United States, users must be Registered Radiologic Technologists meeting criteria to perform mammography. The mammography users must meet all applicable MQSA personnel requirements under FDA guidelines for conventional and digital mammography.

The user has options available for training, which include but are not limited to:

- Onsite applications training by a Hologic Clinical Services Specialist
- Onsite on the job training, also known as peer training

Additionally, the user manual is a guide for directions on how to use the system.

All users must make sure that they receive training on correct operation of the system before use on patients.

Hologic does not accept the responsibility for injury or damage from incorrect system operation.

1.14 Quality Control Requirements

Facilities in the United States must use the Quality Control Manual to create a Quality Assurance and Quality Control program. The facility must create the program to meet the requirements of the Mammography Quality Standards Act or to be accredited by ACR or another accreditation body.

Facilities outside the United States can use the Quality Control Manual as a guide to create a program to meet the local standards and regulations.

1.15 Where to Find the Installation Instructions

Installation instructions are available in the Service Manual.

1.16 Where to Find Technical Description Information

Technical description information is available in the Service Manual.

1.17 Warranty Statement

Except as otherwise expressly stated in the Agreement: i) Equipment manufactured by Hologic is warranted to the original Customer to perform substantially in accordance with published product specifications for one (1) year starting from the date of shipment, or if Installation is required, from the date of Installation ("Warranty Period"); ii) digital imaging mammography x-ray tubes are warranted for twenty-four (24) months, during which the x-ray tubes are fully warranted for the first twelve (12) months and are warranted on a straight-line prorated basis during months 13-24; iii) replacement parts and remanufactured items are warranted for the remainder of the Warranty Period or ninety (90) days from shipment, whichever is longer; iv) consumable Supplies are warranted to conform to published specifications for a period ending on the expiration date shown on their respective packages; v) licensed Software is warranted to operate in accordance with published specifications; vi) Services are warranted to be supplied in a workman-like manner; vii) non-Hologic Manufactured Equipment is warranted through its manufacturer and such manufacturer's warranties shall extend to Hologic's customers, to the extent permitted by the manufacturer of such non-Hologic Manufactured Equipment. Hologic does not warrant that use of Products will be uninterrupted or error-free, or that Products will operate with non-Hologic authorized third-party products. These warranties do not apply to any item that is: (a) repaired, moved, or altered other than by Hologic authorized service personnel; (b) subjected to physical (including thermal or electrical) abuse, stress, or misuse; (c) stored, maintained, or operated in any manner inconsistent with applicable Hologic specifications or instructions, including Customer's refusal to allow Hologic recommended Software upgrades; or (d) designated as supplied subject to a non-Hologic warranty or on a pre-release or "as-is" basis.

1.18 Technical Support

Refer to the copyright page of this manual for contact information for product support.

1.19 Product Complaints

Report any complaints or problems in the quality, reliability, safety, or performance of this product to Hologic. If the device has caused or added to patient injury, immediately report the incident to the Hologic Authorized Representative and Competent Authority of the respective member state or country. The Competent Authorities for medical devices are usually the individual Member States Ministry of Health or an agency within the Ministry of Health. (Refer to the copyright page for Hologic contact information.)

1.20 Hologic Cybersecurity Statement

Hologic continuously tests the current state of computer and network security to examine possible security problems. When necessary, Hologic provides the updates to the product.

For Cybersecurity Best Practices documents for Hologic products, refer to the Hologic Internet site at <u>www.Hologic.com</u>.

1.21 Antivirus Information

Contact your service representative for antivirus installation.

1.22 Symbols

Symbol	Description	Standard
	Caution—radiation	Hologic
	Non-ionizing electromagnetic radiation	IEC 60417, Reference 5140
4	Warning electricity	ISO 7010, Reference W012
4	Dangerous voltage	IEC 60417, Reference 5036
	Warning	ISO 7010, Reference W001
	Caution	ISO 15223-1:2021, Reference 5.4.4
Ó	Emergency stop	Hologic

This section describes the Symbols on this system.

Symbol	Description	Standard
X -	Discard electrical and electronic equipment separately from standard waste. Send decommissioned material to Hologic or contact your service representative.	WEEE Directive 2012/19/EU
*	Type B applied part	IEC 60417, Reference 5840
\ ↓	Equipotentiality	IEC 60417, Reference 5021
	Protective earth (ground)	IEC 60417, Reference 5019
	Fuse	IEC 60417, Reference 5016
\sim	Alternating Current	IEC 60417, Reference 5032
0	"OFF" (power)	IEC 60417, Reference 5008
	"ON" (power)	IEC 60417, Reference 5007
RXONLY	Prescription use only	FDA 21 CFR 801.109
EC REP	Authorized Representative in the European Community	ISO 15223-1:2021, Reference 5.1.2
C E xxxx	CE Mark with notified body reference number	MDR Regulation (EU) 2017/745
MD	Medical device	ISO 15223-1:2021, Reference 5.7.7
NON STERILE	Non-sterile	ISO 15223-1:2021, Reference 5.2.7
www.hologic.com/package-inserts	Consult instructions for use	ISO 15223-1:2021, Reference 5.4.3

Symbol	Description	Standard
Translations in Box	Translations in box	Hologic
B	Recyclable symbol	Unicode, Reference U+267C
Ē.	Provision for the disabled or handicapped persons	ISO 7000, Reference 0100
Ť	Keep dry	ISO 15223-1:2021, Reference 5.3.4
∎ ⊥	Fragile; handle with care	ISO 15223-1:2021, Reference 5.3.1
	Do not stack	ISO 7000:2014, Reference 2402
	Temperature limit	ISO 15223-1:2021, Reference 5.3.7
Temp Logger Activated	Temperature logger activated	Hologic
Temp Logger SN	Temperature logger serial number	Hologic
<u>11</u>	This way up	ISO 7000:2014, Reference 0623
	Do not use if package is damaged	ISO 15223-1:2021, Reference 5.2.8
P/N	Part number	Hologic
REV	Revision	Hologic

Symbol	Description	Standard
Model	Model number	Hologic
REF	Catalog number	ISO 15223-1:2021, Reference 5.1.6
LOT	Batch code	ISO 15223-1:2021, Reference 5.1.5
SN	Serial number	ISO 15223-1:2021, Reference 5.1.7
QTY	Quantity	Hologic
SFW SHIPPED	Software version shipped	Hologic
SFW Version	Software version number	Hologic
CASE	Case (box or package)	Hologic
VC	Vendor code	Hologic
	Manufacturer	ISO 15223-1:2021, Reference 5.1.1
<u></u>	Date of manufacture	ISO 15223-1:2021, Reference 5.1.3
Made in USA	Made in United States of America	Hologic
Patents	Patents	Hologic
	Use-by Date	ISO 15223-1:2021, Reference 5.1.4
\otimes	Do not re-use	ISO 15223-1:2021, Reference 5.4.2

1.23 Descriptions of Warnings, Cautions, and Notes

Descriptions of Warnings, Cautions, and Notes used in this manual:



WARNING!

The procedures that you must follow accurately to prevent possible dangerous or fatal injury.



Warning:

The procedures that you must follow accurately to prevent injury.



Caution:

The procedures that you must follow accurately to prevent damage to equipment, loss of data, or damage to files in software applications.



Note Notes show additional information.

1.24 Document Conventions

When prompted to add text, enter the text printed in **monospaced** font exactly as shown.

Chapter 2 General Information

2.1 System Overview

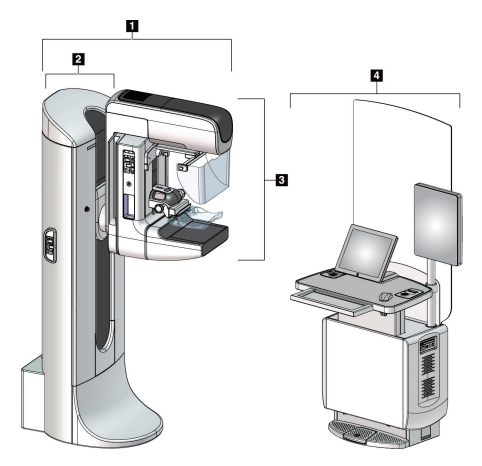


Figure 1: 3Dimensions[™] System

Figure Legend

- 1. Tubestand (Gantry and C-arm)
- 2. Gantry
- 3. C-arm (Tube Arm and Compression Arm)
- 4. Universal Acquisition Workstation

2.1.1 C-Arm Overview

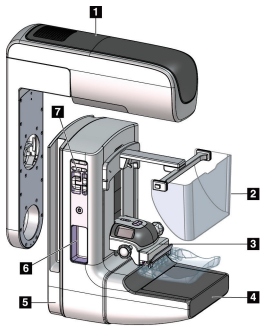


Figure 2: C-arm Overview

Figure Legend

- 1. Tube Arm
- 2. Patient Face Shield
- 3. Compression Device
- 4. Image Receptor
- 5. Compression Arm
- 6. Patient Handle
- 7. C-arm Control Panel

2.2 Safety Information

Read and understand this manual before you use the system. Keep the manual available during system operation.

Always follow all the instructions in this manual. Hologic does not accept responsibility for injury or damage from incorrect system operation. For training options, contact your Hologic representative.

The system has safety interlocks, but the user must understand how to operate the system safely. The user must also understand the health hazards of x-ray radiation.

2.3 Warnings and Precautions



This system is classified as CLASS I, TYPE B APPLIED PART, IPX0, permanently connected equipment, continuous operation with short term loading per IEC 60601-1. There are no special provisions to protect the system from flammable anesthetics or ingress of liquids.



WARNING!

Risk of electric shock. Only connect this equipment to supply mains with Protective Earth.



WARNING!

For North American electrical safety requirements, use a Hospital Grade receptacle to supply a correct Ground.



WARNING!

Electrical equipment used near flammable anesthetics can cause an explosion.



WARNING!

To correctly isolate the system, attach only approved accessories or options to the system. Only approved personnel can change the connections.



WARNING!

Keep a 1.5 meter safe distance between the patient and any non-patient devices.

Do not install non-patient system components (like the Workflow Manager, a diagnostic review workstation, or a hard copy printer) in the Patient Area.



WARNING!

Only trained Service Engineers authorized through Hologic can open any of the panels. This system contains lethal voltages.



WARNING!

The user must correct problems before the system is used. Contact an approved service representative for preventive maintenance.



WARNING!

After power failure, remove the patient from the system before you apply power.



Warning:

This device contains dangerous material. Send decommissioned material to Hologic or contact your service representative.



Warning:

C-arm movement is motorized.



Warning:

You increase the patient dose to high levels when you increase the AEC exposure adjustment. You increase the image noise or decrease image quality when you decrease the AEC exposure adjustment.



Warning:

Control the access to the equipment according to local regulations for radiation protection.



Warning:

The disk drives installed in this system are a Class I Laser Product. Prevent direct exposure to the beam. Hidden laser radiation exists if the case to a disk drive is open.



Warning:

The bar code scanner installed in this system is a Class II Laser Product. Prevent direct exposure to the beam. Hidden laser radiation exists if the cover is opened.



Warning:

Keep your full body behind the radiation shield during the exposure.



Warning:

Do not move the C-arm while the system retrieves the image.



Warning:

If a paddle touches possible infectious materials, contact your Infection Control Representative to remove contamination from the paddle.



Warning:

Do not leave the patient unattended during the procedure.



Warning:

Keep the hands of the patient away from all buttons and switches at all times.



Warning:

Place each footswitch in a position where, when used, they remain in reach of the Emergency Off Switches.



Warning:

Position the footswitches to prevent accidental operation by a patient or wheelchair.



Warning:

To prevent a higher radiation dose to the patient, only put approved materials in the x-ray beam path.



Warning: This system can be dangerous to the patient and the user. Always follow the safety precautions for x-ray exposures.



Warning:

For exposures except magnification case studies, always use the Face Shield.



Warning:

The Face Shield does not protect the patient from radiation.



Warning:

Risk of entrapment. Make sure that the C-arm has 50 cm (20 inches) of clearance to any object during C-arm rotation. Do not use Auto Rotation when C-arm clearance is less than 50 cm (20 inches).



Caution:

The system is a medical device and not a normal computer. Only make approved changes to the hardware or software. Install this device behind a firewall for network security. The computer virus protection or network security for this medical device is not supplied (for example, a computer firewall). The network security and anti-virus provisions are the responsibility of the user.



Caution:

Do not turn off the Acquisition Workstation circuit breaker except in emergency. The circuit breaker can turn off the Uninterruptible Power Supply (UPS) and risk data loss.



Caution:

Risk of data loss. Do not put any magnetic media near or on devices that create any magnetic fields.



Caution:

Do not use any heat source (like a heating pad) on the image receptor.



Caution:

To prevent possible damage from thermal shock to the Digital Image Receptor, follow the recommended procedure to turn off the equipment.

Caution: The disp

The display is calibrated for compliance to DICOM standards. Do not make any brightness or contrast adjustments to the display.



Caution:

Use the least possible amount of cleaning fluids. The fluids must not flow or run.



Caution:

To prevent damage to the electronic components, do not use disinfectant sprays on the system.



Note:

Hologic does not supply the Gantry power cable for some countries. If the power cable is not supplied, the installed cable must meet the following requirements and all local codes that apply: 3 conductor, 8 AWG (10 mm²) copper not more than 25 feet (7.62 meters) in length.

2.4 Emergency Off Switches

The Emergency Off switches remove the power from the Gantry and the Acquisition Workstation Lift Mechanism (on the workstations that include that option). Do not routinely use the Emergency Off switches to turn off the system.

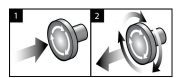


Figure 3: Emergency Off Switch Functionality

There are three Emergency Off switches, one on each side of the Gantry and one on the Acquisition Workstation.

- To turn Off the Gantry and disable the Acquisition Workstation Lift Mechanism, press any of the Emergency Off switches.
- 2. To reset the Emergency Off switch, turn clockwise approximately one-quarter turn until the switch pops back out.

2.5 Interlocks

The system has safety interlocks:

- C-arm vertical movement and rotation are disabled when compression force is applied. A Service Engineer can configure the lockout force from 22 Newtons (5 pounds) to 45 Newtons (10 pounds).
- If the x-ray button and/or x-ray footswitch* is released before the end of the exposure, the exposure stops and an alarm message is displayed.

* (The x-ray footswitch option is available only on the Universal Acquisition Workstation console.)

- When in Tomo mode, the system does not allow an exposure if the Grid is in the x-ray field (Tomosynthesis option).
- Mirror and Filter position flags also prevent x rays when the Light Field Mirror or the Filter Assembly is not correctly positioned.

2.6 Compliance

This section describes the mammography system compliance requirements and the responsibilities of the manufacturer.

2.6.1 Compliance Requirements

The manufacturer has the responsibility for the safety, reliability, and performance of this equipment with the following provisions:

- The electrical installation of the room meets all requirements.
- The equipment is used according to the *User Guide*.
- The assembly operations, extensions, adjustments, changes, or repairs are performed only by authorized persons.
- The network and communication equipment is installed to meet IEC Standards. The complete system (network and communications equipment and the mammography system) must be in compliance with IEC 60601-1.



Caution:

Medical Electrical Equipment needs special precautions about EMC and must be installed, put into service and used according to the EMC information provided.



Caution:

Portable and mobile RF communications can affect medical electrical equipment.



Caution:

The use of unauthorized accessories and cables can result in increased emissions or decreased immunity. To keep the isolation quality for the system, attach only approved Hologic accessories or options to the system.



Caution:

The Medical Electrical (ME) Equipment or ME System should not be used adjacent to or stacked with other equipment. If adjacent or stacked use is necessary, make sure that the ME Equipment or ME System operates correctly in this configuration.



Caution:

This system is intended for use by healthcare professionals only. This system may cause radio interference or may disrupt the operation of nearby equipment. It may be necessary to take mitigation measures, such as re-orienting or relocating the equipment or shielding the location.



Caution:

Changes or modifications not expressly approved by Hologic could void your authority to operate the equipment.



Caution:

This equipment has been tested and found to comply with the limits for a Class A digital device, pursuant to part 15 of the FCC Rules. These limits are designed to provide reasonable protection against harmful interference when the equipment is operated in a commercial environment. This equipment generates, uses, and can radiate radio frequency energy and, if not installed and used in accordance with the instruction manual, may cause harmful interference to radio communications. Operation of this equipment in a residential area is likely to cause harmful interference at his own expense.



Caution:

Based on ENCLOSURE PORT IMMUNITY to RF WIRELESS COMMUNICATIONS EQUIPMENT test level per IEC 60601-1-2 ed. 4, this device shows susceptibility to GMRS 460, FRS 460 transmission frequency. Provide a minimum separation of 30 cm between GMRS and FRS radios and the system.

2.6.2 Compliance Statements

The manufacturer states this device is made to meet the following requirements:



Medical – Applied electromagnetic radiation equipment as to electrical shock, fire and mechanical hazards only in accordance with ANSI/AAMI ES 60601-1 (2005) + A1: 2012, C1: 2009 / (R)2012, A2: 2010 / (R)2012 and CAN/CSA-C22.2 No. 60601-1 (2014)

- CAN/CSA ISO 13485-03 Medical Devices Quality Management Systems Requirements for Regulatory Purposes (Adopted ISO 13485:2003 second edition, 2003-07-15)
- CAN/CSA C22.2 NO. 60601-1:2014 Medical Electrical Equipment Part 1: General Requirements for Basic Safety and Essential Performance
- EN 60601-1: 2006 / A1: 2013 Medical Electrical Equipment. General Requirements for Basic Safety and Essential Performance
- ETSI EN 300 330-1: V1.3.1, and ETSI EN 300 330-2: V1.5.1: 2006—Electromagnetic compatibility and Radio spectrum Matters (ERM); Short Range Devices (SRD); Radio equipment in the frequency range 9 kHz to 25 MHz and inductive loop systems in the frequency range 9 kHz to 30 MHz
- ETSI EN 301 489-1: V1.6.1, and ETSI EN 301 489-3: V1.8.1: 2008—Electromagnetic compatibility and Radio spectrum Matters (ERM); ElectroMagnetic Compatibility (EMC) standard for radio equipment and services
- FCC, 47 CFR Part 15, Subpart C, Section 15.225: 2009
- FDA, 21 CFR [Parts 900 and 1020]
- IEC 60601-1: 2005 / A1: 2012 ed 3.1 Medical Electrical Equipment Part 1: General Requirements for Basic Safety and Essential Performance
- IEC 60601-1-2 Ed. 4th: 2014 Medical Electrical Equipment Part 1-2: General Requirements for Basic Safety and Essential Performance Collateral Standard: Electromagnetic Compatibility Requirements and Tests
- IEC 60601-1-3 Ed. 2.0: 2008 / A1: 2013 Medical Electrical Equipment Part 1-3: Particular Requirements for the Safety of X-Ray source assemblies and X-Ray tube assemblies for medical diagnosis
- IEC 60601-2-28: 2017 Medical Electrical Equipment Part 2-28: Particular Requirements for the Basic Safety and Essential Performance of X-ray Tube Assemblies for Medical Devices
- IEC 60601-2-45: 2011 / AMD1: 2015 Medical Electrical Equipment Part 2-45: Particular Requirements for the Basic Safety and Essential Performance of Mammographic X-Ray Equipment and Mammographic Stereotactic Devices
- RSS-210: Issue 7, 2007 Radio Standards Specification Low-power License-exempt Radiocommunication Devices: Category I Equipment
- ANSI/AAMI ES 60601-1: +A1: 2012, C1: 2009 / (R)2012 and A2: 2010 / (R)2012 Medical Electrical Equipment, Part 1: General Requirements for Basic Safety and Essential Performance, includes amendment (2010)

2.7 System Label Locations

For the location of the system labels and serial numbers, refer to the figures that follow.



Figure 4: Location of the System Label for the Gantry



Figure 5: Location of the System Label for the Universal Acquisition Workstation

Chapter 3 System Controls and Indicators

3.1 System Power Controls

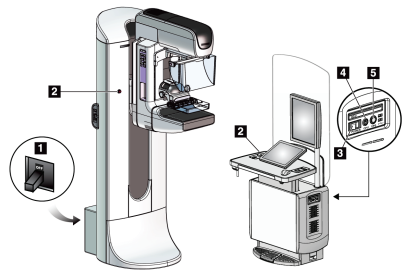
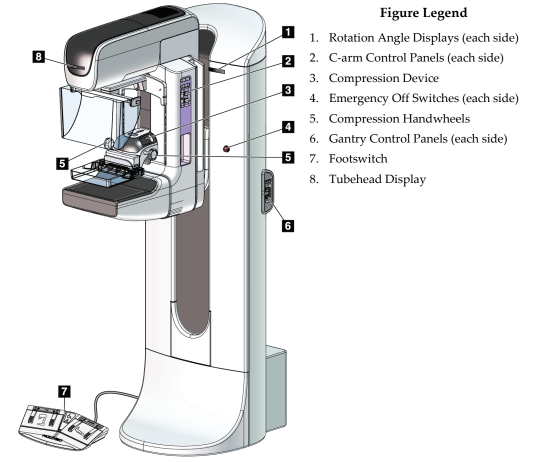


Figure 6: System Power Controls

Figure Legend

- 1. Gantry Power Circuit Breaker
- 2. Emergency Off Switch (two on the Gantry, one on the Acquisition Workstation)
- 3. Acquisition Workstation Power Switch
- 4. Computer Power On/Reset Button
- 5. Uninterruptible Power Supply (UPS) Power Button



3.2 Tubestand Controls and Indicators

Figure 7: Tubestand Controls and Indicators

3.2.1 Tubehead Display

The Tubehead Display shows:

- SID
- Filter Type
- Collimator Setting
- Paddle Position



Figure 8: Tubehead Display

3.2.2 Compression Device Controls and Display

Figure Legend

- 1. Manual Compression Handwheels
- 2. Paddle Shift Buttons
- 3. AEC Sensor Buttons
- 4. Compression Device Display
- 5. FAST Compression Mode Slide
- 6. Paddle Clamp

The Compression Device Display shows:

- AEC Sensor Position
- Compression Force (displays 0.0 when force is less than 4 pounds)
- Compression Thickness
- Angle of the C-arm after rotation (for 5 seconds)

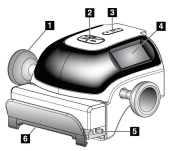


Figure 9: Compression Device



Figure 10: Compression Device Display

3.2.3 C-Arm Control Panels

There is a C-arm control panel on both the left side and right side of the C-arm. These buttons provide the Collimator and C-arm functions.



Figure 11: C-arm Control Panel

Figure Legend

- 1. MLO Rotation
- 2. C-arm Zero
- 3. Light Field Lamp
- 4. Motor Enable
- 5. Collimator Override
- 6. Clockwise C-arm Rotation
- 7. C-arm Up and Down
- 8. Counterclockwise C-arm Rotation
- 9. Compression Up
- 10. Compression Down

3.2.4 Gantry Control Panels

Control panels on the Gantry provide additional access to C-arm functions. There is a C-arm control panel on both the left side and right side of the Gantry.



Figure 12: Gantry Control Panel

Figure Legend

- 1. Motor Enable
- 2. C-arm Zero
- 3. Clockwise C-arm Rotation
- 4. C-arm Up and Down
- 5. Counterclockwise C-arm Rotation

3.2.5 Dual Function Footswitch



Warning:

Place each footswitch in a position where, when used, they remain in reach of the Emergency Off Switches.



Warning:

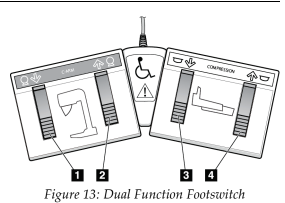
Position the footswitches to prevent accidental operation by a patient or wheelchair.

To use the footswitch:

- Press the footswitch to actuate.
- Release the footwitch to stop the movement.

Figure Legend

- 1. C-arm Down
- 2. C-arm Up
- 3. Compression Down
- 4. Compression Up



3.3 Universal Acquisition Workstation Controls and Displays



Note

Hologic configures some systems to meet specific requirements. Your system configuration may not have all the options and accessories included in this manual.

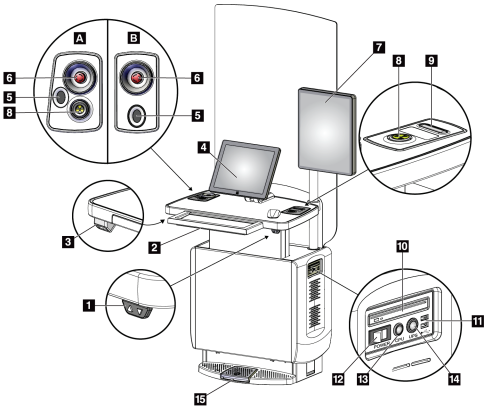


Figure 14: Universal Acquisition Workstation Controls and Displays

Figure Legend

- 1. Height Adjustment Switch
- 2. Keyboard (in drawer)
- 3. Bar Code Scanner
- 4. Control Monitor
- 5. Fingerprint Scanner*
- 6. Emergency Off Switch*
- 7. Image Display Monitor
- 8. X-ray Activation Button*
- 9. Compression Release Button

- 10. CD/DVD Drive
- 11. USB Ports
- 12. Acquisition Workstation Power Switch
- 13. Computer Power On/Reset Button
- 14. Uninterruptible Power Supply (UPS) Power Button
- 15. X-ray Footswitch

*A = series II Universal Acquisition Workstation layout; B = series I Universal Acquisition Workstation layout

\sum

Note

The controls for the Universal Acquisition Workstation installed in a mobile environment are the same as the controls for the Universal Acquisition Workstation.

Chapter 4 Startup, Functional Tests, and Shutdown

4.1 How to Start the System



Note

If the system remains on overnight, reboot the system daily to guarantee best performance.

- 1. Make sure that there are no obstructions to C-arm movement or to the view of the Operator.
- 2. Make sure that all three Emergency Off switches are in the reset position (unpushed).



Figure 15: Turn to Reset the Emergency Off Switches

- 3. Make sure that the Gantry circuit breaker is in the ON position.
- 4. If the UPS was turned OFF, press the UPS power button to apply power to the UPS (see the following figure).

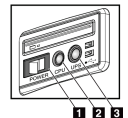


Figure 16: Universal Acquisition Workstation Power Buttons

Figure Legend

- 1. Acquisition Workstation Power Switch
- 2. Computer Power On/Reset Button
- 3. UPS Power Button
- 5. Turn ON the Acquisition Workstation power switch (see the previous figure).

6. Press the computer power button (see the previous figures). The computer powers on and the *Windows 10 Login* screen opens on the Acquisition Workstation control monitor.

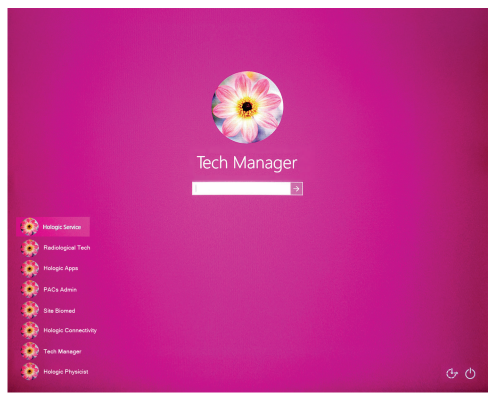


Figure 17: Windows 10 Login Screen

7. Select your user name and enter your password.

8. The *Startup* screen for the 3Dimensions system opens on the Acquisition Workstation control monitor. The Gantry then automatically powers on.

Shutdown Reboot Log Out

3Dimensions[™]

Figure 18: Startup Screen

No User



Note

To log out of the Windows 10 operating system, select the Log Out button.



Note

The *Startup* screen includes a **Shutdown** button that turns off the system, and a **Reboot** button that restarts the system.



Note

The system can require between five minutes and fifteen minutes to prepare for image acquisition. The wait time depends on the detector power configuration. A timer in the taskbar displays the wait time before the system is ready. Do not acquire clinical or QC images unless the System Status Icon indicates that the system is Ready.

4.2 Log In

- 1. Select the **Log In** button on the *Startup* screen.
- 2. The *Select an Operator* (System Log In) screen opens and shows a list of Manager and Technologist user names. If necessary, select the **Show All** button to list the Service, Applications, and Physicist user names.

Select an Operator or Authenticate	Fingerprint						
Manager, Tech							Log In
Tech, Radiological							
							Show All
							Username
	Password						
							Exit
0 🚺 No User	: 0	ي 54	۹ ۵	e 🖉	🥥 🥥	P	11:15:37 AM

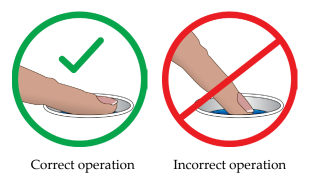
Figure 19: System Log In Screen

- 3. Use one of the following methods to log into the system:
 - Select the applicable user name button. Type in your password, then select the **Log In** button.
 - Select the **Username** button. Type in your user name and password, then select the **Log In** button.
 - Validate your fingerprint by pressing your finger in the fingerprint scanner.



Note Most of the unique rep

Most of the unique, repeatable fingerprint information is from the pad of the finger, not the fingertip. Flat finger placement as shown in the left image results in fast and accurate fingerprint identification.





Note

If Quality Control tasks are due, the *Select Function to Perform* screen opens. You can perform quality tasks or you can select **Skip**.

4.3 Perform the Functional Tests

To make sure that the controls operate correctly, do the Functional Tests as part of your monthly visual checklist. Make sure that all buttons work correctly (Gantry and C-arm control panels, left and right sides).

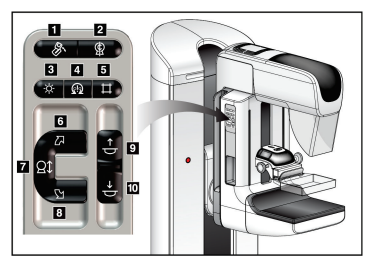


Figure Legend

- 1. MLO Rotation
- 2. C-arm Zero
- 3. Light Field Lamp
- 4. Motor Enable
- 5. Collimator Override
- 6. Clockwise C-arm Rotation
- 7. C-arm Up and Down
- 8. Counterclockwise C-arm Rotation
- 9. Compression Up
- 10. Compression Down

Figure 20: C-arm Control Panel (left side shown)



Note

C-arm vertical movement and rotation are disabled when compression force is applied. A Service Engineer can configure the lockout force from 22 Newtons (5 pounds) to 45 Newtons (10 pounds).

4.3.1	Compression	Functional Tests
-------	-------------	-------------------------

Table 1: Compression Tests		
Function	Functional Test	
Compression Down	 Press a Compression Down button: The compression brake engages. The light field lamp illuminates. The compression device lowers. Note When you press the Compression Down button, the compression brake remains engaged until the Compression Release button is pressed. Compression down movement stops: When you release the button. When you reach the Down Force limit. When you reach the lower travel limit. 	
Compression Up	 Press a Compression Up button: The Compression Device moves toward the top. Note The Compression Up button does NOT release the Compression Brake. Compression up movement automatically stops: When you release the button. When you reach the upper travel limit. 	

Tab	le 2: C-arm Up and Down Movement
Function	Functional Test
C-arm Up	 Press the C-arm Up button. C-arm movement stops when the button is released. C-arm movement stops when the C-arm reaches the upper travel limit. C-arm vertical movement and rotation are disabled when compression force is applied. A Service Engineer can configure the lockout force from 22 Newtons (5 pounds) to 45 Newtons (10 pounds).

4.3.2 C-Arm Movement Functional Tests

C-Arm Up and Down Movement

FunctionFunctional TestC-arm DownPress the C-arm Down button.•C-arm movement stops when the button is released.•C-arm movement stops when the C-arm reaches the lower travel limit.•C-arm vertical movement and rotation are disabled when compression force is applied. A Service Engineer can configure the lockout force from 22 Newtons (5 pounds) to 45 Newtons (10 pounds).
 C-arm movement stops when the button is released. C-arm movement stops when the C-arm reaches the lower travel limit. C-arm vertical movement and rotation are disabled when compression force is applied. A Service Engineer can configure the lockout force from

Table 2: C-arm Up and Down Movement

C-Arm Rotation

Function	Functional Test
Counterclockwise C-arm Rotation	Press the Counterclockwise C-arm Rotation button (item 1) to start counterclockwise C-arm rotation. C-arm movement stops when the switch is released. To fine-tune the C-arm rotation in one degree segments, hold the button down for a second, then let go quickly. Repeat as necessary.

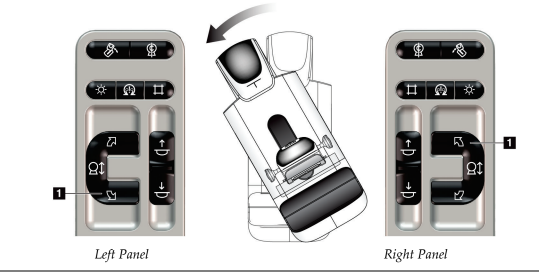
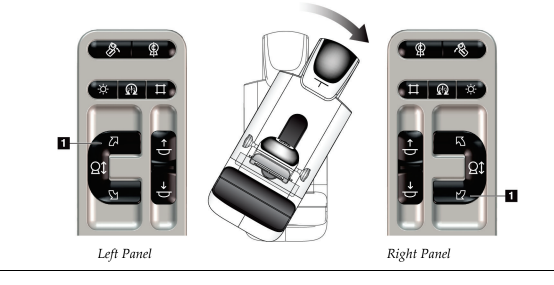
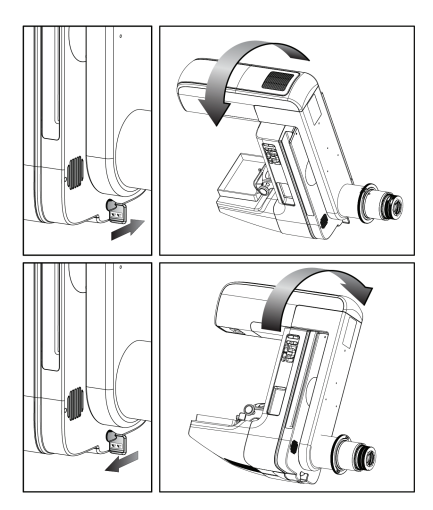


Table 4: C-arm Clockwise Rotation		
Function	Functional Test	
Clockwise C-arm Rotation	Press the Clockwise C-arm Rotation button (item 1) to start clockwise C-arm rotation. C-arm movement stops when the switch is released. To fine-tune the C-arm rotation in one degree segments, hold the button down for a second, then let go quickly. Repeat as necessary.	



Function	Functional Test
C-arm Rotation Switch	Push the C-arm Rotation switch away from you to move the C-arm toward you. Pull the C-arm Rotation switch toward you to move the C-arm away from you. C-arm movement stops when the switch is released.

Table 5: C-arm Rotation Switch



C-Arm Automatic Rotation (Auto-Rotate)



Warning:

Risk of entrapment. Make sure that the C-arm has 50 cm (20 inches) of clearance to any object during C-arm rotation. Do not use Auto Rotation when C-arm clearance is less than 50 cm (20 inches).

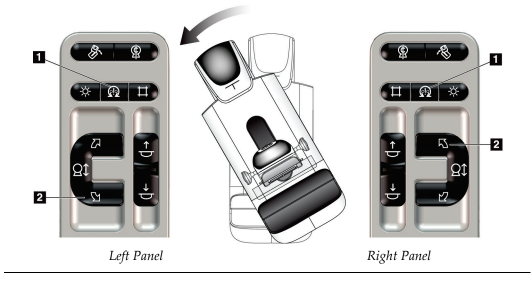


Note To stop the C-arm automatic rotation movement, press any button or the Emergency

Stop switch.

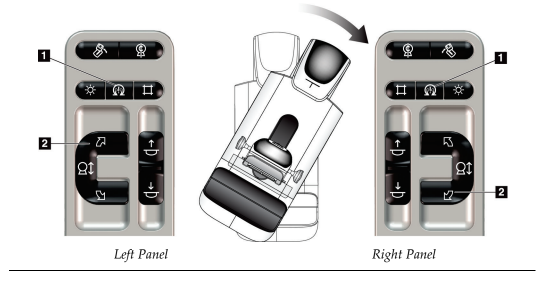
Motor Enable	Press the Motor Enable button and a Rotation button at the same time to enable auto-rotation movement clockwise or counterclockwise.
C-arm Zero	Press the Motor Enable button and the C-arm Zero button at the same time. Release the buttons. The machine beeps and the C-arm returns automatically to the zero position.

Function	Functional Test
Automatic	Press the Motor Enable button (item 1) and the
Counterclockwise C-arm	Counterclockwise C-arm Rotation button (item 2) at the
Rotation	same time. Release the buttons. The machine beeps and
	the C-arm moves automatically counterclockwise.



10010	7. Automatic C-urm Clockwise Rotation
Function	Functional Test
Automatic Clockwise	Press the Motor Enable button (item 1) and the
C-arm Rotation	Clockwise C-arm Rotation button (item 2) at the same
	time. Release the buttons. The machine beeps and the
	C-arm moves automatically clockwise.

Table 7: Automatic C-arm Clockwise Rotation



MLO Automatic Rotation

Table 8: Automatic MLO Rotation		
Function	Functional Test	
Automatic Tubehead Rotation to Zero Position	Position the C-arm at an angle greater than 15 degrees. Press the MLO Rotation button (item 1). The tubehead moves automatically to the zero position. The Compression Arm stays in its angled position. Press the MLO Rotation button again to return the tubehead to its original position.	



4.3.3 Collimation

Table 9: C-arm Collimation		
Function	Functional Test	
Collimator Override	The Collimator Override button changes the collimation through the different x-ray fields. Press the Light Field Lamp button to show the x-ray field, then press the Collimator Override button to select an x-ray field.	
Light Field Lamp	For compression paddles: Press the Light Field Lamp button to see the x-ray field for two minutes. For localization paddles: Press the Light Field Lamp button to see the x-ray field for the configured amount of time. The default time for biopsy or needle localization paddles is ten minutes. Press the Light Field Lamp button again to turn off the light field lamp. The light field lamp automatically illuminates with the start of the Compression Down movement.	

4.3.4 Shifting Paddles



Note

Most paddles can be used with the Shifting Paddle function. The large 24 x 29 cm Frameless Screening Paddle, the large 24 x 29 cm SmartCurve[®] system paddle, and the Magnification paddles are not compatible with the Shifting Paddle function.

Table 10: Shifting Paddle				
Function	Functional Test			
Paddle Shift	The 18 x 24 cm Screening Paddle moves approximately 2.5 cm into the left, center, or right position. While compression is applied, you cannot move the paddle. The collimator is programmed to follow the position of the paddle.			
	To test this function:1. Install the 18 x 24 cm paddle in the Compression Device.			
	2. Select a view.			
	3. Use the Paddle Shift buttons on the <i>Procedure</i> screen to override the position.			
	4. Verify that the paddle automatically moves to the new position.			
	5. Turn on the light field lamp.			
	Confirm that the collimator position matches the paddle position.			
	7. Repeat this procedure for the other paddle positions.			

Use the 18 x 24 cm Frameless Screening Paddle to test the Shifting Paddle function.

4.4 Emergency Off Switches Functionality

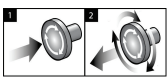


Figure 21: Emergency Off Switch Functionality

There are three Emergency Off switches, one on each side of the Gantry and one on the Acquisition Workstation.

- To turn Off the Gantry and disable the Acquisition Workstation Lift Mechanism, press any of the Emergency Off switches.
- 2. To reset the Emergency Off switch, turn clockwise approximately one-quarter turn until the switch pops back out.

4.5 How to Turn Off the System

- 1. Close any open patient procedures. Refer to <u>Close a Patient</u> on page 76.
- 2. From the *Select Patient* screen, select the **Log Out** button.
- 3. From the *Startup* screen, select the **Shutdown** button.
- 4. Select **Yes** in the confirmation dialog box.

4.6 How to Remove All Power from the System

See the following figure for illustrations of the buttons and switches referenced in the following procedures.

3. UPS Power Button

Figure Legend

1. Acquisition Workstation Power Switch

2. Computer Power On/Reset Button

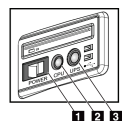


Figure 22: Universal Acquisition Workstation Power Buttons

To remove all power from the system:

- 1. Turn off the system.
- 2. If your system includes the UPS, press the UPS button (item 3).
- 3. Turn OFF the acquisition workstation power switch (item 1).
- 4. Unplug the acquisition workstation power cable from the AC outlet.
- 5. Turn OFF the Gantry circuit breaker.
- 6. Turn OFF the Facility Mains circuit breaker.

Chapter 5 User Interface

5.1 Select Function to Perform Screen

After login, the *Select Function to Perform* screen opens. This screen shows the quality control tasks that are due.



Note

The *Select Patient* screen opens when no quality control tasks are scheduled to be done.

Name	Last Performed	Due Date	
DICOM Printer Quality Control		4/11/2017	Skip
Gain Calibration	2/23/2017	2/27/2017	
CEDM Gain Calibration	3/1/2017	3/6/2017	
Artifact Evaluation		4/11/2017	Start
Phantom Image Quality		4/11/2017	Start
SNR/CNR		4/11/2017	Mark Complete
Compression Thickness Indicator		4/11/2017	
Diagnostic Review Workstation Quality Control		4/11/2017	
/iewboxes and Viewing Conditions		4/11/2017	
/isual Checklist		4/11/2017	
Reject Analysis		4/11/2017	
Repeat Analysis		4/11/2017	
Compression Test		4/11/2017	
AEC Testing		4/11/2017	
			Admin
		Number of results: 14	Log Out

Figure 23: An Example Select Function to Perform Screen

To Complete a Scheduled Quality Control Task:

- 1. Select a Quality Control task from the list.
- 2. Select the **Start** button. Follow the messages to complete the procedure. (The **Start** button is not available for all types of tests.)

- OR -

Select the **Mark Completed** button to mark the status of this procedure as finished. Select **Yes** to confirm the selected procedure is completed.

3. Select End QC.

Note

Note

To Proceed Without Completing All Scheduled Quality Control Tasks:

If all Quality Control tasks are not completed at this time, select the Skip button.



If you select the **Skip** button, the *Select Patient* screen opens. Refer to <u>Select Patient Screen</u> on page 59 for information about this screen.

If you select the **Admin** button, the *Admin* screen opens. Refer to <u>Admin Screen</u> on page 139 for information about this screen.



Quality Control tasks are available to complete at any time. Select the **Admin** button then the **Quality Control** button to access the list of Quality Control tasks.

5.2 About the Taskbar

The taskbar at the bottom of the screen displays additional icons. Select the icon to access information or perform system tasks.



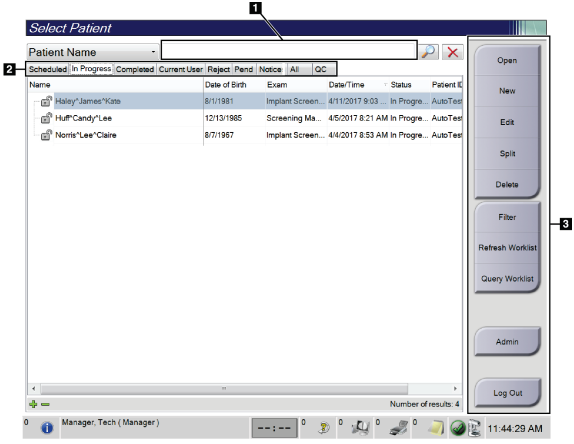
Figure 24: Taskbar

Table 11:	Taskbar Menus

	Description	Menu	
1	Information Icon	No Alarms	
	Select the Information icon to display the Alarms menu.	Acknowledge All	
	This section of the taskbar flashes a yellow color when an alarm exists.	Manage Alarms	
	Select Acknowledge All to stop the flashing indication.		
	Select Manage Alarms to display and close any open alarms.		
2	Current User Name	Users Menu	
	Select the user name section of the taskbar to display the Users Menu.	Log Out	
	Log Out returns you to the <i>Startup</i> screen.	My Settings	
	My Settings lets you select user settings and workflow preferences.	Print	
	Print sends the displayed patient list to a connected printer.		

	Table 11: Taskbar Menus Description	Menu
	Reading Priority IndicatorThe Reading Priority Indicator is displayed if you have Genius AIDetection software. The icon color changes to yellow if there arepatients marked as a high priority for reading per Genius AI Detectionsoftware results. Select the Reading Priority icon to show the list ofpatients that have been marked with high Reading Priority by GeniusAI Detection software. Select the X button to clear the patient from theReading Priority list. Refer to About Genius AI Detection Softwareon page 9 for more information.Patient NameCompletion TimePatient, Test 200.02:55 agoPatient, Test 200.02:44 agoTest, Patient00.01:52 ago	
02:35	Contrast Agent Timer The Contrast Agent Timer is displayed if you have <u>I-View Contrast</u> <u>Enhanced 2D Images</u> on page 100. The timer is activated during a 2D Contrast procedure. The timer shows the amount of time remaining in each contrast phase. The color of the timer indicates the contrast phase. (Initial) Yellow = Waiting Period, contrast agent is not fully dispersed. Green = Optimal Imaging Period, contrast agent is fully dispersed. (Latter) Yellow = Late Period, contrast agent is dissipating. Output Device Icons	
	Select any output device icon to display the <i>Manage Queues</i> screen. This screen displays the status of jobs in the queue, job information for the selected output, and allows you to filter the queue display. The number on the icon shows the number of jobs that remain in the queue.	
4	Notices Icon Select the Notices icon to display the <i>Patients with Unviewed Notices</i> screen. The number on the icon shows the number of patients with Unviewed Notices.	

	Description	Menu
5	System Status Icons	No Alarms
	If the System Status (tubehead) icon has a green check mark next to it, the system is ready for use. Select the System Status icon to display the Faults menu.	Clear All Faults X-Ray Tube, 0 Degrees X-Ray Tube, -15 Degrees X-Ray Tube, +15 Degrees
E min	If the System Status icon is red with a number next to it, the system needs to wait the number of minutes shown before the next image can safely be taken.	System Diagnostics System Defaults About
	If the System Status icon has a yellow exclamation mark next to it and this taskbar section flashes yellow, there is an error. Select the System Status icon for more information about the error. Clear All Faults deletes all fault messages.	
	X-ray Tube, 0 Degrees puts the tubehead at zero degrees of rotation for the next exposure.	
	X-ray Tube, -15 Degrees puts the tubehead at -15 degrees of rotation for an exposure.	
	X-ray Tube, +15 Degrees puts the tubehead at +15 degrees of rotation for an exposure.	
	System Diagnostics accesses Subsystem settings.	
	System Defaults opens the <i>Gantry Defaults</i> screen to set the Compression and Generator default values.	
	About displays information about the Acquisition Workstation (refer to <u><i>About Screen</i></u> on page 141).	



5.3 Select Patient Screen

Figure 25: Select Patient Screen

Table 12: The Select Patient S	Screen
--------------------------------	--------

Item	Description	
1. Quick Search	Search the selected tab for the Patient Name, Patient ID, or Accession Number.	
2. Tabs	The tabs at the top of the screen are configurable. A user with the correct permissions can delete tabs and create new tabs.	
	• Scheduled tab shows the scheduled procedures.	
	• In Progress tab shows the procedures not complete.	
	• Completed tab shows the completed procedures.	
	• Current User tab shows the procedures for the current operator.	
	• Reject tab shows the procedures with rejected views.	
	• Pend tab shows the procedures with pending views.	

Item	Description					
	All tab shows all procedures for all users.					
	 Notices tab shows the patients with unviewed notices by default. Refer to <u>About the Notices Tab</u> on page 61 for complete information. 					
	• QC tab shows the Qua	lity Contro	ol procedure	es.		
	• Enterprise tab is displayed if you have the <u>Advanced Workflow Manager</u> on page 67. This tab shows a list of all studies acquired on systems within an Advanced Workflow Manager cluster. Distribution of the studies is managed from this tab.					
	Scheduled In Progress Completed Curre	1	1	h	Enterprise	
	Name	Date of Birth	Exam	Patient ID		Date/Time
		1/1/2001			DIMENSIONS	6/22/2011 1
	Figure 26: Enterprise Tab					
3. Buttons	Many functions are accessed from this screen by selecting a particular button:					
• Open : Open the selected patient.						
	 New: Add a new patient — refer to <u>Add a New Patient</u> on page 61. Edit: Edit the patient information — refer to <u>Edit the Patient Information</u> on page 62. Split: Remove the images from a procedure or a patient and move the images to a different procedure or patient — refer to <u>Split the Patient</u> <u>Records</u> on page 62. 					e 61.
• Delete : Delete the selected patient — refer to <u>Delete a Patient</u>					<u>e a Patient</u> c	on page 64.
	• Filter : Use a Patient Fil	lter — refe	r to <u>Filters f</u> a	or Patie	<u>nts</u> on pag	e 64.
	 Refresh Worklist: Update the Scheduled Patient Worklist information refer to <u>Refresh the Worklist</u> on page 66. 				ormation	
	Query Worklist: Searce to <u>Query the Worklist</u> or	-	ient in the N	/lodali	ty Worklis	t — refer
	• Admin: Access the Adr		– refer to	<u>Admin</u>	Screen on p	oage 139.
	• Log Out: Exit the syste				-	

5.3.1 About the Notices Tab

- After selecting the **Notices** tab from the *Select Patient* screen, a list of the Patients with Notices is shown.
- The default value for the list is patients with Viewed and Unviewed Notices.
- You can change the value to show patients with Viewed Notices or patients with Unviewed Notices.
- Select **Open** to access the *Procedure* screen for the selected patient.

5.3.2 Open a Patient

- 1. Select a tab to show the desired list of patients.
- 2. Select a patient from the list. The **Open** button becomes active.
- 3. Select **Open** to access the *Procedure* screen for that patient.

5.3.3 Add a New Patient

1. In the Select Patient screen, select the New button. The Add Patient screen opens.

Add Patient		
Last	Ор	en
First		
Middle		
Patient ID*		
Date of Birth*		
Gender*	Female	
Accession Number		
Procedure*	Conventional -	
	Standard Screening - Conventional	
	Ва	ick
(i) Manager, Tech (Manager)	: ⁰) ⁰ 💭 ⁰ <i>I</i> ⁰ <i>I</i> ⁰ <i>I</i> ⁰	8:02 AN

Figure 27: Add Patient Screen

- 2. Enter the new patient information and select a procedure.
- 3. Select the **Open** button. The *Procedure* screen for the new patient opens.

5.3.4 Edit the Patient Information

- 1. In the *Select Patient* screen, select the patient name, then select the **Edit** button.
- 2. In the *Edit Patient* screen, make changes, then select the **Save** button.
- 3. Select **OK** to the *Update Successful* message.

5.3.5 Split the Patient Records

The Split function lets you move images when the images were acquired in an incorrect procedure or an incorrect patient.



You cannot split images for protected patients.



Note

Note

Archived patient records still need correction on PACS or other archive systems after using the Split function.

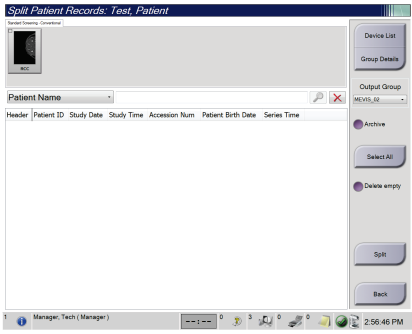


Figure 28: Split Patient Records Screen

Images Acquired in the Wrong Procedure

- 1. In the Select Patient screen, select the patient.
- 2. Select the **Split** button. The *Split Patient Records* screen opens.
- 3. Select the images to move. If all images are being moved, select Select All.
- 4. Select your search criteria (for example, Patient Name or Accession Number) using the drop-down list below the images.
- 5. Enter the same patient information and select the **Search** button (the magnifying glass).
- 6. When the Patient name appears, select the correct procedure for the images.
- 7. If ALL the images are being moved and you want to delete the incorrect procedure, select **Delete Empty**.
- 8. If you want to archive the images or save them to a stored device, select **Archive**. Verify that the correct outputs are selected.
- 9. Select **Split**, then select **OK** in the *Split Successful* message.

Images Acquired in the Wrong Patient

- 1. In the *Select Patient* screen, select the patient that has the images that need to be moved.
- 2. Select the **Split** button. The *Split Patient Records* screen opens.
- 3. Select the images to move. If all images are being moved, select Select All.
- 4. Select your search criteria (for example, Patient Name or Accession Number) using the drop-down list below the images.
- 5. Enter the search criteria and select the Search button (the magnifying glass).
- 6. When the Patient name appears, select the correct procedure.

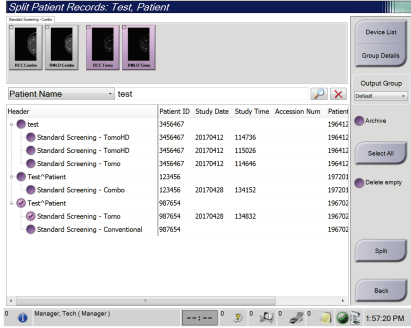


Figure 29: Select the Correct Procedure to Split Patient Records

- 7. If ALL images are being moved and you want to delete the incorrect patient from the Patient List, select **Delete Empty**.
- 8. If you want to archive the images or save them to a stored device, select **Archive**. Verify that the correct outputs are selected.
- 9. Select **Split**, then select **OK** in the *Split Successful* message.

5.3.6 Delete a Patient



Note

Note

Reclamation normally removes the requirement to delete patients manually. Refer to <u>*About Reclamation*</u> on page 137.

- 1. In the *Select Patient* screen, select one or more patients.
- 2. Select the **Delete** button.
- 3. When the confirmation prompt is shown, select **Yes**.



The Technologists do not have permissions to delete patients.

5.3.7 Filters for Patients

After selecting the **Filter** button in the *Select Patient* screen, the *Patient Filter* screen for the selected patient list opens.

Pati	ient Filter:	Schedulea	/						
_	Columns							_	
-	atient Name							^	Open
	atient ID								
-	ccession Numbe	er							
Ø Ra	ange	Today		-				Ξ	
Di	isposition	 Accepted Pended Rejected 							Refresh Worklis
Ro	ole	Me		¥					Query Worklist
🖉 So	ource	☑ Local☑ Worklist							
No	otice State	Unviewed						-	Save
lesult	s			_					Save As
ame			Date of Birth	Exam	Date/Time	/ Prior	Status	Pat	
F	⁰ Test [^] Patient		1/2/1972	Standard Scre		No	Scheduled	123	Delete Tab
									Order Tabs
•			m			Nu	mber of result	۲ 5: 1	Back
0	Manager, Te	ch (Manager)		:) 🦻 🕯 🎜	ų ° 🦨	8 ° 🥥		😧 11:55:02 A

Figure 30: Filter Tab in the Patient Filter Screen

Filter Tab

Use the **Filter** tab to change the filter options for the patient list. When you select or cancel an option, the change shows in the Results area of the screen.



Note

You must have Manager level access to save these new filters to the selected tab in the *Select Patient* screen. (Refer to <u>Other Functions of the Filter Tab</u> on page 65.)



Note

When you select a line in the results list then select the **Open** button, the *Procedure* screen for the selected patient opens.

Other Functions of the Filter Tab

The **Filter** tab allows users with access privileges to add, change, or delete tabs in the *Select Patient* screen. Refer to the following table.

Change the current patient	1.	Select a tab on the <i>Select Patient</i> screen.
filter parameters.	2.	Select the Filter button.
	3.	Select the filter options.
	4.	Select the Save button.
	5.	Make sure the name of the tab you selected is in
		the name box.
	6.	Select OK.
Create a new tab for the Select	1.	Select a tab on the <i>Select Patient</i> screen.
Patient screen.	2.	Select the Filter button.
	3.	Select the filter options for the tab.
	4.	Select the Save As button.
	5.	Enter a new name for the tab.
	6.	Select OK.
Delete a tab from the Select	1.	Select a tab on the <i>Select Patient</i> screen.
Patient screen.	2.	Select the Filter button.
	3.	Select the Delete button.
	4.	Select Yes in the confirmation dialog box.

Table 13: Filter Tab Options (Require Access Privileges)

Columns Tab

Use the **Columns** tab to add more search options (for example, Age, Gender, Notices) to the filtered list. The options show as columns in the results area. To add more columns to a filtered list, select the **Columns** tab then select the options.



You must have Manager level access to save these new columns to the patient filter.



Note

Note

When you select a line in the results list then select the **Open** button, the *Procedure* screen for the selected patient opens.

Order Tabs Button

Select the Order Tabs button to change the order of the patient list tabs.

5.3.8 Refresh the Worklist

Select the **Refresh Worklist** button to update the Scheduled patient list from the Modality Worklist Provider.

5.3.9 Query the Worklist

Select the **Query Worklist** button to search the Modality Worklist Provider for a patient or a list of patients.

There are two methods to enter the query information:

- **Keyboard**—Type the query information into one or more fields. The scheduled procedure is displayed, and the patient is added to the local database. All fields to query are configurable. The default fields are Patient name, Patient ID, Accession Number, Requested Procedure ID, and Scheduled Procedure Date.
- **Bar Code Scanner**—Scan the bar code for the configured query field. The scheduled procedure is displayed, and the patient is added to the local database. The field in which the bar code scanner scans is configurable. The default query field can be the Patient ID, Accession Number, or Requested Procedure ID.

5.3.10 Admin

Select the **Admin** button to access the *Admin* screen and the system administration functions. Refer to *System Administration Interface* on page 139 for more information.

5.3.11 Log Out

Select the Log Out button to exit the system and return to the *Startup* screen.

5.3.12 Advanced Workflow Manager

The Advanced Workflow Manager is a workflow engine that allows compatible Hologic systems to communicate and exchange images with each other. The Advanced Workflow Manager keeps track of all patients, procedures, and images acquired on all systems within an Advanced Workflow Manager cluster. In addition, the Advanced Workflow Manager synchronizes notices and provides images to all systems within the cluster.

Systems with Advanced Workflow Manager show the **Enterprise** tab on the <u>Select Patient</u> <u>Screen</u> on page 59. This tab lists the procedures acquired on all systems within an Advanced Workflow Manager cluster.



5.4 Procedure Screen

Figure 31: Procedure Screen

Item	Description
1. Image Status	The view icon shows the current selected view.
	Implant Present button—select when the patient has an implant.
	Accept button—select to accept the image.
	Reject button—select to reject the image.
	Pend button—select to save the image for future review.
2. Tabs	Select the Generator tab to adjust the exposure techniques for the procedure.
	Select the Tools tab to review the images.
	Select the Biopsy tab to create targets.
	Select the Contrast tab to perform I-View TM 2D Contrast procedures.
3. Buttons	Many functions are accessed from this screen by selecting a particular button:
	Add Procedure: Add a new patient.
	Add View: Add a new view.
	Edit View: Assign a different view to an image.
	Archive/Export: Send images to an output.
	Print: Print images.
	Retrieve: Query the configured devices with the current patient information.
	Paddle Shift: Bypass the default paddle position for the selected view.
	Close Patient: Exit the patient and procedure.
	Trash Can: Delete a view.
4. Thumbnails	Select a tab to show the thumbnail views or the thumbnail images for that procedure.

Table 14: The Procedure Screen

5.4.1 How to Use the Implant Present Button

The **Implant Present** button is above the **Accept** button on the *Procedure* screen. This button applies special implant processing to the implant and implant displaced views and changes the "Implant Present" DICOM tag in the image header. When this button is selected, a check mark appears on the button.



Select the **Implant Present** button for both implant and implant displaced views before you acquire the image.



Note

The **Implant Present** button is automatically selected if any procedure tabs contain an ID view.

5.4.2 How to Use the Paddle Shift Feature

Refer to *Paddle Shift* on page 118 for information about the Paddle Shift feature.

- 1. In the *Procedure* screen, select an unexposed thumbnail image view. The paddle moves to the default position for that view.
- 2. Select a paddle shift button to bypass the default paddle position for the selected view. The paddle moves to the new position.

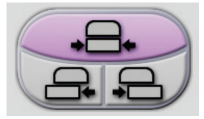


Figure 32: Paddle Shift Buttons

5.4.3 Procedure Information Dialog Box

To show procedure information, select the **Procedure Information** button located below the **Delete View** (trash can) button. The *Procedure Info* dialog box opens with the following information:

- Procedure Name
- Accession Number
- Procedure Status
- Procedure Start and End Date and Time
- Dose Information (each Breast and Cumulative)

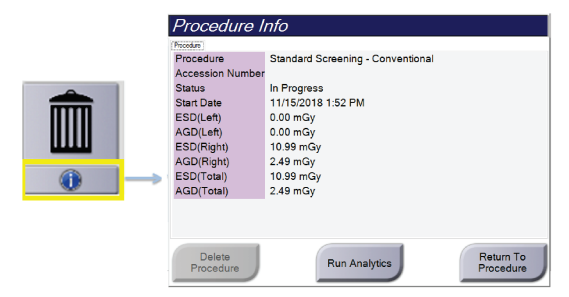


Figure 33: Procedure Info Dialog Box

Note

For procedures that contain no exposed views, select the **Delete Procedure** button to remove the selected procedure from the patient.

To run the image analytics manually, select the **Run Analytics** button.



The **Run Analytics** button only shows if you have a license for ImageChecker[®] CAD, Quantra[™], or Genius AI Detection software.

Select **Return to Procedure** to exit the dialog box.

About the Run Analytics Button

ImageChecker[®] CAD, Quantra[™] volumetric breast density assessment software, and Genius AI Detection software are available on the Selenia Dimensions and 3Dimensions systems.

When you are licensed for one of these features:

- Image analytics are run automatically when you select the **Close Procedure Complete** button to close a patient after acquiring new images. Image analytics are run automatically after editing a patient, editing a view, or adding a view when you select the **Close Procedure Complete** button to close a patient.
- Image analytics can be run manually by selecting the **Run Analytics** button in the *Procedure Info* dialog box. Run image analytics manually as needed on images that are received or imported from an external source.
- Image analytics run only on images that were acquired using the same software version or the immediate prior minor software version as the Acquisition Workstation.
- The *Procedure Info* dialog box includes tabs that show the success or failure status of the image analytics.
- Thumbnail images for the image analytics results are displayed in the thumbnail image section of the *Procedure* screen.

About the Genius AI Detection Tab

Genius AI Detection software is an available feature on the Selenia Dimensions and 3Dimensions systems. To show Genius AI Detection software results, select the **Procedure Information** button located below the **Delete View** (trash can) button. The *Procedure Info* dialog box opens. Select the **Genius AI Detection** tab to show the following information:

- Results
- Read Time Indicator
- Reading Priority
- Date and time of the Genius AI Detection software analysis
- Case Score
- CAD Complexity
- View name, time, and if the view is included for case processing

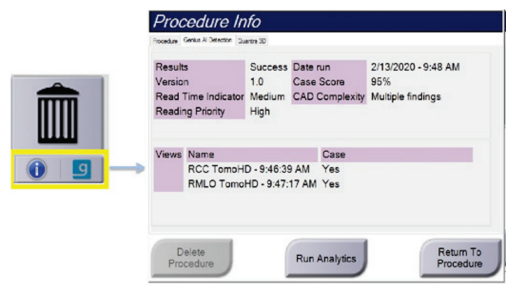


Figure 34: Genius AI Detection Tab

Select **Return to Procedure** to exit the dialog box.

5.4.4 Add a Procedure

1. To add another procedure, select the **Add Procedure** button on the *Procedure* screen to access the Add Procedure dialog box.

Procedure	
Conventional	
Standard Screening - Conventional	
Procedure Info	
Inherit Accession Number	
Accession Number	
	ок

Figure 35: Add Procedure Dialog Box

- 2. Use the drop-down menus to select the type of procedure to add.
- 3. The current Accession Number is automatically used. To use a different Accession Number, deselect the "Inherit Accession Number" check box and enter the desired number.
- 4. Select the **OK** button. A new tab is added with the thumbnail views for the procedure that was selected.

Available Procedures

The table that follows shows the major groups of procedures that are available on the system.



Note

Some procedures in the list show only when the system is licensed for that procedure.

Procedure Group	Description				
Conventional	Digital mammography (2D) imaging only				
Combo	Digital mammography (2D) and breast tomosynthesis imaging				
Tomo	Breast tomosynthesis imaging only				
TomoHD	Breast tomosynthesis imaging and a generated 2D image				
ComboHD	Digital mammography (2D) and breast tomosynthesis imaging and a generated 2D image				
Stereo Biopsy	Biopsy that uses stereotactic targeting				
Tomo Biopsy	Biopsy that uses tomosynthesis targeting				
Specimen	Specialized specimen imaging				
2D Contrast	Contrast enhanced digital imaging				
Contrast Stereo Biopsy	Biopsy that uses contrast stereotactic targeting				

5.4.5 Add (or Remove) a View

To Add a View

Note

1. Select the **Add View** button to access the *Add View* screen.



Depending on the license settings for your system, you may see different tabs.

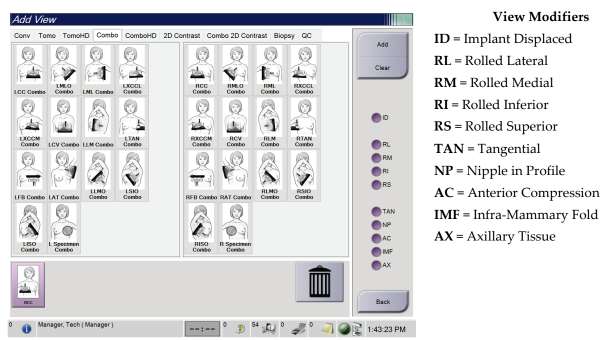


Figure 36: Add View Screen

- 2. Select the tab, then select the view. You can select a maximum of three View Modifiers from the right panel of the screen.
- 3. Select the **Add** button. A thumbnail image of each selected view is added in the bottom panel of the screen.

To Remove a View

- To remove a single view from the added list: select the view, then select the trash can icon.
- To remove all views from the added list: select the **Clear** button.

5.4.6 Edit a View

Use the *Edit View* screen to assign a different view to an image.

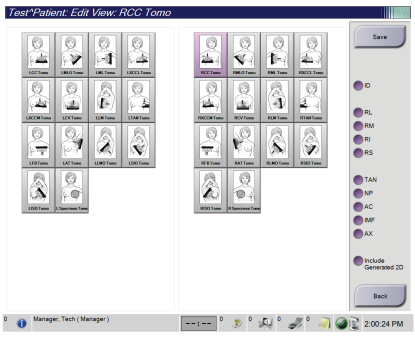


Figure 37: Edit View Screen

- 1. In the *Procedure* screen, select an exposed thumbnail image view.
- 2. Select the Edit View button.
- 3. Select the view from the screen. You can select a maximum of three View Modifiers. Refer to *Add (or Remove) a View* on page 74 for a description of the View Modifiers.
- 4. Select **Save**, then select **OK** to the *Update Successful* message.



Note

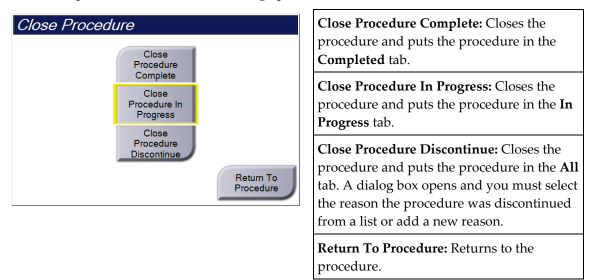
The *Edit View* screen for biopsy procedures shows different View Modifiers. Refer to the Affirm[®] breast biopsy guidance system *User Guide*.

5.4.7 Retrieve

Select the **Retrieve** button to query the configured devices with the current patient information. The **Retrieve** button performs the same function as the **Query Retrieve** button in the *Admin* screen.

5.4.8 Close a Patient

Select the **Close Patient** button. If images were acquired, the *Close Procedure* dialog box opens. Select one of the following options:



If there are images marked as Pend, a dialog box opens with the following options:

- Accept All and Close Complete
- Close Procedure In Progress
- Return to Procedure

If MPPS is activated, procedure status information is sent to the output devices when you select **Complete** or **Discontinue**. You can also select and hold the tab above the thumbnail images to resend a message about the procedure status during the procedure. A *Procedure Action* dialog box opens with buttons to resend a status or to return to the procedure.



Important

When you are licensed for 3DQuorum software, follow the standard procedure to close a patient. If there are SmartSlices that have not completed processing, a dialog box opens with the following message:



Wait for processing to complete before you open the next patient.



Note

When you are licensed for Genius AI Detection software, ImageChecker CAD, or Quantra software, image analytics are run automatically when you select the **Close Procedure Complete** button after acquiring new images. The results are automatically sent to the Output Devices. When you reopen the patient, thumbnail images for the results are displayed in the *Procedure* screen, and status tabs are available in the *Procedure Info* dialog box.

5.5 How to Access the Image Review Features

Select the **Tools** tab on the *Procedure* screen to access the image review features. Refer to *Image Review Tools Tab* on page 96 for information.

5.6 Output Groups

The accepted images are sent automatically to the output devices in the selected Output Group. The system configuration controls if the images are sent after a patient is closed or after the image is accepted.



Note

Tomosynthesis images are not sent to a print device in the selected Output Group. You can print selected tomosynthesis images from the *Print* screen.

5.6.1 Select an Output Group

Output Groups			
SV_02 •			

Figure 38: Output Groups Field

Select an output device group like PACS, Diagnostic Workstations, CAD devices, and printers from the Output Groups drop-down list in the *Procedure* screen.



Note

Images are not sent if an Output Group is not selected.

5.6.2 Add or Edit an Output Group



Note

The configuration of Output Groups occurs during installation, but you can edit existing groups or add new groups.

To add a new Output Group:

- 1. Access the *Admin* screen.
- 2. Select the Manage Output Groups button.
- 3. Select the New button, enter the information, then select the output device(s).
- 4. Select **Add**, then select **OK** in the *Update Successful* message.
- 5. You can select any group to set as the default.

To edit an Output Group:

- 1. Access the *Admin* screen.
- 2. Select the Manage Output Groups button.
- 3. Select the **Edit** button, then make the changes.
- 4. Select **Save**, then select **OK** in the *Update Successful* message.

5.6.3 SmartSlices Output Groups



Note

The configuration of Output Groups occurs during installation, but you can edit existing groups or add new groups.

During the installation of 3DQuorum software, new default Output Groups are created in the system to incorporate output options for SmartSlice images. The new output groups allow you to control whether SmartSlices, tomosynthesis reconstructed slices, or both are sent to the preferred devices automatically. See the following table for a description of the output groups.

Output Group Name	Explanation of the Configuration
SmrtSlces,DBT,2D	Sends all configured data types.
SmartSlices,2D	Sends SmartSlices and 2D data types (synthesized 2D or FFDM). No 1-mm reconstructed slices are sent.
DBT,2D	Sends 1-mm reconstructed slices and 2D data types (synthesized 2D or FFDM). No SmartSlices are sent.
Custom	Provides on-demand selection of outputs and data types.

Table 16: Output Groups for 3DQuorum Software

To create the three new output group types, each ImageStorage output device is configured to have three Installed Device entries. This allows maximum flexibility for experimenting with whether SmartSlices are always used in diagnostic reading, are always archived, or are only sent on-demand. See the following table for a description of the installed output device settings.

Output Name	Explanation of the Configuration
Device name	Sends all desired data types except SmartSlices and 1-mm reconstructed slices.
Device name - 3D 1mm Slices	Sends only 1-mm reconstructed slices.
Device name - SmartSlices	Sends only SmartSlices.

Table 17: Output Device Settings for 3DQuorum Software

ImageStorage[PACS]
ImageStorage[PACS-3D 1mm Slices]
ImageStorage[PACS-SmartSlices]

Figure 39: Example of the Output Device Settings

As decisions are made regarding which data to send to workstations for diagnosis, which data to archive, and which data to only send on-demand, the default configurations can be further customized. Changes can be made during Clinical Applications Training or by contacting Hologic Technical Support.

5.6.4 Custom Output

The Custom Output Group option lets you make an output group from the *Procedure* screen. The custom output group that you make stays as the Custom option until another custom output group is made.

To Make a Custom Output Group from the *Procedure* screen:

- 1. In the Procedure screen, select Custom from the Output Groups drop-down list.
- 2. In the *Output Group* dialog box, select from the list of available devices, then select **OK**.

Output Group	
Please choose the outputs for this custom out	put group.
ImageStorage[NIKOS_SV]	
ImageStorage[MEVIS_02]	
	ок
	Cancel

Figure 40: An Example Custom Output Group

5.7 On-Demand Outputs

The On-Demand Outputs are **Archive/Export** or **Print**. You can manually Archive, Export, or Print the currently opened patient until the procedure is closed.

When you select an **On-Demand Output** button, you have the option to send images from the patient that is open to any of the configured output devices.

5.7.1 Archive

- 1. Select the Archive/Export button.
- 2. Select the procedure or views in the On Demand Archive screen:
 - Select All button selects all items that show in this screen.
 - **Clear** button deselects items that are selected in this screen.
 - Priors button shows previous procedures and views for this patient.
 - Rejected button shows rejected views for this patient.
- 3. Select a storage device:
 - Select the **Device List** button and make your selection from the options in the *Storage Device* drop-down menu.

-OR-

- Select an output group from the *Output Group* drop-down list.
- 4. Select the Archive button to send the selected images to the selected archive.



Note

Use the Manage Queue utility in the taskbar to review the archive status.

5.7.2 Export

- 1. Select the Archive/Export button (on the right side of the *Procedure* screen).
- 2. Select the images to export, then select the **Export** button.

On Demand Archive						
Header	Patient ID	Study Date	Study Time	Accession Num	Patient	
	123456				197201	Device List
🛛 🕢 Standard Screening - Conventional	123456	20170411	141951		197201	Group Details
						Group Details
						Output Group
						MEVIS_02 -
						MLV13_02
						Select All
						Clear
						Display Priors
						Rejected
					•	
Standard Screening - Conventional						Export
						Archive
RCC						Back
						Dack
1 👔 Manager, Tech (Manager)	-	-: 0	2 3 🛴	0 _ 0		2:40:21 PM

Figure 41: Select the Images for Export

3. In the *Export* dialog box, select the target from the drop-down list of media devices.

Export	
Target	Removable Disk (E:)
Progress	
Anonymize	Start
Eject USB device after write	Close

Figure 42: Export Dialog Box

- To anonymize patient data, select **Anonymize**.
- To automatically eject the removable media storage device when the export is complete, select **Eject USB device after write**.
- To select a folder on your local systems for storage of your selections and to select the Export types for the image, select **Advanced**.
- 4. Select the **Start** button to send the selected images to the selected device.

5.7.3 Print

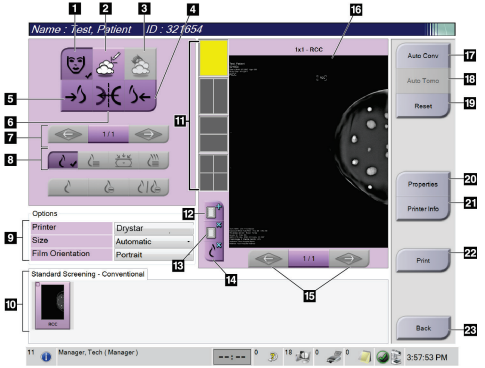


Figure 43: Print Screen

Figure Legend

- 1. Show or hide patient data.
- 2. Show or hide markings and annotations.
- 3. Show or hide targets in images from a biopsy.
- 4. Print the image from a dorsal perspective.
- 5. Print the image from a ventral perspective.
- 6. Reverse (mirror) the image.
- Go to previous or next tomosynthesis slice or projection (Tomosynthesis option).
- Select Conventional, Projection, or Reconstruction views (Tomosynthesis option). The C-View button shows if the system is licensed for C-View imaging.
- 9. Select the printer options.
- 10. View Thumbnail images.
- 11. Select the film format (number of tiles).
- 12. Create a new film.

- 13. Delete a film.
- 14. Delete an image from a film.
- 15. Step through the film pages.
- 16. Print preview area.
- 17. Print Conventional (and C-View images, if system is licensed) with the default setup.
- 18. Print tomosynthesis images (slices or projections) Tagged for Print (Tomosynthesis option).
- 19. Return the Print screen to default settings.
- 20. Open the Properties screen.
- 21. Show the printer IP address, AE Title, Port, and capability for True Size print.
- 22. Start the print process.
- 23. Return to the Procedure screen.
- 1. From the *Procedure* screen, select the **Print** button to show the *Print* screen.
- 2. Select the film format from the film format area of the screen.
- 3. Select a thumbnail image.
- 4. Select the image mode (Conventional, Projection, or Reconstruction). The **C-View** button shows if the system is licensed for C-View imaging.

- 5. Select the print preview area (item 16) in the *Print* screen. The image that shows in this area is the image that prints on the film.
- 6. To put other images on the same multiformat film, repeat steps 3 through 5.
- 7. To print a different film format of the same images, select the **New Film** button (item 12), then complete steps 2 through 6.
- 8. Use the buttons in the top left area of the *Print* screen (item 1 through item 6) to hide or show patient data, markings, and annotations, and to change the orientation of the image.
- 9. Select the **Print** button to print your films.

Chapter 6 Imaging

6.1 Image Display Screen

After you make an exposure, the acquired image opens on the Image Display monitor. Patient and procedure information can be displayed on the screen. To turn the information on or off, go to the **Tools** tab in the *Procedure* screen and select the **Patient Information** button.

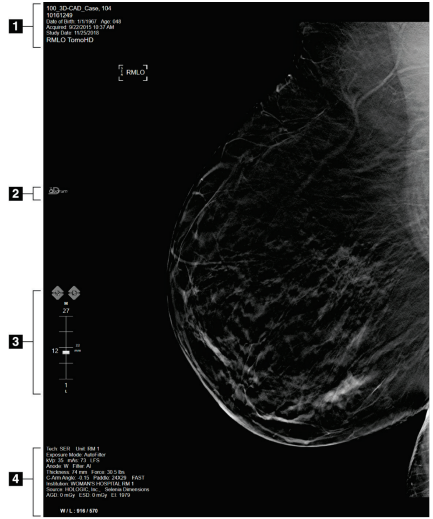


Figure 44: Image Display Screen (SmartSlices option shown)

Figure Legend

- 1. Patient information and exam date
- 2. 3DQuorum software logo
- 3. Slice Indicator
- 4. Exposure information

6.2 How to Set the Exposure Parameters

6.2.1 Select the Image Acquisition Mode (Tomosynthesis Option)

- Standard For routine Tomosynthesis screening procedures
- Enhanced For diagnostic Tomosynthesis views



Warning:

Enhanced acquisition mode with combo mode imaging (DM + BT) can produce a radiation dose that is higher than the MQSA screening limit of 3.0 mGy and therefore should only be used in diagnostic evaluation.

6.2.2 Select the Exposure Mode

Use the Automatic Exposure Control modes (AEC) to let the system control the exposure techniques. AEC modes are available from 20-49 kV.

- Manual The user selects the kV, mAs, Focal Spot, and Filter.
- AEC: Auto-Time The user selects the kV, Focal Spot, and Filter. The system selects the mAs.
- AEC: Auto-kV The user selects the Focal Spot. The system selects the kV, mAs, and Filter (Rhodium).
- AEC: Auto-Filter The user selects the Focal Spot. The system selects the kV, mAs, and Filter.

6.2.3 How to Use the AEC Sensor

The AEC Sensor has seven manual positions and an automatic position. The manual positions start at the chest wall edge (position 1) and reach toward the nipple edge (position 7). The automatic position selects two regions within an area that extends from the chest wall to the nipple.

Use the plus (+) and minus (-) keys on the Compression Device or in the AEC Sensor area of the screen to change the sensor position. You can select Auto AEC to allow the system to calculate the best exposure for the breast.

6.3 How to Acquire an Image

Refer to <u>Clinical Procedures</u> on page 127 for information about clinical procedures.

- 1. Select a view from the thumbnail views at the bottom of the *Procedure* screen.
- Press and hold the x-ray button and/or the x-ray footswitch for the full exposure.
 During the exposure:
 - The System Status bar shows the radiation symbol and a yellow background (see the following figure).

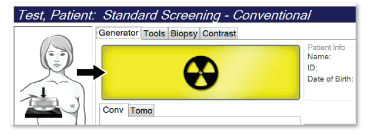


Figure 45: Exposure In Progress

• An audible tone continues to sound during the exposure

The audible tone is a continuous sequence of tones. The tone sounds during the entire combo acquisition from the initiation of the exposure to the end of the conventional view. There is no interruption of the audible tone between breast tomosynthesis and conventional digital mammography exposures. Do not release the exposure switch during the audible tone.

3. When the tone stops and the System Status bar shows **Standby** (see the following figure), release the **x-ray** button and/or the **x-ray footswitch**.

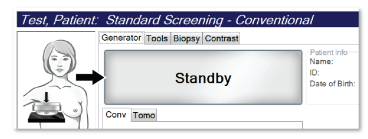


Figure 46: Exposure Complete

4. When the x-ray is finished, the image appears on the Image Display monitor. The *Procedure* screen automatically changes to the **Tools** tab.

Select one of the following options to complete the acquisition:

- Accept the image. The image transmits to output devices with all attributes.
- **Reject** the image. When the dialog box opens, select the reason for rejecting the image. The *Image Display* screen closes. You can repeat the rejected view or select another view. If you select **Reject**, an "X" appears on the thumbnail image.
- **Pend** the image. The image saves for future review. If you select **Pend**, a question mark "?" appears on the thumbnail image.



Figure 47: Marked Images in a Procedure

5. Repeat steps 1 through 4 for each view.



A Manager user can configure the system to Auto-Accept or Auto-Pend new images.

6.3.1 Conventional Imaging Sequence of Events

- 1. Review the image after the exposure and add a comment, if necessary.
- 2. Accept, Reject, or Pend the image. A thumbnail image replaces the thumbnail view in the Case Study area of the screen.



Note

Note

Note

A Manager user can configure the system to Auto-Accept or Auto-Pend new images.

6.3.2 Tomosynthesis Imaging Sequence of Events

- 1. Wait for the image reconstruction to complete.
- 2. Review the projection images for motion.
- 3. Accept, Reject, or Pend the images. A thumbnail image replaces the thumbnail view in the Case Study area of the screen.



A Manager user can configure the system to Auto-Accept or Auto-Pend new images.

6.3.3 How to Accept a Rejected Image

If a rejected image is better than the new image, you can retrieve and use the old image. Select the thumbnail image on the *Procedure* screen to review the image, then **Accept** the image.

6.3.4 How to Accept or Reject a Pended Image

To accept or reject a Pended image, select the Pended thumbnail image, then select the **Accept** button or the **Reject** button.

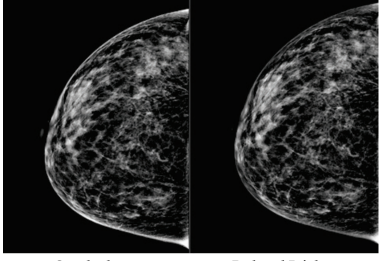


Note

A Manager user can configure the system to Auto-Accept or Auto-Pend new images.

6.3.5 Skin Line Brightness

An optional image processing setting is available for 2D imaging to reduce the brightness and overshadow of the skin on screening views. To enable this option, please contact your Hologic representative.



StandardReduced BrightnessFigure 48: Comparison of Standard Brightness and Reduced Brightness

6.4 How to Acquire an Image with an Implant

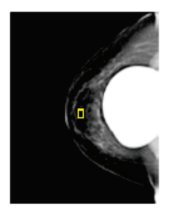
\land

Note

ALWAYS select the **Implant Present** button for all implant views. This button processes the implant images correctly.

6.4.1 Implant Away from the Nipple

- 1. Select a view from the thumbnail views at the bottom of the *Procedure* screen.
- 2. For the AEC mode, select **Auto Filter**.
- 3. Use the + and buttons to move the AEC sensor to the tissue behind the nipple.



- 4. Acquire the exposure.
- 5. Accept, Reject, or Pend the images.



Note

A Manager user can configure the system to Auto-Accept or Auto-Pend new images.

6. Repeat steps 1 through 5 for each view.

6.4.2 Implant Near the Nipple

- 1. Select a view from the thumbnail views at the bottom of the *Procedure* screen.
- 2. For the AEC mode, select Manual.
- Use the + and buttons to select the kVp and mAs values. Refer to the tables in <u>Values</u> <u>Tables for Implant Near the Nipple</u> on page 91 for the recommended values based on the type of view and the compression thickness.





Note

The CEDM view consists of two separate exposures. Only the techniques for the first, lower energy exposure can be set manually. The system automatically sets the techniques for the second exposure based on the techniques from the first exposure.



Note

For Combo Contrast views, select values for the tomosynthesis exposure from Table 17 and values for the contrast enhanced (CEDM) exposure from Table 18.

For more information on acquiring contrast enhanced images, refer to <u>*I-View Contrast*</u> <u>*Enhanced 2D Images*</u> on page 100.

- 4. Acquire the exposure.
- 5. Accept, Reject, or Pend the images.



Note

A Manager user can configure the system to Auto-Accept or Auto-Pend new images.

6. Repeat steps 1 through 5 for each view.

Values Tables for Implant Near the Nipple

Table 18: Conventional 2D Imaging						
Compression Thickness	kV	mAs	Filter			
< 4 cm	28	100	Rh			
4 - < 6 cm	28	120	Rh			
6 - < 8 cm	28	140	Rh			
8 - 10 cm	28	160	Rh			
> 10 cm	28	180	Rh			

Table 19: Tomosynthesis Imaging

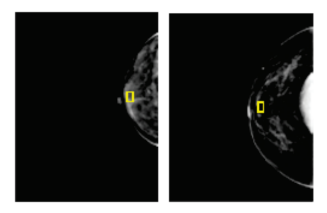
Compression Thickness	kV	mAs	Filter
< 4 cm	29	60	Al
4 - < 6 cm	31	70	Al
6 - < 8 cm	33	90	Al
8 - 10 cm	35	100	Al
> 10 cm	38	100	Al

Compression Thickness	kV	mAs	Filter
< 3.5 cm	26	30	Rh
3.5 - < 4 cm	27	40	Rh
4 - < 5 cm	28	40	Rh
5 cm - < 5.5 cm	29	60	Ag
5.5 cm - < 6 cm	30	60	Ag
6 - < 7.5 cm	31	80	Ag
7.5 - < 8.5 cm	32	120	Ag
8.5 - 12 cm	33	120	Ag
> 12 cm	33	160	Ag

Table 20: Contrast Enhanced (I-View) Imaging

6.4.3 Implant Displaced Views

- 1. Select a view from the thumbnail views at the bottom of the *Procedure* screen.
- 2. For the AEC mode, select **Auto Filter**.
- 3. Use the + and buttons to move the AEC sensor to the tissue behind the nipple.



- 4. Acquire the exposure.
- 5. Accept, Reject, or Pend the images.



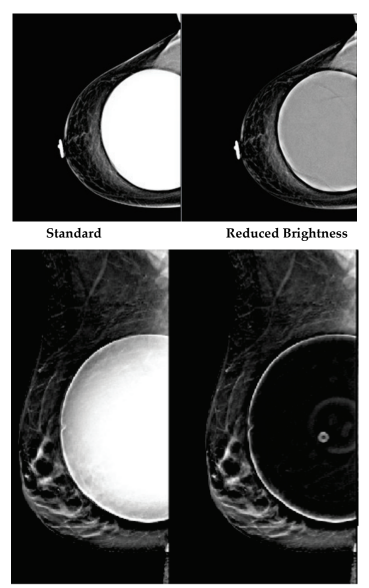
Note

A Manager user can configure the system to Auto-Accept or Auto-Pend new images.

6. Repeat steps 1 through 5 for each view.

6.4.4 Implant Brightness

An optional image processing setting is available for both 2D and 3D imaging with implants. The implant brightness can be reduced to improve visibility of the implant/breast boundary region. To enable this option, please contact your Hologic representative.



Standard Reduced Brightness *Figure* 49: *Comparison of Standard Brightness and Reduced Brightness*

6.5 How to Correct and Reprocess Implant Images

You must correct the image if you acquire an implant or an implant displaced view without the **Implant Present** button activated.

6.5.1 If the Image Is Not Accepted

Select the **Implant Present** button on the *Procedure* screen to indicate an implant exists. A check mark appears on the button and the image reprocesses.



6.5.2 If the Image Is Accepted

- 1. Select the image.
- 2. Select the **Implant Present** button on the *Procedure* screen to correct the image. A check mark appears on the button and the image reprocesses.
- 3. Select the **Accept** button to accept the changes.



Note

The corrected image is sent automatically to the selected output devices if the system is set to send images when the **Accept** button is selected.

6.6 How to Review the Images

Review of the images involves use of the thumbnails, image review tools, and display modes.

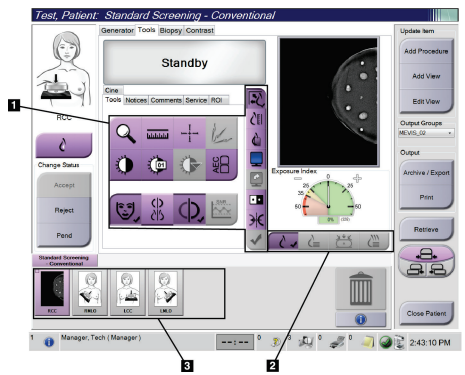


Figure 50: Tools Tab (Tomosynthesis option shown)

- 1. Image Review Tools refer to *Image Review Tools Tab* on page 96.
- 2. Image Display Modes refer to *Display Modes* on page 98.
- 3. Thumbnail Views and Thumbnail Images. Select any thumbnail image to display that image on the Image Display monitor.

6.6.1 Image Review Tools Tab

The **Tools** tab in the *Procedure* screen provides the image review tools. A check mark appears on an active tool.

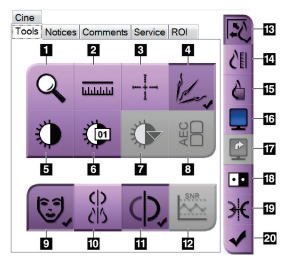


Figure 51: Image Review Tools

- 1. The **Zoom** tool magnifies a section of the image.
- 2. The **Ruler** tool measures the distance between two points.
- 3. The **Crosshair** tool displays a crosshair on the *Image Display* screen.
- 4. The **Demetalizer** tool for processing tomosynthesis images that contain metal objects.
- 5. The **Window/Level** tool changes the brightness and contrast.
- 6. The **Window/Level Fine Adjustment** tool allows the entry of specific window and level values.
- The LUT Selection tool scrolls through available Window/Level settings for a displayed image with LUTs attached.
- 8. The **AEC** button displays the AEC Sensor areas used for the exposure calculation. The sensor areas display on the *Image Display* screen.
- 9. The Patient Information button activates the patient information display.
- 10. The **Auto-Hanging** button automatically hangs the currently selected study in a 4-up configuration.
- 11. The **Auto-Pairing** button turns off auto-pairing for the currently selected image in a multi-up configuration.
- 12. The **SNR/CNR** button calculates the signal-to-noise ratio and contrast-to-noise ratio on the ACR Phantom.
- 13. The **Fit-to-Viewport** button fits the image within the image tile.
- 14. The **True Size** button displays the image in the actual size of the breast.
- 15. The View Actual Pixels button displays the image in full resolution.
- 16. The **Multi-Up Display** button selects the number of tiles to display.
- 17. The Image Tile Advance button sets the active Multi-Up tile.
- 18. The Invert Image tool changes blacks to whites and whites to black.
- 19. The Mirror button reverses (mirrors) the image.
- 20. The **Tag for Print** button tags the projection or reconstruction images of a tomosynthesis image to print later.

6.6.2 Notices Tab

The tools on the **Notices** tab allow you to mark and annotate images and send notices about the displayed image or procedure. The Notice Output Groups drop-down menu at the bottom of the tab allows you to select the destination for the Notices.

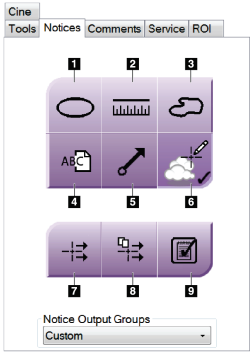


Figure 52: Tools on the Notices Tab

- 1. The **Oval** tool draws an oval marking on the image.
- 2. The **Ruler** tool measures the distance between two points.
- 3. The Freehand tool draws a freehand marking on the image.
- 4. The Text tool adds a text note to the image without a marking.
- 5. The **Arrow** tool draws an arrow marking on the image.
- 6. The Markings tool shows or hides markings and annotations on an image.
- 7. The **Send Notice** tool sends a notice for the current image to the selected destination.
- 8. The **Send All Notices** tool sends all notices for all images in the opened procedure to the selected destination.
- 9. The Viewed tool changes the Patient Notices status to viewed.

6.6.3 Other Image Review Tools

Other Tabs

- **Comments**: Add comments to an image.
- Service: Mark an image for service use.
- **ROI**: Draw a Region of Interest on the Image Display monitor.
- Cine: Show a series of images as a movie (Tomosynthesis option).

Exposure Index

The Exposure Index is an image quality guide. When the Exposure Index indicates the red or yellow area, review the selected image for noise and decide whether to reacquire the image.



Figure 53: Exposure Index

Display Modes

Use the buttons in the Display Modes area of the screen to select the type of view to show on the Image Display monitor. You can change between conventional, generated 2D, projections, reconstructions, and SmartSlices to show the combination images.

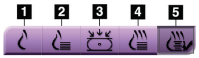
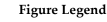


Figure 54: Display Modes

- 1. Conventional button shows conventional 2D images.
- 2. **Generated 2D** button shows a 2D image generated from the acquired tomographic image.
- 3. **Projections** button shows the projection images from the 15° sweep.
- 4. Reconstructions button shows 1-mm reconstructed slices.
- 5. SmartSlices button shows 6-mm slices from the 3DQuorum software.

6.6.4 Projection Indicator

The Projection Indicator on the Image Display monitor shows only on tomosynthesis procedures. Select the Projections display mode to show the indicator.



- 1. Use the scroll wheel on the mouse to move through the projections.
- 2. Number of the projection for the image
- 3. Angle of the tube for the projection

1/15 -7.49°

Figure 55: Projection Indicator

6.6.5 Slice Indicator

1

The Slice Indicator on the Image Display monitor shows only on tomographic reconstructions, including SmartSlices.

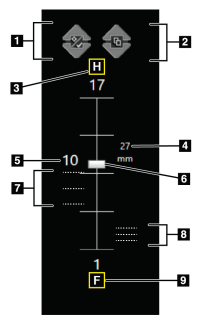


Figure 56: Slice Indicator (*SmartSlices option shown*)

Figure Legend

- 1. **Up** and **Down** arrows let you change between slices that contain a lesion target and slices that are tagged for printing.
- 2. **Up** and **Down** arrows let you change between slices that contain notices.
- 3. "H" (anatomical reference to head direction)
- 4. Height within the breast volume (for SmartSlices)
- 5. Number of the tomosynthesis reconstructed slice or SmartSlice image
- 6. Scroll bar moves through the slices of the reconstruction.
- 7. Slices that contain targets or are tagged for printing.
- 8. Slices that contain notices.
- 9. "F" (anatomical reference to foot direction)

6.7 How to Send the Images to the Output Devices

You can send the images to output devices or use the Archive/Export function to copy images to temporary storage media. Refer to <u>Output Groups</u> on page 77 and <u>On-Demand</u> <u>Outputs</u> on page 80 for instructions.

6.8 I-View Contrast Enhanced 2D Images



Warning:

Patients can have adverse reactions to contrast agents. Refer to the Instructions for Use for the contrast agent for complete information.



Warning:

Contrast enhanced mammography uses contrast agents that are injected intravenously. Allergic reactions can occur.



Note

Hologic configures some systems to meet specific requirements. Your system configuration may not have all the options and accessories included in this manual.



Note

Do not use SmartCurve system paddles for I-View procedures.

- Test, Patient: Standard 2D Contrast Generator Tools Biopsy Contrast Add Procedure READY Add View Edit View Output Groups RCC CEDM Ò 0 Archive / Export Image Status Print Accept Retrieve Reject Pend Standard 2D Contrast Tube Loading Close Patient Manager, Tech (Manager) ----- h 0 0 0 0 0 2 2 12:50:42 PM
- 1. Select the **Contrast** tab to access the I-View[®] 2D Contrast (CEDM) function.

Figure 57: I-View 2D Contrast Screen

2. Set the contrast settings. For more information, refer to <u>*How to Configure the Contrast*</u> <u>*Settings*</u> on page 105.



Contrast information is included in the DICOM header of I-View images.

To begin the contrast dose and start the timer, select the Start (arrow) button.
 The timer begins in the Waiting Period, which appears with a yellow background.



Note You can adjust the length of time for th

You can adjust the length of time for the **Waiting Period** and **Optimal Imaging Period** phases of the timer for each procedure. Use the left and right arrows.

Note

Note

You can adjust the default length of time for the **Waiting Period** and **Optimal Imaging Period** phases of the timer. Refer to <u>Set the Contrast Defaults</u> on page 150.



Figure 58: I-View 2D Contrast Screen, Waiting Period



Note

The timer function does not allow you to stop the timer, only to start and reset it. The timer stops only when you exit the patient procedure.

After the **Waiting Period**, the timer begins the **Optimal Imaging Period**, which appears with a green background.

4. Acquire the images during the **Optimal Imaging Period**.

After you acquire an image, a marker will appear below the timer.

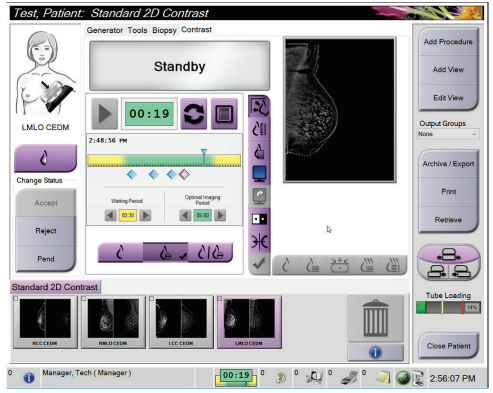


Figure 59: I-View 2D Contrast Screen, Optimal Imaging Period

- 5. Select the low energy and subtraction buttons to display both low energy and dualenergy subtracted images.
 - Low Energy: Conventional low-energy image
 - **Subtraction**: Contrasted, subtracted image
 - Show Low Energy and Subtraction images in 2-up display: Shows both the low and sub screen images side-by-side in a split screen.

Caution:

Performing an I-View 2D Contrast procedure increases the recovery time the machine needs before more images are acquired. When the System Status icon appears red

(^{min}), a recommended wait time is shown. This wait time lets the x-ray tube temperature decrease to avoid damage that can void the x-ray tube warranty. If the System Status icon turns red during a procedure, follow the guidance provided in section <u>Tube Loading Indicator</u> on page 103.

The x-ray tube heat load is tracked by the tube loading indicator. Be aware of the tube loading status when acquiring Contrast Enhanced images. Refer to <u>*Tube Loading Indicator*</u> on page 103 for more information.

6.8.1 Tube Loading Indicator

The **Generator** tab of the *Procedure* screen includes a Tube Loading indicator. This indicator shows the current heat load of the X-ray tube.

The Tube Loading indicator displays one of the following three statuses:

• A green status means that the heat loading of the x-ray tube is below what is considered steady state heat loading. During the green status, the system can accommodate 4 patients per hour of I-View contrast enhanced procedures.

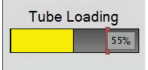




Note

Steady state heat loading of the Dimensions x-ray tube is defined as operating at 55% of x-ray tube heat capacity continuously. Steady state represents continuous Combo imaging at 8 patients per hour.

• The yellow status means that the x-ray tube heat loading is above what is considered steady state operation (see note above). Under yellow status, the system still has plenty of heat capacity to accommodate a full I-View contrast enhanced procedure and sustain such acquisitions for up to 4 patients per hour.





Note

The number displayed at the corner of the Tube Loading indicator shows the estimated tube heat loading with 55% representing steady state. It is expected that the Tube Loading indicator will turn yellow during an I-View contrast enhanced procedure and will remain yellow for the duration of the procedure. It is also possible, but rare, that the Tube Loading indicator appears yellow when opening a patient to start an I-View contrast enhanced procedure. In either case, the I-View procedure can continue uninterrupted. While the Tube Loading indicator remains yellow, the tube heat loading is considered safe.



Note

During a yellow Tube Loading status, the System Status icon may flash red with a number representing the number of minutes it will take to drop below the steady state heat loading. There is no need to wait before acquiring images under a yellow Tube Loading indicator despite the flashing System Status icon. The System Status icon is only informational and does not require action under a yellow Tube loading indicator.

The red status means that the x-ray tube heat loading just passed a threshold, which is considered potentially risky when attempting to acquire additional images. Under red status, the system still has some heat capacity left (threshold is set to 75% of tube heat loading capacity). However, waiting is recommended for the suggested number of minutes indicated under the Tube Loading indicator prior to acquiring an additional image. Typically, this time is very short (approximately one minute). As soon as the Tube Loading indicator returns to yellow status, an additional image can be acquired.





Note

Under certain circumstances (for example, large compressed breast thickness or more than four contrast views), it is normal to see the Tube Loading indicator turn red after the last exposure of an I-View contrast enhanced procedure. The system will quickly come back to equilibrium, and it will return to yellow or even green status within minutes. Even if the Tube Loading indicator turns red occasionally, the system can still maintain a rate of 4 patients per hour undergoing I-View contrast enhanced exams. There is still space for additional heat loading to the x-ray tube.



Caution

Excessive heat buildup can damage the X-ray tube.

6.8.2 How to Configure the Contrast Settings

1. From the **Contrast** tab, select the **Configure Contrast** button to edit the contrast settings. The *Contrast Information* dialog box opens.

Contrast Entry Route	Intra-arteria	l route
Contrast Agent	Diatrizoate	•
Contrast Agent Concentration	240	► mg/ml
Total Bolus Volume	0.0	ml
Patient Weight	lb 0.0	= 0.0
Contrast Concentration Per Body Weight	0.0	ml/kg
Amount of Contrast Agent	0.0	📄 ml

Figure 60: I-View 2D Contrast Settings

- 2. Use the drop-down lists to select the appropriate settings.
- 3. Enter data for the 'Contrast Concentration Per Body Weight' field or the 'Amount of Contrast Agent' field. The other field automatically populates with the correct information.
- 4. Select Save & Close.

6.9 Specimen Images

The system can be used for imaging biopsy specimens.

To acquire specimen images:

- 1. Select a patient from the worklist, or manually add a new patient.
- 2. Add or select the required specimen imaging procedure.

Patient, Test: Right Specimen - Conventional						
	Generator Tools	Generator Tools Biopsy Contrast				
	READY			Patient Info Name: Patient, Test ID: 987321 Date of Birth: 2/1/1974		Add Procedure Add View
	Conv Tomo	AEC Mode	Focal Spot	Comp Release	Manual	Edit View
R Specimen				Comp Mode	Dual	Output Groups
	[STANDARD]	Auto Time	LFS	Force	0.0 lbs	Store ~
3				Thickness	18.4 cm	
	kVp	mAs	Filter	Stereo Mode	Manual	Output
Image Status					_	
	26	[Auto]	Rh .	Collimation	24x29	Archive / Export
Accept	- +		- +	Paddle	None	Print
Reject	AEC Sensor	AEC Comp.	Grid	Mode	Normal	
	Auto	4	In	Position	Center	
Pend	- +	- +	- +	Мад	None	Retrieve
Right Specimen - Conventional						* *
- Conventional					Î	Close Patient
0 🚺 Manager, Teo	ch (Manager)		: 0	82 40 0	🦪 ° 🥥 🥥	3:14:01 PM

Figure 61: Procedure Screen for Specimens

- 3. Make sure that the C-arm is set to 0 degrees.
- 4. Prepare the specimen and place it on the image detector.

Patient, Test:	· Left Specimen - Conventional	
	Generator Tools Biopsy Contrast	Update Item
	Standby	Add Procedure Add View
1000	Cine Tools Notices Comments Service ROI	Edit View
L Specimen		Output Groups None ~
Change Status		Output
Accept	Exposure Index	Archive / Export
Reject		Print
Pend		Retrieve
Left Specimen - Conventional		
L Specimen		Close Patient
0 🚺 Manager, Te	ech (Manager) 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	3:09:53 PM

5. Acquire the images. For more information on acquiring images, refer to <u>*How to*</u> <u>*Acquire an Image*</u> on page 87.

Figure 62: Specimen Imaging Screen

6. Review the images as necessary.

Chapter 7 Accessories

The system can perform screening or diagnostic applications with specified accessories. This chapter describes how to use all possible system accessories.



Note

Hologic configures some systems to meet specific requirements. Your system configuration may not have all the options and accessories included in this manual.

7.1 How to Install Accessories on the C-Arm

The Retractable Face Shield, Magnification Stand, and Localization Crosshairs are installed in slots on the C-arm. The slots have labels with icons to indicate the accessory that attaches in the slot. Each accessory has two lines. Align the accessory with the related line on the C-arm. When the hook on the accessory is at the correct depth, the second, thinner line aligns with the line on the C-arm. The next sections contain installation instructions for each accessory.

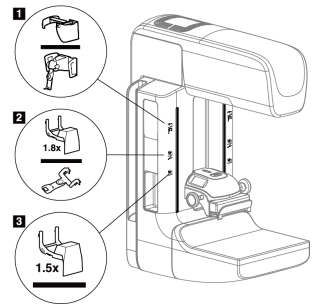


Figure 63: C-arm Accessories

- 1. Slot for Retractable Face Shield (Tomosynthesis option) or the Affirm® breast biopsy guidance system (biopsy option)
- 2. Slot for 1.8x Mag Stand and Localization Crosshairs
- 3. Slot for 1.5x Mag Stand

7.2 Patient Face Shields

The Face Shield keeps the head and face of the patient away from the x-ray field during the examination. Inspect the shield each day before use.



Warning:

The Face Shield must be attached for all exposures except magnification case studies.



Warning: The Face Shield does not protect the patient from radiation.

7.2.1 How to Install and Remove the Retractable Face Shield

To install the Retractable Face Shield:

- 1. Completely extend the Face Shield to the outer position.
- 2. Align the hooks of the Face Shield with the mounting slots on the C-arm, indicated by a face shield icon.
- 3. Put the hooks on both sides of the Face Shield into the mounting slots on the C-arm. The Unlock Lever (see item 1 in the following figure) is in the Up position.
- 4. Push the Face Shield to the down and locked position. The Unlock Lever is in the Down position when the Face Shield locks.

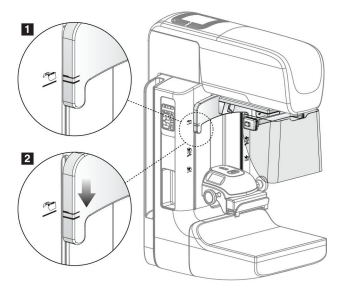


Figure 64: Align the Retractable Face Shield on the C-arm

To remove the Retractable Face Shield:

- 1. Press and hold the Unlock Lever (see item 1 in the previous figure) in the Up position.
- Lift the Face Shield from the slots and remove from the C-arm. 2.

7.2.2 How to Use the Retractable Face Shield



Note Before you make an exposure, make sure the Face Shield is completely extended or

To extend the Face Shield, pull the Face Shield away from the C-arm until the device latches in the outer position.

To retract the Face Shield:

completely retracted.

- 1. Press a Latch Release (see item 2 in the following figure—one on each side).
- 2. Push the Face Shield toward the C-arm until the device stops.

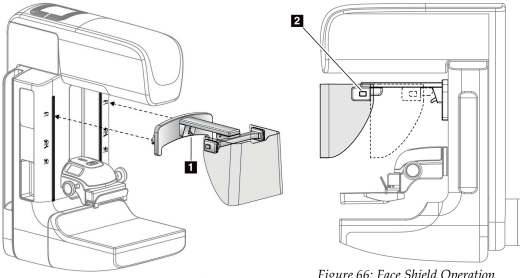
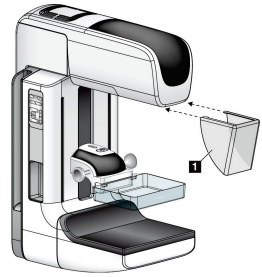


Figure 65: Face Shield Installation

Figure 66: Face Shield Operation



7.2.3 How to Install and Remove the Conventional Face Shield

Figure 67: How to Install the Conventional Face Shield

To install the Conventional Face Shield:

- 1. Carefully put the tab ends of the Face Shield (item 1 in the previous figure) into the slots at the front of the tubehead mount.
- 2. Slide the Face Shield on the tubehead mount until the Face Shield locks.

To remove the Conventional Face Shield:

- 1. Pull the sides of the Face Shield in a horizontal direction (away from the tubehead).
- 2. Remove the Face Shield.

7.3 Compression Paddles



Some paddles are optional and may not be included with your system.

The system can identify each paddle and automatically adjust the collimator. Available accessories depend on your system configuration.

Acce	ssory	2D/BT	2D Screening
Routine Screening Paddles	18 x 24 cm	*	*
	24 x 29 cm	*	*
	Small Breast	*	*
	18 x 24 cm SmartCurve®	*	*
	24 x 29 cm SmartCurve	*	*
	SmartCurve Mini	*	*
Contact and Spot	10 cm Contact	*	
Compression Paddles	15 cm Contact	*	
	7.5 cm Spot Contact	*	See Note
	Frameless Spot Contact	*	
Magnification Paddles	7.5 cm Spot Mag	*	
	10 cm Mag	*	
	15 cm Mag	*	
Localization Paddles	10 cm Rectangular Open	*	
	15 cm Rectangular Open	*	
	10 cm Perforated	*	
	15 cm Perforated	*	
	10 cm Mag Perforated Loc	*	
	10 cm Mag Localization	*	
Ultrasound Paddle	15 cm Large Ultrasound	*	
Patient Face Shield	*	*	
Magnification Stand		*	
Localization Crosshair Devic	re	*	
Magnification Crosshair Device		*	

Table 21: Available Accessories



Note On the 2D screening system, only use the 7.5 cm Spot Contact Paddle for compression thickness calibration.



Note

The 24 x 29 cm frameless screening paddle, the 24 x 29 cm SmartCurve system paddle, the magnification paddles, and the localization paddles are NOT compatible with the paddle shift function.

7.3.1 Routine Screening Paddles





18 x 24 cm Frameless Screening Paddle

24 x 29 cm Frameless Screening Paddle



Small Breast Frameless Paddle

SmartCurve System Paddles







18 x 24 cm SmartCurve System Frameless Screening Paddle

24 x 29 cm SmartCurve System Frameless Screening Paddle

SmartCurve System Mini Screening Paddle



Note

The SmartCurve system paddles may not be suitable for all patients. If the breast cannot be properly immobilized or compressed due to the curvature of the paddles, use the standard flat screening paddles.



Note

The SmartCurve system paddles are not recommended for cleavage views, rolled views, or mosaic views of very large breasts. Use the standard flat screening paddles for these views.



Note

The SmartCurve system paddles accommodate most breast sizes. Due to the curvature of the paddles, some patients who would use the smaller standard flat paddle may be more easily positioned using the larger SmartCurve paddle.



Note

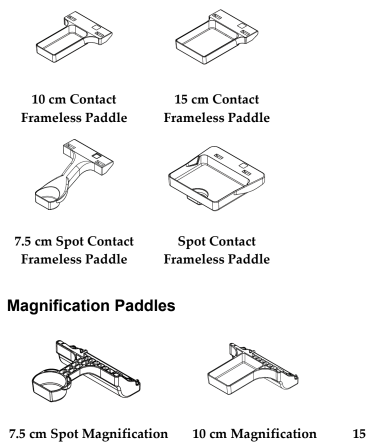
SmartCurve system paddles are not compatible with FAST Compression mode.



Note

Do not use SmartCurve system paddles for I-View procedures.

7.3.2 **Contact and Spot Compression Paddles**





Paddle

Paddle

15 cm Magnification Paddle



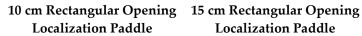
7.3.3

Note

You cannot acquire Tomosynthesis images with the Magnification paddles.

Localization Paddles 7.3.4







Localization Paddle



10 cm Magnification **Localization Paddle**





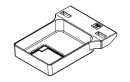
10 cm Perforated **Localization Paddle**

15 cm Perforated Localization Paddle



10 cm Magnification Localization Perforated Paddle

Large Ultrasound Paddle 7.3.5



15 cm Large Ultrasound Paddle

7.3.6 How to Install and Remove a Compression Paddle

To install a Compression Paddle:

- 1. Hold the front of the paddle with one hand in front of the Compression Device.
- 2. Tilt the paddle (between 30 and 45 degrees), then put the rear of the paddle on the groove in the rear of the Compression Device (item 1 in the following figure).
- 3. Slide the paddle along the groove until the slots on the top of the paddle are under the locks on the Paddle Clamp (item 2 in the following figure).
- 4. Compress the Paddle Clamp (item 3 in the following figure) with your free hand.
- 5. Rotate the paddle up (item 4 in the following figure).
- 6. Release the Paddle Clamp to lock the paddle.

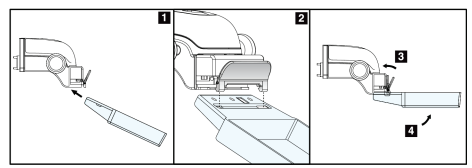


Figure 68: Installation of a Compression Paddle

To remove a Compression Paddle:

- 1. Hold the paddle with one hand while you use the free hand to compress the Paddle Clamp to release the lock (item 1 in the following figure).
- 2. Lower the paddle (item 2 in the following figure) and remove the paddle from the Compression Device (item 3 in the following figure).
- 3. Release the Paddle Clamp.

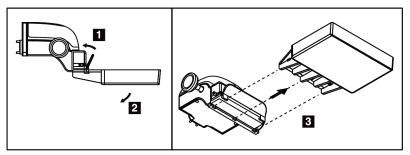


Figure 69: How to Remove a Compression Paddle

7.3.7 Paddle Maintenance and Cleaning

Clean the paddles after each use. Refer to <u>Maintenance and Cleaning</u> on page 133 for cleaning instructions.

7.3.8 Paddle Shift

The system allows most paddles to move to the left or right of the center position. The feature helps small-breast examinations with lateral views. When a lateral view is selected, the system automatically moves the collimator for the selected paddle position.



Note

The 24 x 29 cm frameless screening paddle, the 24 x 29 cm SmartCurve system paddle, and the magnification paddles are NOT compatible with the paddle shift function.

7.3.9 FAST Compression Mode

About FAST Compression Mode

The Fully Automatic Self-adjusting Tilt (FAST) Compression Mode is for use when the composition of the breast tissue does not allow uniform compression across the complete breast with a flat compression paddle. For these patients, not enough compression can cause an image to appear to be out of focus at the anterior region from both involuntary motion and not enough compression.

The FAST Compression mode used with this type of breast provides these features:

- Reduced motion artifacts, because the compression is more effective
- More uniform compression, from the chest wall to the nipple
- Maximum patient comfort, because over compression at the chest wall is prevented

When the FAST Compression mode is selected, the paddle automatically tilts when compression is applied. The paddle starts at the flat position until some compression force is applied. The paddle then tilts until its maximum angle is reached.

The FAST Compression mode does not require excessive compression, but you must use enough compression to prevent the movement of the breast. You should use a consistent amount of compression, especially for related left and right views.

The FAST Compression mode may not be best for breasts that are equal or symmetrical in thickness from the chest wall to the anterior area of the breast.



Note

Note

Only the 18 x 24 cm Frameless Screening Paddle and the 24 x 29 cm Frameless Screening Paddle are compatible with FAST Compression Mode.

$\widehat{}$

The system beeps when FAST Compression Mode is engaged but is not compatible with the current paddle.

How to Use the FAST Compression Mode Slide

To engage FAST Compression Mode, push the slide (from either side) until the "F" is visible and the slide clicks into position.

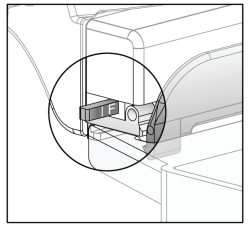


Figure 70: The FAST Compression Mode Slide

7.4 Magnification Stand

The Magnification Stand has a breast platform and an abdominal shield. When the Magnification Stand is installed, the grid automatically retracts and the x-ray exposure techniques are set to the Magnification default values. Only use the Magnification paddles when the Magnification Stand is installed (refer to <u>Magnification Paddles</u> on page 115).

7.4.1 How to Install and Remove the Magnification Stand

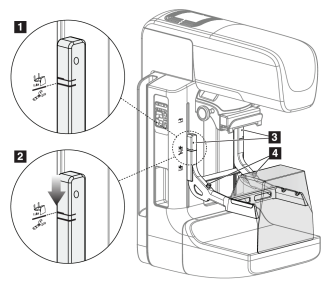


Figure 71: Installation of the Magnification Stand

To Install the Magnification Stand:

- 1. Remove the Face Shield (refer to *Patient Face Shields* on page 110).
- 2. Remove the compression paddle (refer to *How to Install and Remove a Compression Paddle* on page 117).
- 3. Move the Compression Device completely to the top.
- 4. Hold the stand on each side just below the black buttons (see item 4 in the previous figure). Do not press the black buttons.



The black buttons are used only when removing the Magnification Stand.



Note

Note

There are two sets of mounting slots for the Magnification Stand—One set is for 1.8x, and the other set is for 1.5x. Refer to *How to Install Accessories on the C-Arm* on page 109 for more information.

- 5. Align the thick black lines on the Magnification Stand with the thick black lines on the C-arm. When these lines meet, the hooks of the Magnification Stand align to the mounting slots on the C-arm. See item 1 in the previous figure.
- 6. Put the hooks of the Magnification Stand into the C-arm slots. Slide the Magnification Stand down until the thin black lines on the Magnification Stand and the black line of the C-arm meet. See item 2 in the previous figure.
- 7. The locking pins slide into holes and lock the device. You hear an audible click.



Note

If the Magnification Stand is not installed correctly, there is an indicator with a red shaft which protrudes (see item 3 in the previous figure). When the stand is installed correctly, the indicator is retracted.

To Remove the Magnification Stand:

- 1. Remove the Magnification paddle.
- 2. Hold the handles of the Magnification Stand and press the black buttons.
- 3. Lift and remove the device from the C-arm.

7.5 Crosshair Devices

7.5.1 How to Install and Remove the Localization Crosshair Device

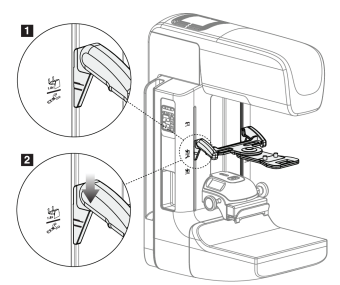


Figure 72: Installation of the Localization Crosshair Device

To Install the Localization Crosshair Device

- 1. Remove the face shield (refer to <u>Patient Face Shields</u> on page 110).
- Move the Compression Device below the mounting slots, indicated by a crosshair icon. Refer to <u>*How to Install Accessories on the C-Arm*</u> on page 109 for more information.
- 3. Hold the crosshair device by the handles and align the thick lines on the device with the line on the C-arm. Compress the release levers.
- 4. Put the hooks into the C-arm slots.
- 5. Slide the hooks toward the bottom until the thin black lines on the crosshair meet the black line on the C-arm.
- 6. Release the levers. The locking pins slide into holes and lock the device in position.

To Remove the Localization Crosshair Device:

- 1. Compress the release levers.
- 2. Lift the frame toward the top and remove the hooks from the C-arm slots.

7.5.2 How to Use the Localization Crosshair Device

- 1. The crosshair device rotates to the left or right of the tubehead. Rotate the device away from the x-ray beam during the exposure acquired with the localization paddle.
- 2. When you rotate the device back to the front for use, make sure the rotation continues until the device clicks into position.
- 3. Turn on the light field lamp.
- 4. Rotate the two crosshair knobs until the shadow on the breast matches the crosshairs on the image that identifies the suspect lesion.

7.5.3 How to Install and Remove the Magnification Crosshair Device

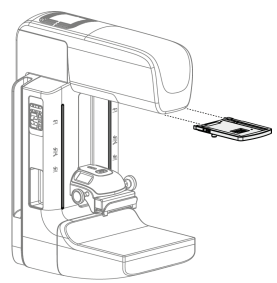


Figure 73: How to Install and Remove the Magnification Crosshair Device

To Install the Magnification Crosshair Device:

- 1. Remove the face shield (refer to *How to Install and Remove the Conventional Face Shield* on page 112).
- 2. Align the Magnification Crosshair Device with the tubehead.
- 3. Slide the crosshair device on the rails on each side of the tubehead that are used by the Conventional Face Shield. Make sure the device locks into position.
- 4. Install the remaining magnification devices.

To Remove the Magnification Crosshair Device:

- 1. Hold the sides of the device.
- 2. Pull the device toward you and remove from the tubehead.

7.5.4 How to Align the Crosshair Device

Note

If the crosshair light rectangle appears skewed to the opening in the paddle, perform this alignment procedure.

- 1. Install the rectangular localization paddle.
- 2. Loosen the adjustment lock screw on the bottom of the Crosshair Device.
- 3. Put a piece of white paper on the image receptor to make the shadows of the crosshairs easier to see.
- 4. Move the localization paddle approximately 6 cm above the image receptor.
- 5. Turn on the light field.
- 6. Rotate the Crosshair Device until the rectangle of light aligns with the opening in the localization paddle.
- 7. Tighten the adjustment screw.

7.6 MammoPad Breast Cushion Application Instructions

MammoPad[®] breast cushions are single-use disposable medical products. They are radiolucent and latex free. Use a fresh pad for each patient. Follow this procedure to make sure the pad is properly installed.

1. Open the bag containing the MammoPad breast cushions, remove one pad, and reseal the bag.



Note

Pads must be stored in a sealed plastic bag to prevent contamination that could affect image quality. Store out of direct sunlight and seal bag when not in use.

2. Peel the paper backing from the pad and discard it.

3. Apply the adhesive side to the breast platform in accordance with the instructions in the following note for your type of system. Do not apply the pad over painted decals.

Note

Selenia[®] and most other FFDM systems - Orient the rounded corner at the lower left side of the detector. Smooth the pad on from the back to the front of the breast platform.

Dimensions[®] and GE systems - Orient the rounded corner at the lower right side of the detector. Smooth the pad on from the front to the back of the breast platform.

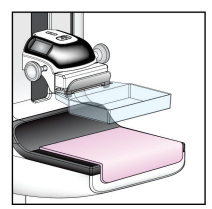


Figure 74: MammoPad Installed on Breast Platform

4. Follow your normal positioning routine for all screening views.

- 5. When the exam is complete, remove the pad from the image platform and discard it.
 - a. Grip one corner of the MammoPad and carefully pull upward and diagonally backward from the breast platform.



Figure 75: Removing MammoPad from Breast Platform

b. Once the MammoPad is detached from the top of the breast platform, carefully pull diagonally to remove from the front of the breast platform. Continue until the MammoPad has been completely removed.



Figure 76: Removing MammoPad from Breast Platform (continued)



Note

If the pad was exposed to blood or body fluids, follow the biohazard materials disposal procedure for your facility.

Chapter 8 Clinical Procedures



Warning: C-arm movement is motorized.



Warning:

Keep the hands of the patient away from all buttons and switches at all times.



Warning:

Place each footswitch in a position where, when used, they remain in reach of the Emergency Off Switches.



Warning:

Position the footswitches to prevent accidental operation by a patient or wheelchair.

8.1 Standard Workflow

8.1.1 Preparation

- 1. Select a patient from the worklist, or manually add a new patient.
- 2. Identify the required procedures.
- 3. Select the output device set if a different or additional device is needed.
- 4. Install the paddle.
- 5. Select the first view.

8.1.2 At the Gantry

- 1. Set C-arm height and rotation angle.
- 2. Make sure the light field illuminates the correct area.
- 3. Position the patient and compress the breast.

8.1.3 At the Acquisition Workstation

- 1. Set the exposure technique.
- 2. Acquire the image.
- 3. Release the patient.
- 4. Preview the image. Look at the Exposure Index to make sure that the exposure is within acceptable range.

- 5. You can use the Window/Level tool or other image review options during image preview.
- 6. Accept, Reject, or Pend the image.
- 7. Perform the Acquisition cycle as required for the requested procedures.
- 8. If necessary, add an additional view or procedure.
- 9. Make sure that the patient is safely away from the system after you complete the examination.
- 10. Close the procedure.

8.2 Screening Procedure Example

Test, Patient:	Standard S	Screening -	Тото			
	Generator Tools	Biopsy Contras	t			Update Item
	F	READ	Y	Patient Info Name: Test ID: 9876 Date of Birth: 2/3/1		Add Procedure Add View
	Conv Tomo					
RCC Tomo	Acq Mode STANDARD	AEC Mode Auto Filter	Focal Spot [LFS]	Comp Release Comp Mode Force	Manual Dual 28.3 N	Edit View Output Groups
8	- +	- +		Thickness	4.6 cm	Donuur
	kVp	mAs	Filter	Stereo Mode	Manual	Output
Accept	[30]	[Auto]	[AI]	Collimation	24x29	Archive / Export
, coopt				Paddle	24X29	Print
Reject	AEC Sensor	AEC Comp.	Grid	Mode	Normal	
	Auto	[0]	[Out]	Position	Center	Retrieve
Pend	- +			Mag	None	Retrieve
Screening - Tomo -		MLO Tomo				Close Patient
0 🚺 Manager, Te	ch (Manager)		: 0		🧈 ° 🦪 🥝	1:47:49 PM

Figure 77: Example of a Screening Procedure Screen

8.2.1 **Position the Patient**

- 1. Lift or lower the breast platform for the patient.
- 2. Move the tubehead to the projection angle.
- 3. Move the patient to the C-arm.
- 4. Position the patient as required.
- 5. Put the arm or hand of the patient on the Patient Handle or against the side of the body.

- 6. Tell the patient to keep away from system controls.
- 7. Compress the breast.
 - When possible, use the footswitch controls to provide hands-free compression control and C-arm height adjustment.
 - Use the light field lamp as necessary to see the x-ray field.
 - Apply compression slowly. As necessary, stop and make the adjustments to patient position.
 - Use the handwheels for final compression.

8.2.2 Set the Exposure Techniques

Select the exposure techniques for the procedure. Refer to <u>*How to Set the Exposure*</u> <u>*Parameters*</u> on page 86 for information.

8.2.3 Acquire the Exposure

- 1. Confirm that all exposure factors are set correctly.
- 2. If the system does not display Ready in 30 seconds, verify that the accessories are correctly installed and the paddle is locked into position. When the generator status displays **Ready**, the system is ready for exposure.



Warning:

This system can be dangerous to the patient and the user. Always follow the safety precautions for x-ray exposures.

3. Press and hold the **x-ray** button and/or the **x-ray footswitch** for the full exposure.

During the exposure:

- A System Message with the radiation symbol and a yellow background is displayed (see the following figure)
- An audible tone continues to sound during the exposure

The behavior of the audible tone during a combo exposure has changed to avoid early releases of the x-ray button and/or the x-ray footswitch by users. The audible tone is now a continuous sequence of tones. The tone sounds during the entire combo acquisition from the initiation of the exposure to the end of the conventional view. There is no interruption of the audible tone between breast tomosynthesis and conventional digital mammography exposures. Do not release the exposure switch during the audible tone.

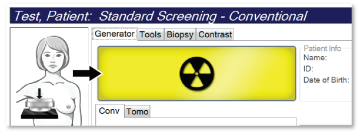


Figure 78: Exposure In Progress

4. When the tone stops and the System Message shows **Standby** (see the following figure), you can release the **x-ray** button and/or the **x-ray footswitch**.

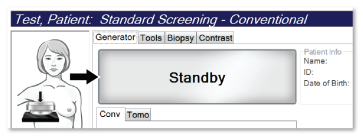


Figure 79: Exposure Complete

5. Release the compression device. If the automatic release feature is set, the compression device automatically lifts after the exposure.

8.3 **Procedure for Needle Localization with Tomosynthesis**

- 1. Install a Localization Paddle, and install the Crosshair Device at the tubehead. Be sure that the crosshair guides are out of the x-ray field.
- 2. Open a new procedure with a Tomo or TomoHD view for your approach.
- 3. Position the patient and apply compression.
- 4. Acquire a Tomo Scout. Make sure that the ROI is visible inside the Localization Paddle opening. If not, reposition the patient and repeat.
- 5. Note the Compression Thickness, and note the thickness of the excess tissue through the opening of the Localization Paddle.
- 6. Scroll through the reconstruction slices to identify where the lesion is best seen. Note the slice number (each slice is 1 mm in thickness).
- 7. Place the Acquisition Workstation crosshair on the lesion.
- 8. To find the coordinates for the Gantry Crosshair Device, scroll through the reconstructions until you can identify the alpha numeric coordinates.
- 9. Calculate the needle depth:

Value	Example
Breast Compression Thickness	50 mm
(+) Thickness of the tissue through the opening of the paddle	+ 7 mm
(-) Slice number where the lesion is found	- 30 mm
(+) Optional distance past the ROI for the wire	+ 5-15 mm
(=) Needle depth of the localization wire	32– 42 mm

- 10. Turn on the collimator light and align the Crosshair Device at the tubehead to match the Acquisition Workstation crosshair.
- 11. Position and insert the needle.
- 12. Move the Crosshair Device guides out of the x-ray field.
- 13. Acquire another Tomo image to be sure that the needle is in the correct location. To calculate if a correction is necessary, compare the slice number of the point of the needle and the slice number of the lesion.
- 14. Insert the guide wire through the needle, then remove the needle, if desired, leaving the wire in position.
- 15. If desired, complete the following steps:
 - a. Acquire a Conventional or Tomo view to be sure of correct wire placement.
 - b. Take the orthogonal view to document wire or needle placement (either in Tomo or conventional).
- 16. Only add one view icon at a time for orthogonal views to remove the possibility of paddle shift due to possible minimal compression.

Example: Calculating Needle Depth with Tomosynthesis

In this example, use the values from the table on the previous page and see the following figure.

Calculate the needle depth from the tissue skin line (item 1) rather than from the localization paddle (item 9). Insert the needle a minimum of 27 mm (breast compression + bulging tissue).

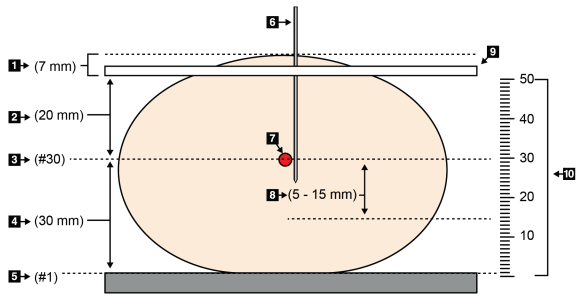


Figure 80: Calculating needle depth

Item	Description	Example
1	Thickness of the tissue through the opening of the localization paddle	7 mm
2	Thickness measured from the localization paddle to the lesion	
3	Lesion slice number (the slice number where lesion is best seen (clearest))	30 mm
4	Thickness measured from the detector to the lesion	
5	Slice number 1	
6	Needle	
7	Lesion	
8	Advancing the needle 5 - 15 mm more than the lesion (optional)	5 - 15 mm
9	Localization paddle	
10	Thickness of the breast compression from the detector (0 mm) to the localization paddle (50 mm in this example)	50 mm

Chapter 9 Maintenance and Cleaning

9.1 Cleaning

9.1.1 General Information About Cleaning

Before each examination, clean and use a disinfectant on any part of the system which touches a patient. Give the attention to the paddles and the image receptor.



Caution:

Do not use any hot source (like a heating pad) on the image receptor.

Be careful with the compression paddles. Inspect the paddles. Replace the paddles when you see damage.

9.1.2 For General Cleaning

Use a lint-free cloth or pad and apply a diluted dishwashing liquid.



Caution: Use the least possible amount of cleaning fluids. The fluids must not flow or run.

If more than soap and water is required, Hologic recommends any one of the following:

- 10% chlorine bleach solution and water with one part commercially available chlorine bleach solution (normally 5.25% chlorine and 94.75% water) and nine parts water. Mix this solution daily for best results.
- Commercially available isopropyl alcohol solution (70% isopropyl alcohol by volume, not diluted)
- 3% maximum concentration of hydrogen peroxide solution

After you apply any of the above solutions, use a pad and apply a diluted dishwashing liquid to clean any parts that touch the patient.



Warning:

If a paddle touches possible infectious materials, contact your Infection Control Representative to remove contamination from the paddle.



Caution:

To prevent damage to the electronic components, do not use disinfectant sprays on the system.

9.1.3 To Prevent Possible Injury or Equipment Damage

Do not use a corrosive solvent, abrasive detergent, or polish. Select a cleaning/disinfecting agent that does not damage the plastics, aluminum, or carbon fiber.

Do not use strong detergents, abrasive cleaners, high alcohol concentration, or methanol at any concentration.

Do not expose equipment parts to steam or high temperature sterilization.

Do not let liquids enter the internal parts of the equipment. Do not apply cleaning sprays or liquids to the equipment. Always use a clean cloth and apply the spray or liquid to the cloth. If liquid enters the system, disconnect the electrical supply and examine the system before returning it to use.



Caution:

Wrong cleaning methods can damage the equipment, decrease imaging performance, or increase the risk of electric shock.

Always follow instructions from the manufacturer of the product you use for cleaning. The instructions include the directions and precautions for the application and contact time, storage, wash requirements, protective clothing, shelf life, and disposal. Follow the instructions and use the product in the safest and most effective method.

9.1.4 Acquisition Workstation

How to Clean the Image Display Screen:

Avoid touching the display screen of the Image Display monitor.

Use care when cleaning the outer surface of the LCD screen. Always use a clean, soft, lint-free cloth to clean the display area. Microfiber cloths are recommended.

- Never use a spray or flow a liquid on the display.
- Never apply any pressure to the display area.
- Never use a detergent with fluorides, ammonia, alcohol, or abrasives.
- Never use any bleach.
- Never use any steel wool.
- Never use a sponge with abrasives.

There are many commercially available products to clean LCD displays. Any of the products free of the listed ingredients and used according to the directions of the manufacturer can be used.

How to Clean the Touchscreen Display:

Use a window or glass cleaning product to clean the Touchscreen display. Apply the cleaning product to a cloth, then clean the Touchscreen display. Do not apply the cleaning product to the display without the cloth.

How to Clean the Keyboard:

Wipe the surfaces with a CRT wipe. If necessary, clean the keyboard with a vacuum. If liquids enter the keyboard, contact Technical Support for a replacement.

How to Clean the Fingerprint Scanner:



Caution:

To protect the Fingerprint Scanner:

- Do not apply any liquid product directly on the Fingerprint Scanner window.
- Do not use products that contain alcohol.
- Never put the Fingerprint Scanner under liquid.
- Never apply any pressure to the Fingerprint Scanner window with abrasive material.
- Do not push the Fingerprint Scanner window.

To clean the Fingerprint Scanner window, do one of the following:

- Apply the adhesive side of cellophane tape, then remove the tape.
- Apply a product with ammonia base to a cloth and clean the Fingerprint Scanner window.

9.2 Maintenance

9.2.1 Preventive Maintenance Schedules

Table 22: User Preventive Maintenance						
	Recommended Frequency					
Maintenance Task Description	Each Use	Weekly	Biweekly	Monthly	Bimonthly	Semiannually
Clean & disinfect paddle	\checkmark					
Clean & disinfect breast platform	\checkmark					
Visually inspect all paddles for damage	✓					
Detector Flat Field Calibration *		✓				
Artifact Evaluation *		✓				
Phantom Image *		✓				
Signal to Noise / Contrast to Noise Measurements *		✓				
Geometry Calibration (Tomosynthesis Option) *						✓
Compression Thickness Indicator *			✓			
Visual Checklist *				✓		
Compression *						✓
* Refer to Quality Control Manual						

Table 22: User Preventive Maintenance

Table 23: Service Engineer Preventive Maintenance

Meintenne Tel Descision	Recommended Frequency	
Maintenance Task Description	Semiannually	Annually
Clean and Inspect the Gantry and Acquisition Workstation	✓	
Inspect the radiation shield for chips, cracks, breaks, and for tight attachments.	✓	
Check all primary power connections	✓	
Check interlocks, safety and limit switches	✓	
Inspect/Lubricate C-arm	✓	
C-arm / Verify all C-arm buttons	✓	
Verify C-arm and Rotational Calibration	✓	

Maintonanao Task Description	Recommended Frequency		
Maintenance Task Description	Semiannually	Annually	
Replace Breast Platform Filter	✓		
Verify Compression Force Calibration	✓		
Verify Compression Thickness Calibration	✓		
Inspect LED Collimator Lamp for dust and dirt	✓		
Clean & lubricate collimator, and worm screws	✓		
Perform Rotational Brake verification	✓		
Verify X-ray Field / Light Field Calibration	✓		
Verify kV Calibration and Tube Current Calibration	✓		
Check HVL Evaluation	✓		
Verify Target Dose Verification	✓		
Verify AEC Exposure Compensation 2D	✓		
Perform System Resolution Test *	✓		
Perform Phantom Image Quality Evaluation *	✓		
Perform Image Artifact Evaluation *	✓		
Backup Acquisition Workstation files	✓		
Evaluate UPS Performance Status/ Batteries Status	✓		
Backup all Calibration Data	✓		
* Refer to Quality Control Manual			

 Table 23: Service Engineer Preventive Maintenance

9.2.2 About Reclamation

Reclamation is an automatic function that makes disk space available for storing newly acquired images. Configurable parameters let a given number of images collect before reclamation starts and older images are removed from the system.

Chapter 10 System Administration Interface

10.1 Admin Screen

Note

This section describes the functions available in the *Admin* screen. To access all the functions in this screen, log in to the system as a user with Administrator, Manager, or Service permissions.

Refer to the table on the following page for descriptions of the Admin screen functions.



Depending on the license settings for your system, you may see different buttons.



Figure 81: Admin Screen

Table 24: Admin Screen Functions

Group	Button	Function
Operators	Manage Operators	Add, delete, or change Operator information.
	My Settings	Change the information for the current Operator.
Procedures	Procedure Editor	Add or Edit the procedures, or change the view order for each user.

Group	Button	Function		
	Procedure Order	View or change the order of the procedures within a procedure group.		
	View Editor	Set the default view order for a procedure and edit individual views.		
	Contrast	Access the contrast enhanced digital mammography functionality and configure the default settings.		
Quality Control	Quality Control	Select a Quality Control task to perform or mark completed.		
	QC Report	Create a QC Report.		
	Test Patterns	Select and send the test patterns to output devices.		
	Reject and Repeat Report	Create a Reject and Repeat Report.		
System	System Tools	The Interface for Service for the configuration of or the identification of problems in the Acquisition Workstation.		
	System Defaults	Set the Gantry default values.		
	System Diagnostics	Shows the status of all subsystems.		
	Preferences	Set the system preferences.		
	About	Describes the system. Refer to <u>About Screen</u> on page 141.		
	Exposure Report	Create a radiation Exposure Report.		
	System Security	Access account security settings, Computer Management, Local Security Policy, Local Users and Groups, and Local Group Policy in the Windows OS		
	Biopsy Devices	View and configure available biopsy devices.		
	QAS	Access the QAS Needle Test screen.		
	Lateral QAS	Access the Lateral QAS Needle Test screen.		
	Eject USB	Eject a media storage device connected to the USB port.		
Connectivity	Query Retrieve	Query and retrieve patient information from the configured devices.		
	Import	Import the data from a DICOM source.		
	Manage Output Groups	Add, delete, or edit output groups.		
	Archive	Send local studies to networked storage or export to removable media devices.		

Table 24: Admin Screen Functions

and change.

10.2 About Screen

The *About* screen provides information about the machine, such as system level, IP address, and Gantry serial number. This type of data can be useful when you are working with Hologic to resolve a system issue or configure the system. To access the *About* screen, select **About** from the System group in the *Admin* screen.



Note You can also access the *About* screen through the taskbar. Select the **System Status** (tubehead) icon, then select **About**....

System Licensing Institutio	n Copyright UDI	
Computer System AWS Computer Rev Build Date Last Boot Time IP Address GIP2D GIP3D Filter GIP3D Filter GIP3D CadScience GIP3D CadScience Contrast IP Auto SNR/CNR M35 Dose Calculation Method PCI Driver	2 0 0.2 1 0 0.2 (200 - 2 200 - 2 20	Gantry Serial Number AIO BKY BKY CPLD CDI CRM DET DTC GCB GEN GEN PMC THD CPLD VTA
PCI Firmware HARI Video Card #0 NVidia	8 - 1 - 92 1 - 82 80 (104: Galfanas (171: 1070 (1 - 1 - 1 - 108)	Detector Detector Temperature Serial Number Read Out Sequence Model Id Hardware Revision CPU Firmware Analog Firmware Version DTC Firmware

Figure 82: System Tab of the About (the Acquisition Workstation) Screen

There are five tabs on the *About* screen:

- System Tab (default) lists system configuration information
- Licensing Tab lists the Hologic-licensed options installed on this machine
- **Institution** Tab lists the name and address of the organization and the Station Name assigned to this machine
- **Copyright** Tab lists the copyrights of Hologic and third-party software installed on this machine
- **UDI** Tab lists the unique device identifier(s) of this machine

10.2.1 Licensing Tab

The Licensing tab of the *About* screen shows all the licenses installed on your system.



Note

Hologic configures some systems to meet specific requirements. Your system configuration may not have all the options and accessories included in this manual.



Note

The features shown in this manual may not be available in all regions. Contact your Hologic representative for information.

ystem Licensing Instit	isition Workstation	
Computer		Refresh
Dimensions	Licensed	
BDQuorum	Licensed	
Advanced BioMed	Licensed	
dvanced Connectivity	Licensed	
iopsy	Licensed	
iopsy Specimen	Licensed	
larity HD	Licensed	
ontrast Biopsy	Licensed	
-View 1	Licensed	
-View Natural Look	Licensed	
iagnostic	Licensed	
nageChecker CAD	Licensed	
ntelligent 2D	Licensed	
telligent 2D Map	Licensed	
View CE2D	Licensed	
ateral Arm	Licensed	
laster	Licensed	
lotices	Licensed	
Quantra	Licensed	
SmartCurve	Licensed	
SmartCurve 2	Licensed	
System	Licensed	
omo	Licensed	
omo Biopsy	Licensed	
		Back

Figure 83: Licensing Tab of the About Screen

10.3 Change the User Language Preference

Users can set the language on the user interface to automatically change to their individual preference when logging in.

1. In the Operators group of the *Admin* screen, select **My Settings**.



You can also access **My Settings** through the taskbar. Select the User Name area then select **My Settings** in the pop-up menu.

- 2. The **Users** tab of the *Edit Operator* screen opens. From the Locale field, select a language from the drop-down list.
- 3. Select **Save**, then select **OK** to the *Update Successful* message. The user interface changes to the selected language.

10.4 Set Auto-Hanging and Auto-Pairing

To set the system for Auto-Hanging and Auto-Pairing of images:

1. In the Operators group of the Admin screen, select My Settings.

Note

You can also access **My Settings** through the taskbar. Select the User Name area, then select **My Settings** in the pop-up menu.

- 2. The *Edit Operator* screen opens. Select the **Workflow** tab.
 - Select the Auto-Hanging check box to show a prior study in 4-up mode automatically.
 - Select the Auto-Pairing check box to show a prior view in multi-up mode next to a newly captured image.

Edit Operator: Manager, Tech	,
User Workflow	^
Auto Hanging	
Enable Auto Hanging	\bigcirc
CC image placement	Тор
Left laterality image placement	Right
Auto Hanging repeated views	Hang older
Auto Pairing	
Enable Auto Pairing	\bigcirc
Auto Pairing protocol	On thumbnail selection
Auto Pairing repeated views	Pair older
Match Chest Walls	\bigcirc

Figure 84: Enable Auto-Hanging and Auto-Pairing

3. Select **Save**, then select **OK** to the *Update Successful* message.

10.5 Set Multi Line Procedure Tabs

You can set the operator preferences to display more of the procedure name in the top of the procedure tabs. To change the procedure tabs from a single line of text to multiple lines of text:

1. In the Operators group of the *Admin* screen, select **My Settings**.



Note

You can also access **My Settings** through the taskbar. Select the User Name area, then select **My Settings** in the pop-up menu.

- 2. The *Edit Operator* screen opens. Select the **Workflow** tab.
- 3. Select the Use Multi Line Procedure Tabs check box.

ser Workflow	
Auto Hanging	
Enable Auto Hanging	\bigcirc
CC image placement	Тор
Left laterality image placement	Right
Auto Hanging repeated views	Hang older
Auto Pairing	
Enable Auto Pairing	\bigcirc
Auto Pairing protocol	On thumbnail selectio
Auto Pairing repeated views	Pair older
Match Chest Walls	
Match Chest Walls Procedure Display	

Figure 85: Enable Multi Line Procedure Tabs

4. Select **Save**, then select **OK** in the *Update Successful* message.

10.6 Enable and Set the Height Memory

Users can enable and set the acquisition workstation height to automatically change to their individual preference when logging in. To enable and set the height adjust memory:

1. In the Operators group of the *Admin* screen, select **My Settings**.



Figure 86: My Settings Button in the Admin Screen



Note

You can also access **My Settings** through the taskbar. Select the User Name area, then select **My Settings** in the pop-up menu.

- 2. In the *Edit Operator* screen, select the **Console** tab.
- 3. To enable the height adjust memory, select the radio button to the right of the 'Auto-Height adjustment on login' field. A check mark appears. (To disable the height adjust memory, clear the radio button.)

Edit Operator: Tech, Radiolog	nical	
User Workflow Console	^	
User's desired console height		
Auto-Height adjustment on login		
Desired console height	83.8 cm (33.0 in)	Apply
Current console height		
Current console height	83.8 cm (33.0 in)	

Figure 87: Console Tab of the Edit Operator Screen

4. Use the ▲ UP and ▼ DOWN buttons on the height adjust control panel to set the desired height (see the following figure).

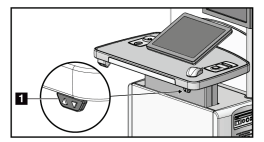


Figure 88: Height Adjust Control Panel

5. The 'Desired console height' field displays the height as it is now positioned. The 'Current console height' field displays the most recently saved height. (See the following figure.) To save your desired height setting, select **Apply**.

Edit Operator: Tech, Radiolog	nical	
User Workflow Console		
User's desired console height		
Auto-Height adjustment on login	\checkmark	
Desired console height	83.8 cm (33.0 in)	Apply
Current console height		
Current console height	99.8 cm (39.3 in)	

Figure 89: Desired Console Height and Current Console Height Fields

6. Select **Save**, then select **OK** to the *Update Successful* message.

10.7 Enable and Set the Default Height

A Manager user can set the acquisition workstation to automatically return to a default height when a user logs out. To enable and set the default height:

1. In the System group of the *Admin* screen, select **Preferences**.

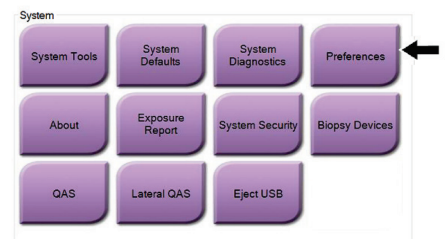


Figure 90: Preferences Button in the Admin Screen

- 2. In the *System Preferences* screen, select the **Console** tab.
- 3. To enable the default height, select the radio button to the right of the 'Auto-Height adjustment on logout' field. A check mark appears. (To disable the default height option, clear the radio button.)

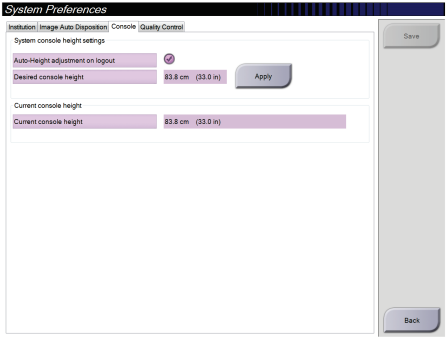


Figure 91: Console Tab of the System Preferences Screen

4. Use the ▲ UP and ▼ DOWN buttons on the height adjust control panel to set the desired height (see the following figure).

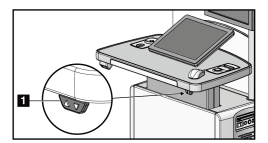


Figure 92: Height Adjust Control Panel

5. The 'Desired console height' field displays the height as it is now positioned. The 'Current console height' field displays the most recently saved height. (See the following figure.) To save your desired height setting, select **Apply**.

System Preferences	
Institution Image Auto Disposition Console Quality Control	
System console height settings	
Auto-Height adjustment on logout	
Desired console height 83.8 cm (33.0 in) Ap	ply
Current console height	
Current console height 121.2 cm (47.7 in)	

Figure 93: Desired Console Height and Current Console Height Fields

6. Select **Save**, then select **OK** to the *Update Successful* message.

10.8 Set Auto-Accept and Auto-Pend Images

A Manager user can configure the system to automatically accept or automatically pend new images.

- 1. In the System group of the *Admin* screen, select **Preferences**. The *System Preferences* screen opens.
- 2. Select the Image Auto Disposition tab.

- 3. Use the drop-down menus to select the auto disposition for each type of image.
 - Select Manual to manually accept, reject, or pend each newly acquired image.
 - Select Accept to automatically accept newly acquired images.
 - Select **Pend** to automatically pend newly acquired images.

stitution Image Auto Disposition Quality Co	ontrol		
mage Auto Disposition			Save
Conventional Images	Accept	•	
Tomosynthesis Images	Accept	•	
Combination Images	Manual	•	
FomoHD Images	Pend	•	
ComboHD Images	Pend	 •	
Biopsy Images	Accept	•	
			Back

Figure 94: Set Image Auto Disposition

4. Select **Save**, then select **OK** to the *Update Successful* message.

10.9 Set the Contrast Defaults

A Manager user can configure the default timer periods and the default contrast information.

Set the Default Timer Periods

1. From the Procedures group in the *Admin* screen, select the **Contrast** button.

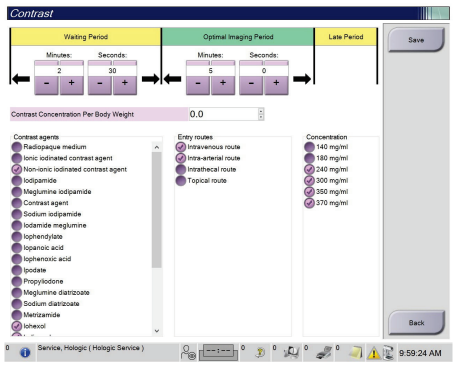


Figure 95: I-View 2D Contrast Default Settings

- 2. Choose the plus (+) or minus (-) buttons to change the Minutes and Seconds for the **Waiting Period** and **Optimal Imaging Period**.
- 3. Select Save.

Your selections appear as the default timer settings on the Contrast tab.

Set the Default Contrast Information

- 1. From the Procedures group in the *Admin* screen, select the **Contrast** button.
- 2. Select one or more **Contrast agents**, **Entry routes**, and **Concentration**. See the previous figure.
- 3. Select Save.

Your selections appear as the default options in the Contrast Information dialog box.

10.10 Set Genius AI Detection Software Preferences

A procedure can occasionally require repeated views. A Manager user can set preferences for the image to use for Genius AI Detection software case processing when there are duplicate views.

To set the default system selection of duplicate views:

- 1. In the System group of the *Admin* screen, select the **Preferences** button. The *System Preferences* screen opens.
- 2. Select the **Genius AI Detection** tab.
- 3. In the 'Genius AI Case Processing Automatic Duplicate View Selection' field, select **Newest** or **Oldest**.
 - **Newest** sets the system to automatically select the newest duplicate view for case processing.
 - **Oldest** sets the system to automatically select the oldest duplicate view for case processing.

System Prefe	erences	
Institution Image Aut	o Disposition Quality Cont	rol Genius Al Detection
Enable User View S Case Processing	election for Genius Al	
Genius Al Case Pro Duplicate View Sele	cessing Automatic ction	Newest

Figure 96: Automatic Selection for Genius AI Case Processing

4. Select **Save**, then select **OK** to the *Update Successful* message.

To enable user selection of the duplicate views:

- 1. In the System group of the *Admin* screen, select the **Preferences** button. The *System Preferences* screen opens.
- 2. Select the Genius AI Detection tab.
- 3. To enable user selection, select the radio button for 'Enable User View Selection for Genius AI Case Processing'. A check mark appears. (To disable user selection, clear the radio button.)

Syster	n Preferences	
Institution	Image Auto Disposition Qualit	Control Genius Al Detection
	ser View Selection for Genius	
	I Case Processing Automatic View Selection	Newest

Figure 97: Automatic Selection for Genius AI Software Case Processing

4. Select **Save**, then select **OK** to the *Update Successful* message.

When a duplicate view is acquired and accepted, the system shows that the user selected the image for Genius AI Detection software case processing. The **Tools** tab of the *Procedure* screen shows a large check mark. The image thumbnail that was selected for case processing is identified with a small check mark. (See the following figure.)

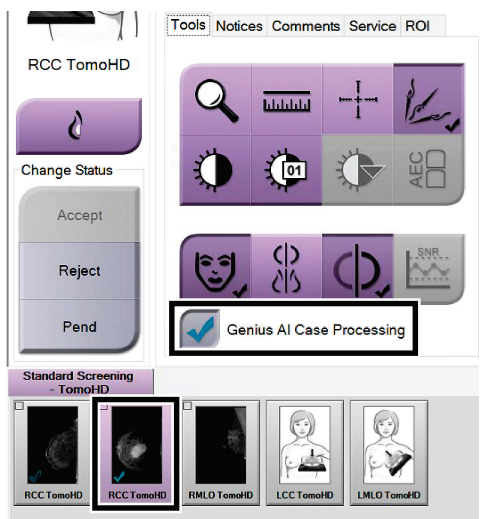


Figure 98: User Selection for Genius AI Detection Software Case Processing

10.11 System Tools

The Radiologic Technologist Managers and users with Service permissions can access the System Tools utility. The System Tools utility contains the configuration information about the system. To access the utility, select **System Tools** from the System group in the *Admin* screen.

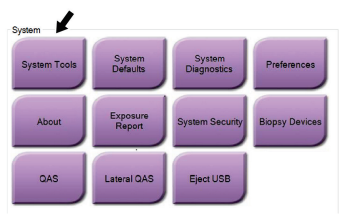


Figure 99: System Tools Button

10.11.1 System Tools for the Radiologic Technologist Manager

() Ill http://localhost/Hologic.Web	ク -	- D
HOLOGIC	System Tools	Logged in as Tech
The Science of Sure	Site Name: abcdefghijklimnopqrstuvwxyzABCDEFGHIJKLMNOPQRSTUVWXYZ0123456 Host Name:	
	K Welcome	
Search	Welcome	
	➤ Getting Started	
Welcome	≻ AWS	
- O Getting Started	➤ Peripherals	
- O AWS - O Peripherals		
- O Hardware	► Hardware	
• Troubleshooting	> Troubleshooting	
	© 2018 - Hologic System Tools	
1		

Figure 100: System Tools Screen

Section	Screen Functions			
Getting Started	About: The introduction to the service tool.			
	FAQ: List of common questions.			
	Glossary: List of terms and descriptions.			
	Platform: List of directories, software version numbers, and system software statistics.			
	Shortcuts: List of Windows shortcuts.			
AWS	Connectivity: List of Installed Devices.			
	Film & Image Information: Create an Image Report*. Create a QC			
	Report. (*You can also access this report from a remote computer.			
	Refer to <u>Remote Access to Image Reports</u> on page 154.)			
	Licensing: List of Installed Licenses.			
	User Interface: Change the options in the Software application.			
	Internationalization: Select the local language and culture.			
Troubleshooting	AWS: Allows for download of images.			
	Computer: System Management and Network Information.			
	Log: Change the event record options.			
	Backups: Control the backups for the system.			

Table 25: Radiologic Technologist Manager—System Tools	Functions
--	-----------

10.11.2 Remote Access to Image Reports

Access image reports via a remote computer networked to the system. This function can be useful for sites that do not permit USB downloads of reports directly from the system.

Follow these steps to access image reports from a remote computer. You must log in to the System Tools as a Manager-level user for this procedure.

- 1. Get the IP Address for the system you want to access. You can get the IP Address from your IT administrator or from the system. From the system, go to the *About* screen and select the **System** tab. Write down the IP Address.
- 2. Using an internet browser on your remote computer, navigate to http:// [IP address]/Hologic.web/MainPage.aspx. Use the IP Address from step 1.

3. The *System Tools Login* screen opens. Type a Manager-level user name and password, then select the **Log in** button.



Figure 101: System Tools Login Screen

4. The *System Tools Welcome* screen opens. Go to **AWS** > **Film & Image Information** > **Create Image Report.**

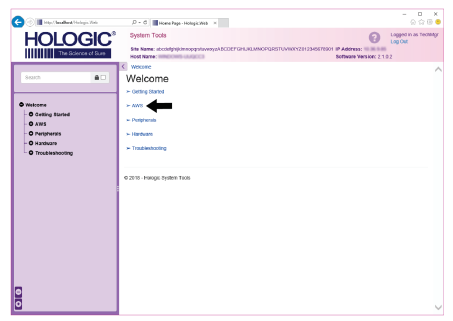


Figure 102: System Tools Welcome Screen

5. Select the parameters for the report and select **Generate**.

	ー ロ クー C III Home Page - Hologic.Web ×	× 00 🙂
HOLOGIC	System Tools Coged in as Tech Site Name: abcdefphildmnopqrstuwwyzABCDEFGHUKLMNOPQRSTUWWYZ0123456778901 IP Address:	Mgr
The Science of Sure	Host Name: Software Version: 2.1.0.2	
Search	Velcone – AWS – Film & Image Information – Create Image Report Create Image Report	^
Welcome O Getting Started O AWS O Peripherals O Hardware	Last Quarter Last 30 Days From: 2018-10-16 To: 2018-11-15 Techs ALL •	
• O Troubleshooting	Type Reject - Format Report - Source Default - Generate Reset	
	© 2018 - Hologic System Tools	
⊖ ⊙		~

Figure 103: Create Image Report Parameters

 The report shows on the screen. Scroll to the bottom of the report and select either Download to (html) or Download to (csv) for the file download type. Select Save when prompted.

🗲 🛞 🏢 http://localhost/Hologic.Web			eb ×							- គេជះ	× @ 0
HOLOGIC® The Science of Sure		System Tools Site Name: abcdefghijkimnopgrstur Host Name:		DEFGHIJKL	MNOPQRS	TUVWXYZ	012345678		ress: re Version: 2.1.0		chMgr
	<	6. Incorrect Patient ID	0	0	0	0	0	0	0	0%	^
Search 🔒 🗌		7. X-ray Equipment Failure	0	0	0	0	0	0	0	0%	
	ΙC	8. Software Failure	0	0	0	0	0	0	0	0%	
• Welcome	ΙC	9. Blank Image	0	0	0	0	0	0	0	0%	
- O Getting Started		10. Wire Localization	0	0	0	0	0	0	0	0%	
- O AWS - O Peripherals	۱L	11. Aborted AEC Exposure	0	0	0	0	0	0	0	0%	
- O Hardware	۱L	12. Other	0	0	0	0	0	0	0	0%	
• Troubleshooting	L						Totals:		0	100%	
				1							
	∣⊢	Total with Reasons:	0								
		Total Exposures:	0								
	ΙL	Ratio (%):	0%	l							
											1
		Remarks:									
	۱ŀ	Corrective Action:									
	L										
0		ownload html (Right click to downloa ownload html (Right click to downloa									
0	© 2	1018 - Hologic System Tools									\sim

Figure 104: Download the Image Report

- 7. Select a folder on the computer, then select **Save**.
- 8. Log out from System Tools when finished.

10.12 Archive Tool

The archive feature in the *Admin* screen lets you:

- Send local studies to an archive.
- Export studies to removable media.

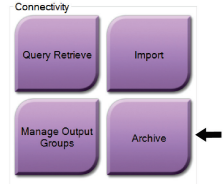


Figure 105: Archive Button

- 1. From the Connectivity group in the *Admin* screen, select the **Archive** button. The *Multi Patient On Demand Archive* screen opens.
- 2. To search for a patient, enter at least two characters in the Search parameters area and select the magnifying glass.

Multi Patient On Demand Archive Patient Name test \gg \times Device List Header Patient ID Study Date Study Time Accessi Group Details 🕢 test 3456467 Standard Screening - TomoHD 3456467 20170412 114736 Output Group Standard Screening - TomoHD 20170412 3456467 115026 Default Standard Screening - Tomo 3456467 20170412 114646 Test^Patient 123456 2-4 Standard Screening - Combo 123456 20170428 134152 Select All Test^Patient 987654 Clear 🕢 Standard Screening - Tomo 987654 20170428 134832 Name Patient ID Date of Birth 3456467 12/31/1964 Export Test, Patient 987654 2/3/1967 Archive 3 5 Back Manager, Tech (Manager) J 0 0 --:--2 0 🔊 🧻 🥥 🔃 1:50:34 PM

A list of patients that match the search criteria is displayed.

Figure 106: Multi Patient On Demand Archive Screen

Figure Legend

- 1. Search parameters
- 2. Patient List area
- 3. Patients to Be Archived or Exported area
- 4. Add selection in the Patient List area to the Patients to Be Archived or Exported area
- 5. Remove the selection from the Patients to Be Archived or Exported area

To Archive:

- 1. Select the patients and the procedures to archive.
 - Select patients from the patient list, or do a search with the search parameters (item 1) and select patients from the search results.



Note

The **Select All** button (on the right side of the screen) selects all the patients in the Patient List area. The **Clear** button (on the right side of the screen) clears selections.

- Select the procedures for each patient.
- Select the **Down Arrow** (item 4) on the screen to move the selected patients to the Patients To Be Archived area (item 3).
- Select the **Up Arrow** (item 5) on the screen to remove the selected patients from the Patients To Be Archived area (item 3).
- 2. Select a storage device.
 - Select an option from the Store Device drop-down menu.
 - -OR-
 - Select the **Group List** button, then select an option.
- 3. Select the **Archive** button. The list in the Patients To Be Archived area copies to the selected archive devices.



Note

Use the Manage Queue utility in the taskbar to review the archive status.

To Export:

- 1. Select the patients and the procedures to export.
 - Select patients from the patient list, or do a search with one of the search parameters (item 1) and select patients from the search results.



Note

The **Select All** button (on the right side of the screen) selects all the patients in the Patient List area. The **Clear** button (on the right side of the screen) clears selections.

- Select the procedures for each patient.
- Select the **Down Arrow** (item 4) on the screen to move the selected patients to the Patients To Be Archived area (item 3).
- Select the **Up Arrow** (item 5) on the screen to remove the selected patients from the Patients To Be Archived area (item 3).
- 2. Select the **Export** button.

3. In the *Export* dialog box, select the Target from the drop-down list of media devices.

Export		
Target	Removable Disk (E	
Progress		
Anonymize		Start
Eject USB device after write Advanced		Close

Figure 107: Export Screen

- 4. Select other options, if necessary:
 - **Anonymize**: to anonymize patient data.
 - **Eject USB device after write**: to automatically eject the removable media storage device when the export is complete.
 - **Advanced**: to select a folder on your local system for storage of your selections, and also to select the Export types of the images.
- 5. Select the **Start** button to send the selected images to the selected device.

Appendix A Specifications

A.1 Product Measurements

A.1.1 Tubestand (Gantry with C-Arm)

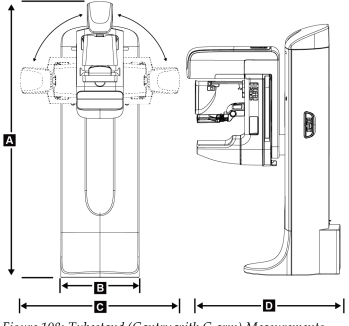


Figure 108: Tubestand (Gantry with C-arm) Measurements

А.	Height	223 cm (87.8 inches)
В.	Width	66 cm (26 inches)
С.	Width	173 cm (68 inches)
D.	Depth	138 cm (54.3 inches)
	Weight	Maximum of 400 kg (882 pounds)

A.1.2 Acquisition Workstations

Universal Acquisition Workstation

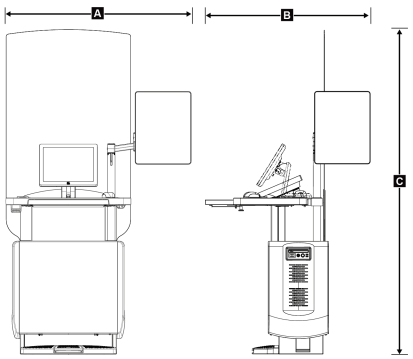


Figure 109: Universal Acquisition Workstation Measurements

А.	Width (maximum) with optional articulated display arm extended	136 cm (53.4 inches) - series I UAWS
		128 cm (50.3 inches) - series II UAWS
	Width (maximum) with standard display arm	94.0 cm (36.9 inches) - series I UAWS
		107 cm (42.0 inches) - series II UAWS
В.	Depth (maximum) with keyboard tray extended and optional articulated display monitor arm	122 cm (48.4 inches) - series I UAWS, rotated to the side
		115 cm (45.1 inches) - series II UAWS, rotated to the side
	Depth (maximum) with keyboard tray extended and standard display arm	83.6 cm (32.9 inches) - series I and II UAWS
С.	Height (nominal)	219 cm (86.1 inches) after August 2017
		204 cm (80.3 inches) before September 2017
	Weight (maximum)	209 kg (460 pounds)



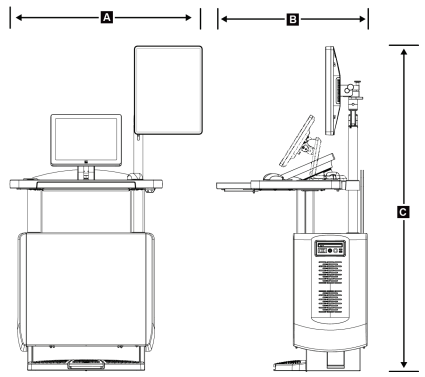


Figure 110: Mobile Universal Acquisition Workstation Measurements

А.	Width (maximum) with mobile display arm	100 cm (39.5 inches) - series I UAWS 107 cm (42.0 inches) - series II UAWS
В.	Depth (maximum) with keyboard tray extended	85 cm (33.5 inches)
С.	Height (maximum)	180 cm (71 inches)
	Weight (maximum)	179 kg (395 pounds)

A.2 Operation and Storage Environment

A.2.1 General Conditions for Operation

Temperature Range	20 °C (68 °F) to 30 °C (86 °F)
Relative Humidity Range	20% to 80% without condensing moisture

A.2.2 Storage Environment

Temperature Range	–10 °C (14 °F) to 40 °C (104 °F)
Relative Humidity Range	10% to 90% without condensing moisture

(Put in a package for storage in a building.)

X-ray Detector

Temperature Range	10 °C (50 °F) to 30 °C (86 °F) indefinitely	
	10 °C (50 °F) to 35 °C (95 °F) for a maximum of 12 hours	
Maximum rate of temperature change	Less than 10 °C (50 °F) per hour	
Relative Humidity Range	10% to 80% without condensing moisture	

(Put in a package for storage in a building.)

Acquisition Workstation

Temperature Range	–10 °C (14 °F) to 40 °C (104 °F)	
Relative Humidity Range	10% to 90% without condensing moisture	
(Put in a package for storage in a building.)		

A.3 Radiation Shield

Radiation Shield Lead (Pb) equivalent 0.5 mm lead for x ray energy to 35 kV

A.4 Electrical Input

A.4.1 Tubestand

Mains Voltage	200/208/220/230/ 240 VAC ±10%
Mains Impedance	Maximum line impedance not to exceed 0.20 ohms for 208/220/230/240 VAC, 0.16 ohms for 200 VAC
Mains Frequency	50/60 Hz ±5%
Average Current over 24 Hours	< 5 A
Peak Line Current	$4 A (65 A maximum for \le 5 seconds)$

A.4.2 Acquisition Workstation

Mains Voltage	100/120/200/208/220/230/240 VAC ±10%
Mains Frequency	50/60 Hz ±5%
Power Consumption	< 1000 watts
Duty Cycle (Standard Acquisition Workstation)	10% ~ 6 minutes per hour or 2 minutes on, 18 minutes off
Overcurrent Protection	8A

A.5 Tubestand Technical Information

A.5.1 C-Arm

Rotation Range	Conventional Mammography:
	+195° +3°/–0.5° to 0° ±0.5° to –155° +0.5°/–3°
	Tomosynthesis option:
	+180° ±0.5° to 0° ±0.5° to -140° ±0.5°
Absolute Angular Position	accurate to $\pm 0.5^{\circ}$
Rotation Acceleration	18°/s² +18/-9%
Rotation Deceleration	18°/s² +18/-9%
Rotational Positioning Angular Velocity	18°/s ±25%



Note

The angular velocity is the average of the velocity of the tube arm rotating clockwise between 0° and 90° or rotating counterclockwise between 90° and 0° . The angular velocity does not include the time to accelerate from zero velocity and decelerate to zero velocity.

Source-to-Image Distance (SID)	70.0 cm ±1.0 cm (27.6 inches ±0.4 inches)
	(Focus position deviation is ±5 mm)
Patient Support (non-magnification)	
Vertical Position Lower Limit	70.5 cm +5.1/-0 cm (27.75 inches +2.0/-0 inches)
Vertical Position Upper Limit	141 cm +0/-17.8 cm (55.5 inches +0/-7.0 inches)

A.5.2 Compression

Manual Compression Force	Maximum of 300 N (67.4 pounds)
Motorized Compression	Functions in three operating modes:
	Pre-compression, Full-Range, Dual Compression.
	User selectable through software.
Pre-Compression Force	15 pounds to 30 pounds (67 to 134 N), motorized
Full Range Compression Force	20 pounds to 40 pounds (89 to 178 N), motorized
Dual Mode Compression	Provides Pre-Compression force upon first activation of compression switch; then, if switch is activated within 2 seconds, the force is increased incrementally for each additional switch activation, up to the user selected full compression force.
Compression Controls	<i>Up/Down controls on both sides of C-arm and on 2-position footswitch (Motorized). Handwheel on both sides of Compression Device (Manual).</i>
Compression Release	Manual Motorized Release controlled by push-buttons on both sides of the C-arm.
Automatic Compression Release	User selectable automatic release mode raises Compression Device upon exposure termination.
Down Motion Variable Speed	4.2 cm/s ±15% (1.66 inches/s ±15%)
Compression Force Display	Two LCDs on the Compression Device show the compression force through the range of 18 N to 300 N in 1 N increments (4 pounds to 67 pounds in 1 pound increments).
Compression Force Display Accuracy	±20 N (±4.5 pounds)
Compression Thickness Display	<i>Two LCDs on the Compression Device measure compression thickness in 0.1 cm increments. The display is visible from both sides of the patient.</i>
Compression Thickness Accuracy	± 0.5 cm (± 0.2 inches) for thicknesses between 0.5 cm and 15 cm (5.9 inches)
Breast Tomosynthesis Compression Thickness	Standard resolution tomosynthesis Maximum: 24 cm (restricted by compression device geometry)
	High resolution tomosynthesis Maximum: 15 cm (restricted by DICOM limitations)
Compression Paddles	Compression Paddles are transparent. The paddles are composed of polycarbonate resin or the equivalent. With compression applied, paddle deflection from a plane parallel to the patient support surface shall be less than or equal to 1.0 cm.

A.5.3 X-ray Tube

Focal Spot	Large (0.3 mm) Nominal
	Small (0.1 mm) Nominal
Tube Voltage	20 kV to 49 kV
Anode Material	Tungsten
X-Ray Window	Beryllium 0.63 mm
Tube leakage test conditions	49 kVp, 2.0 mA

A.5.4 X-ray Beam Filtration and Output

Filtration

Five-position filter wheel: Position 1: Rhodium, 0.050 mm ±10% Position 2: Aluminum, 0.70 mm (nominal) (Tomosynthesis option) Position 3: Silver, 0.050 mm ±10% Position 4: Copper, 0.3mm Position 5: Lead (provided for servicing)

kV/mA Range

kV	LFS mA	SFS mA
20	100	30
21	110	30
22	110	30
23	120	30
24	130	30
25	130	40
26	140	40
27	150	40
28	160	40
29	160	40
30	170	50
31	180	50

Table 26: Maximum mA Setting as a Function of kV

Table 26: Maximum mA Setting as a Function of		
kV	LFS mA	SFS mA
32	190	50
33	200	50
34	200	50
35	200	50
36	190	50
37	180	50
38	180	50
39	180	50
40	170	
41	170	
42	160	
43	160	
44	150	
45	150	
46	150	
47	140	
48	140	
49	140	

mAs Steps (Table 1, default)

4, 5, 6, 7, 8, 9, 10, 12, 14, 16, 18, 20, 22, 25, 30, 32.5, 35, 37.5, 40, 42.5, 45, 47.5, 50, 52.5, 55, 57.5, 60, 62.5, 65, 67.5, 70, 75, 80, 85, 90, 95, 100, 120, 140, 160, 180, 200, 220, 240, 260, $280,\,300,\,320,\,340,\,360,\,380,\,400,\,420,\,440,\,460,\,480,\,500$

Attenuation of Carbon Fiber

Image Receptor	$< 0.3 \ mm \ Al$
Magnification Platform	< 0.3 mm Al

A.5.5 X-ray Collimation

Collimation Fields	7.0 cm x 8.5 cm
	10 cm x 10 cm
	15 cm x 15 cm
	18 cm x 24 cm
	18 cm x 29 cm (Tomosynthesis option)
	24 cm x 29 cm

A.5.6 Light Field Indication

Light Field to X Ray Congruency

A.5.7 X-ray Generator

Туре	Constant Potential High Frequency Inverter
Rating	7.0 kW, maximum (isowatt), 200 mA at 35 kV
Electrical Power Capacity	9.0 kW maximum
kV Range	20 kV to 49 kV in 1 kV increments
kV accuracy	±2%, over range 20-49 kVp
mAs Range	3.0 mAs to 500 mAs in Manual Mode mAs (8 mAs minimum in AEC Mode)
mAs Accuracy	$\pm (10\% + 0.2 mAs)$
mA Range	10 mA to 200 mA, Large Focal Spot
	10 mA to 50 mA, Small Focal Spot

Within 2% of SID

A.6 Imaging System Technical Information

A.6.1 Image Receptor

Fluid Ingress	No fluid from accidental spillage on the Image Receptor may seep inside.
Deflection	Does not exceed 1.0 mm at maximum compression.
Active Imaging Area	Not less than 23.3 cm by 28.5 cm (9.2 inches x 11.2 inches)
DQE Conventional Mammography	Not less than 50% at 0.2 lp/mm
	Not less than 15% at the Nyquist limit

DQE (Tomosynthesis option)	Not less than 30% at 0.2 lp/mm	
	Not less than 15% at the Nyquist limit	
Dynamic Range and Linearity	Detector Subsystem response is linear with linearity of 0.999 over a dynamic range of 400:1 in x-ray exposure.	
Uniformity	Detector Subsystem can correct pixel-to-pixel gain variations.	
	For conventional mammography procedures, the uniformity of flat field image response of the detector shall be no greater than 2% after gain calibration is applied over an exposure range of 0.5 mR to 200 mR.	

Appendix B System Messages and Alert Messages

B.1 Error Recovery and Troubleshooting

Most faults and alert messages are cleared without result to your workflow. Follow the instructions on the screen or fix the condition then clear the status from the taskbar. Some conditions require a system restart or indicate that more action is necessary (for example, to call Hologic Technical Support). This appendix describes the message categories and your actions to return the system to normal operation. If errors repeat, contact Hologic Technical Support.

B.2 Types of Messages

B.2.1 Fault Levels

Each Message has a particular set of the following characteristics:

- Aborts an exposure in progress (yes/no)
- Prevents an exposure from starting (yes/no)
- Displays a message to the user on the Acquisition Workstation (yes/no)
- May be reset by the user (yes/no)
- May be reset automatically by the system (yes/no)

Displayed Messages

All displayed messages will be shown in the user's selected language.

Any message which aborts an exposure or prevents an exposure from starting will always display a message directing the user's actions required to proceed.

Additional Message Information

Technical information about the message is available in the log file.

Some messages always show as a critical fault (a system restart is necessary). These messages result from a condition which prevents an exposure, and which cannot be reset by the user or the system.

B.2.2 System Messages

When the following system messages show, do the step shown in the User Action column to clear the message and allow the next exposure.

Table	27: System Messages

Icon	Message	User Action		
	Paddle is moving	No action needed.		
₽	Sending notice	No action needed.		
8	Invalid use of Magnification Stand	You selected a tomographic view with the Magnification Stand installed. Select a non-tomographic view. (Tomosynthesis option)		
00	Face shield is not secured	Fully extend or fully retract the Face Shield. (Tomosynthesis option)		
8	Invalid use of compression paddle	Remove the Magnification Stand or install the Magnification Paddle.		
	Paddle position does not match selected view	Shift the Paddle to the correct location for the selected view.		
<u>→4.5 cm</u>	Compression is less than 4.5 cm during calibration	Move the Compression Paddle higher than 4.5 cm to complete the calibration procedure.		
	FAST compression is engaged	Disengage FAST compression and install a paddle designated for this mode.		
	License is missing	A license is necessary to use this feature or function. (This message is for your information only. There are no user actions.)		
? 2	Invalid detector calibration	Install the Magnification Stand for Small Focal Spot calibration. Remove the Magnification Stand to do Large Focal Spot calibration.		

Icon	Message	User Action		
?**	Invalid geometry calibration	Repeat the geometry calibration before you try to take an exposure. (Tomosynthesis option)		
	Configuration file is missing	Applies to Service Personnel.		
	Waiting for Detector	No action needed.		
	System in Test Mode	Applies to Service Personnel.		
Ţ₽+	Tube needs to be manually positioned (move to 0 degrees)	Rotate the C-arm to 0 degrees.		
01	Tube needs to be manually positioned (move to -15 degrees)	Rotate the C-arm to -15 degrees.		
<u>i</u> Q	Tube needs to be manually positioned (move to 15 degrees)	Rotate the C-arm to +15 degrees.		
	The Emergency Stop switch has been engaged.	Turn the Emergency Off switch one-quarter turn to reset the switch.		
>0.5 cm	Compression too low for tomo reconstructions.	Move the Compression Paddle higher than 0.5 cm to take tomography exposures.		

Table 27: System Messages

B.3 UPS Messages

Note



The User Guide for the UPS is supplied with the system. Refer to the UPS *User Guide* for complete instructions.

The LCD in the UPS shows the power status.

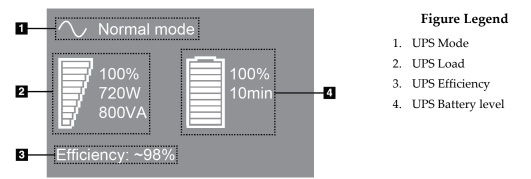


Figure 111: UPS LCD Display

If the UPS battery expires, the Mode icon changes as shown. Contact your service representative to replace the battery.



Appendix C Use of Mobile System

This appendix describes the system installed in a mobile environment.

C.1 Conditions for Safety and Other Precautions

An acceptable, stable, clean VAC power source is required to make sure that the system meets all its performance specifications. Where available, shore power correctly supplied to the system provides the best performance. If a mobile power generator is used, you must keep the specifications for input power during all load conditions.



Warning:

The radiation shield is not approved for mobile use and is not provided. The coach manufacturer must provide adequate shielding.



Caution:

When shore power is unavailable, mobile power sources that provide equivalent performance may be employed. (Refer to <u>Specifications for Mobile Use</u> on page 176.) Proper system function and performance can only be ensured if continuous true sinusoidal VAC power is supplied per the system power input specifications and loading characteristics. Intermittently, the power source must provide 65 Amps at 208 VAC for a minimum of 5 seconds, and 4 Amps maximum continuous otherwise. This load must be supported once every 30 seconds. In the event of shore or mobile power service interruption, the UPS must be capable of providing the operational power described above for a minimum of 4 minutes. Acquisition Workstation and Gantry power must be fed on separate dedicated circuits. The use of an uninterruptible power supply with active line conditioner is recommended on each power circuit. Accordingly, all ancillary mobile coach power should be distributed by other circuits. The electrical installation must be verified to meet system power input specifications and IEC 60601-1 safety requirements after initial installation and upon each relocation of the mobile coach.



Caution:

The temperature and humidity inside the vehicle must be maintained at all times. Do not allow environmental conditions to exceed stated specifications when the unit is not in use.



Caution:

Voltages cannot change by more than ±10% when the x-ray unit or other equipment (for example, heating or air conditioning) is operated.

Caution



To avoid image artifacts from occurring:

- Care should be exercised not to locate or park the mobile coach near sources of high power (such as power transmission lines and outdoor transformers).
- Make sure that any mobile power generator, uninterruptible power system (UPS), or voltage stabilizer is at least 3 meters (10 feet) from the closest point of the image detector travel.

C.2 Specifications for Mobile Use

The following system specifications are for mobile use only. For all other specifications, refer to *Specifications* on page 161.

C.2.1 Shock and Vibration Limits

Vibration Limit	Maximum of 0.30 G (2 Hz to 200 Hz), measured at the point where the system mounts to the coach.
Shock Limit	Maximum of 1.0 G (1/2 sine pulse), measured at the point where the system mounts to the coach. An "air ride" coach suspension is recommended.

C.2.2 Coach Environment

Operation Environment

Temperature Range	20 °C (68 °F) to 30 °C (86 °F)
Relative Humidity Range	20% to 80% without condensing moisture

Non-operating/Transit Environment

Temperature Range	10 °C (50 °F) to 35 °C (95 °F) for a maximum of 12 hours
	10 °C (50 °F) to 30 °C (86 °F) indefinitely
Maximum Rate of Temperature Change	<10 °C/hr.
Relative Humidity Range	10% to 80% without condensing moisture

C.3 Electrical Input

C.3.1 Gantry

Mains Voltage	200/209/220/230/ 240 VAC ±10%		
Mains Impedance	Maximum line impedance not to exceed		
	0.20 ohms for 208/220/230/240 VAC,		
	0.16 ohms for 200 VAC		
Mains Frequency	50/60 Hz ±5%		
Average Current over 24 Hours	< 5 A		
Peak Line Current	4 A (65 A maximum for 3 seconds)		

C.3.2 Acquisition Workstation

Mains Voltage	100/120/200/ 208/220/230/ 240 VAC ±10%
Mains Frequency	50/60 Hz ±5%
Power Consumption	< 1000 watts

C.4 Prepare the System for Travel

Before travel, perform these steps:

- 1. Rotate the C-arm to 0 degrees (CC position).
- 2. Lower the C-arm to its lowest position.
- 3. Turn off the system through the user interface.
- 4. Place the mouse in the keyboard tray.

- 5. Lock the keyboard tray (see the following figures):
 - a. Close the tray.
 - b. Find the lock knob under the tray.

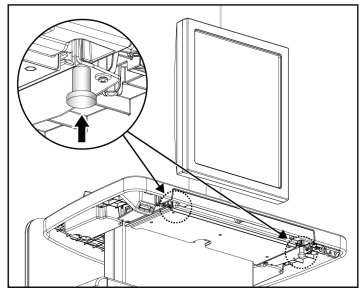


Figure 112: Keyboard Tray Lock Knob (Right or Left Side)

c. Turn the lock knob 90° until the knob fits into the lock. Position A in the following figure shows the locked position.

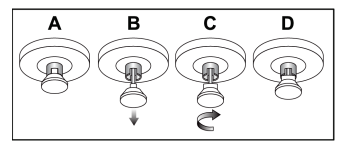


Figure 113: Tray Lock Release from Locked (A) to Unlocked (D)

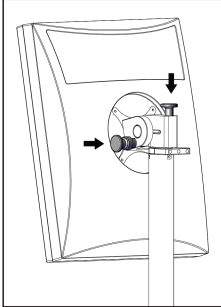


Figure 114: Swivel Monitor Lock Knobs on the Mobile Universal Acquisition Workstation (series I)

Figure 115: Swivel Monitor Lock Knobs on the Mobile Universal Acquisition Workstation (series II)

- 7. Lower the work surface to the minimum height.
- 8. Remove all system accessories.
- 9. Put all accessories in a safe storage area.

C.5 Prepare the System for Use

- 1. Unlock the keyboard tray:
 - a. Find the lock knob under the tray.
 - b. Pull the knob down.
 - c. Turn the knob 90°. This position keeps the latch open. Position D (in the following figure) shows the unlocked position.

6. Lock the swivel monitor using the knobs provided (see the following figures).

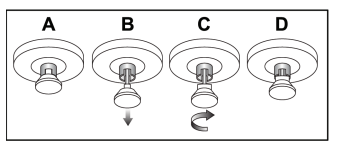


Figure 116: Tray Lock Release from Locked (A) to Unlocked (D)

- 2. Pull the tray out, if needed.
- 3. If using the mobile Universal Acquisition Workstation, unlock the swivel monitor (see the following figures).

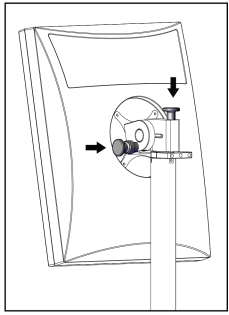


Figure 117: Swivel Monitor Lock Knobs on the Mobile Universal Acquisition Workstation (series I)

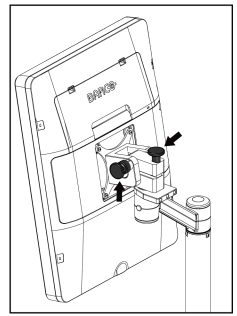


Figure 118: Swivel Monitor Lock Knobs on the Mobile Universal Acquisition Workstation (series II)

C.6 Test the System after Travel

C.6.1 Mobile System Controls and Functional Tests

To test the mechanical integrity of a mobile system, perform the Controls and Functional tests. Refer to *Perform the Functional Tests* on page 41.

- Compression Up/Down
- Compression Release
- C-arm Rotation
- C-arm Up/Down
- Collimator Override
- Light Field Lamp
- Shifting Paddle System
- Emergency Off Switches

Do the Controls and Functional tests each time the mobile system is relocated.

C.6.2 Quality Control for Mobile Systems

To verify the performance integrity of a mobile system, perform the following quality control tests:

- Artifact Evaluation
- SNR/CNR Measurement
- Phantom Image Evaluation
- Compression Thickness

Do the quality control tests each time the mobile system is relocated.

Appendix D Dose Information

D.1 EUREF Dose Tables



Notes

This information is only applicable for the European Union. The following values are for the default dose tables.

The following tables show typical dose values when operating the system in 2D and in BT imaging modes. All dose values have a tolerance of ±30%. The tables follow the procedures given in the *European guidelines for quality assurance in breast cancer screening and diagnosis, Fourth edition*: section **2a.2.5.1 Dosimetry**, and **Appendix 5: Procedure for determination of average glandular dose**.

Phantom	cm	kV	Anode	Filter	EUREF dose (mGy)
2.0 cm PMMA	2.1	25	W	0.05 mm Rh	0.55
3.0 cm PMMA	3.2	26	W	0.05 mm Rh	0.75
4.0 cm PMMA	4.5	28	W	0.05 mm Rh	1.05
4.5 cm PMMA	5.3	29	W	0.05 mm Rh	1.42
5.0 cm PMMA	6	31	W	0.05 mm Rh	2
6.0 cm PMMA	7.5	31	W	0.05 mm Ag	2.7
7.0 cm PMMA	9	34	W	0.05 mm Ag	3.1

Table 28: 2D Dose (EUREF)

Table 29: BT Dose (EUREF)

Phantom	cm	kV	Anode	Filter	EUREF dose (mGy)
2.0 cm PMMA	2.1	26	W	0.7 mm Al	1
3.0 cm PMMA	3.2	28	W	0.7 mm Al	1.15
4.0 cm PMMA	4.5	30	W	0.7 mm Al	1.5
4.5 cm PMMA	5.3	31	W	0.7 mm Al	2.00
5.0 cm PMMA	6	33	W	0.7 mm Al	2.5
6.0 cm PMMA	7.5	36	W	0.7 mm Al	3.9
7.0 cm PMMA	9	42	W	0.7 mm Al	5.15

		1			
Phantom	cm	kV	Anode	Filter	EUREF dose (mGy)
2.0 cm PMMA	2.1	26/45	W	0.05/0.3 mm Rh/Cu	0.83
3.0 cm PMMA	3.2	26/45	W	0.05/0.3 mm Rh/Cu	1.1
4.0 cm PMMA	4.5	28/45	W	0.05/0.3 mm Rh/Cu	1.6
4.5 cm PMMA	5.3	29/49	W	0.05/0.3 mm Rh/Cu	2.1
5.0 cm PMMA	6	31/49	W	0.05/0.3 mm Rh/Cu	3.0
6.0 cm PMMA	7.5	32/49	W	0.05/0.3 mm Ag/Cu	4.1
7.0 cm PMMA	9	33/49	W	0.05/0.3 mm Ag/Cu	4.7

Table 30: CEDM Dose (EUREF)

D.2 EUREF CNR Table



Notes

This information is only applicable for the European Union. The following values are for the default dose tables.

The following table shows typical CNR values when operating the system with Hologic Clarity HD[®] or 15-projection standard resolution BT imaging mode. All CNR values have a tolerance of ±25%. The data in the table was acquired by following the procedure given in the *European Protocol for the Quality Control of the Physical and Technical Aspects of Digital Breast Tomosynthesis Systems*: section **2.4 AEC Performance**.

Phantom	Hologic Clarity HD	Standard Resolution
2.0 cm PMMA	3.2	6.0
3.0 cm PMMA	2.5	4.7
4.0 cm PMMA	2.1	4.0
4.5 cm PMMA	2.1	4.0
5.0 cm PMMA	2.0	3.7
6.0 cm PMMA	1.9	3.5
7.0 cm PMMA	1.6	2.9

Appendix E List of Licenses and Accessories

The system is compatible with the following licenses and accessories:

	Description
1	C-View software
2	Genius AI Detection software
3	Hologic Clarity HD imaging technology
4	Intelligent 2D imaging technology
5	3DQuorum imaging technology
6	I-View software
7	SmartCurve breast stabilization system
8	Advanced Workflow Manager
9	Affirm [®] breast biopsy guidance system
10	Routine Screening Paddles
11	Contact and Spot Compression Paddles
12	Magnification Paddles
13	Localization Paddles
14	Ultrasound Paddle
15	Patient Face Shield
16	Magnification Stand
17	Localization Crosshair Device
18	Magnification Crosshair Device
19	Acquisition Workstations
20	MammoPad Breast Cushion

Table 31: Licenses and Accessories

Glossary of Terms

ACR

American College of Radiology

AEC

Automatic Exposure Control

Annotations

Graphic or text marks on an image to show an area of interest

вт

Breast Tomosynthesis. An imaging procedure that provides information about the breast in three dimensions

CEDM

Contrast Enhanced Digital Mammography

Collimator

A device at the x-ray tube to control the x-ray beam exposure area

Conventional Mammography

Single projection x-ray images of views for screening and diagnostic purposes

C-View

A licensed Hologic feature where a digital mammography (DM) image is generated from data acquired during a breast tomosynthesis (BT) scan

DBT

Digital Breast Tomosynthesis

DICOM

Digital Imaging and Communications in Medicine

DM

Digital Mammography (2D)

EMC

Electromagnetic Compatibility

FAST Paddle

Fully Automatic Self-adjusting Tilt Paddle

FDA

Food and Drug Administration (in the United States)

Grid

An element within the Digital Image Receptor that reduces scatter radiation during the exposure

Image Receptor

Assembly of the x-ray detector, x-ray scatter reduction grid, and carbon fiber cover

Intelligent 2D

A licensed Hologic feature where a highresolution digital mammography (DM) image is generated from data acquired during a high resolution breast tomosynthesis (BT) scan

I-View

A licensed feature for 2D Contrast Enhanced Digital Mammography

MPPS

Modality Performed Procedure Step

MQSA

Mammography Quality Standards Act

Notice

Annotations and comments per image communicated between Diagnostic Review Workstations, Technologist Workstations, and Acquisition Workstations

PACS

Picture Archiving and Communications System. A computer and network system that transmits and archives digital medical images

Pend

The action taken on an image to mark the image if the Technologist is not positive about the image quality (pended images must be Accepted or Rejected before the procedure is closed)

Projection Image

One of a group of breast tomosynthesis images taken at different projection angles and used to produce the final reconstructed image

Reclamation

Automatic removal of patient images and related information to allow storage of new patient image acquisitions

RF

Radio Frequency

ROI

Region of Interest

SID

Source to Image Distance

Tomosynthesis

An imaging procedure that combines a number of breast images taken at different angles. The tomosynthesis images can be reconstructed to show focal planes (slices) within the breast.

UDI

A United States Food and Drug Administration program for Unique Device Identification (UDI). For more information about UDI, go to <u>http://www.fda.gov/MedicalDevices/DeviceRegulationa</u> <u>ndGuidance/UniqueDeviceIdentification/UDIBasics/de</u> fault.htm.

UPS

Uninterruptible Power Supply

USB

Universal Serial Bus

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