

Instructions for Use

Selenia

Digital Mammography System



MAN-02620



Digital Mammography System

Instructions for Use

For Software Version 3.4

Part Number MAN-02620

Revision 004

January 2013

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Preface

The Selenia[®], based on the M-IVTM Mammography X-ray System, uses an Image Acquisition system that includes a digital image receptor. This receptor, which covers an area of 24 cm x 29 cm, is a direct-capture detector using an amorphous selenium photoconductor. At the Acquisition Workstation the user selects x-ray exposure technique factors, adds patient identification data, acquires, processes, and displays the digital images. The images are then either processed for printing and transmitted to peripheral hard copy laser film printer or sent to a diagnostic review workstation or both. Contrast and brightness are set automatically and can be adjusted by the user prior to printing or viewing. Hard-copy printers and a diagnostic review workstation are available as options.

The patient is prepared for the procedure in the same manner as for a film-based system. The Acquisition Workstation provides direct digital acquisition system advantages in system efficiency and productivity including:

- Patient demographics are associated with the electronic image.
- Technologist comments can be associated with the image.
- The technologist is assisted through the steps of image acquisition.
- Possible image destinations are provided via configurable settings.
- The technologist may review past acquisitions (including priors), to see previous captures, and then resend, and/or remark them.

1.0 Intended Uses

 $R_{x^{Only}}$ United States federal law restricts this device to use by, or on the order of, a physician.

The Selenia is intended to produce radiographic images of the breast. Its intended use is for the production, storage, and diagnostic review of digital screening, diagnostic, and needle localization mammography.

The system is to be used in a radiology or clinic exam room environment in a hospital, outpatient clinic, or a breast imaging center. Mammography technologists operate the system for the production and storage of digital mammograms. The system may also be used for quality control purposes and other clinical or research related activities by medical physicists and radiologists certified in accordance with MQSA standards.

- The Selenia Acquisition Workstation display is not approved for final interpretation of examinations. Final interpretations should be done from either films or with the diagnostic review workstation. Images shown on the Acquisition Workstation display are for quality assurance or confirmation purposes only.
- Only images produced by recommended laser printers, or an approved diagnostic review workstation, should be used for final interpretation of examinations. For compatible printers, see the latest product data sheets for the system, which can be obtained from Hologic® or your sales representative.



1.1 Indications For Use

The Selenia Full Field Digital Mammography System generates digital mammographic images that can be used for screening and diagnosis of breast cancer. The Selenia Full Field Digital Mammography System is intended for use in the same clinical applications as traditional screen-film mammographic systems. Mammographic images can be interpreted on either hard copy film or the diagnostic review workstation.

1.2 Contraindications

There are no known contraindications.

1.3 Potential Adverse Effects

The following is a list of potential adverse effects that apply to mammography and are also applicable to digital mammography using the Selenia.

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

2.0 Summary of Clinical Studies

This information is available in the document MAN-02233.

3.0 Product Complaints

Report any complaints or problem 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 Hologic. (See the title page for contact information.)

4.0 Technical Support

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

5.0 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.



6.0 Quality Control Requirements

The 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.

The 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.

7.0 Installation Instructions

Installation instructions are available in the Service Manual.

8.0 User Profiles

8.1 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.

8.2 Radiologist

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

8.3 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.



9.0 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, and facility on the job training also known as peer training. Additionally, the user's instruction manual is a reference for directions on how to use the system.

Your Hologic representative can arrange for training by a clinical services specialist.

All users must ensure that they receive training on proper use of the system prior to use on patients.

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

Accession Number	A DICOM term that refers to a RIS-created number that
	uniquely identifies a visit to a site by a patient.
ACR	American College of Radiology
Ag	Silver
AEC	Automatic Exposure Control. A method that limits the amount
	of radiation a patient receives.
collimator	Device at the x-ray tube that limits the area of the receptor that is exposed.
dialog box	A pop-up window that requires a user decision and a mouse or trackball click before another activity can occur.
DICOM	Digital Imaging and Communications in Medicine. An industry standard specification for communication between medical imaging equipment.
digital marker	A software mechanism that marks an image to show some information (normally orientation).
FAST Paddle™	Fully Automated Self-Adjusting Tilt Paddle system
Grid	Element within the Digital Image Receptor that reduces the scatter radiation during exposure
HIS/RIS	Hospital Information System/Radiology Information System. Generic term for non-PACS systems which track the Patient demographics and ordered radiological studies
HTC TM	High Transmission Cellular Grid
kV	Kilovolts: One of the x-ray exposure settings.

10.0 Terms and Definitions

Selenia
Digital Mammography System

login/logout	The process of logging into and out of the Operating System of
	the Acquisition Workstation
LUT	Look-Up Table (LUT). An image processing function that replaces one image pixel value with a different image pixel
	value.
Mag	Magnification
mA	Milliamperes. One of the x-ray exposure settings.
mAs	Milliampere-seconds. An electrical term used in x-ray exposure settings
Мо	Molybdenum
Modality Worklist (MWL)	A list of scheduled procedures normally kept by a RIS or PACS.
MPPS	Modality Performed Procedure Step. A DICOM service to allow the RIS (or another device) to know about work performed on the AWS.
MQSA	Mammography Quality Standards Act
Operating System (OS)	The software control system which runs all functions of a computer.
Outputs	A list of devices to which the accepted image is sent. The
1	devices can be a combination of archives, workstations and film
	printers.
PACS	Picture Archiving and Communications System. A computer
	and network system for the transfer and archive of digital
	medical images
PPS Status	The condition of a Performed Procedure Step being "in
	progress", "completed", or "discontinued".
Procedure	A generic medical protocol which contains a set of images
	(Views) which are acquired under certain conditions, and are
	performed together for a one purpose (for example standard
	screening). There is no Procedure instance UID item because a
	Procedure is a generic item in DICOM. DICOM supports the
	identification of requested Procedures.
Rh	Rhodium
ROI	Region of Interest
RIS	Radiology Information System
Series	A set of images acquired by a single tech for a single Patient and
	Procedure on a particular modality with a fixed body part,
	laterality and view position. DICOM uniquely identifies the
	series with a globally unique instance UID. Based on this
	description, each individually acquired DR image on the
	Acquisition Workstation is also an individual series.
Sign-in	The process of user identification to the Acquisition
	Workstation application.



Sign-out	The process in which a user exits the Acquisition Workstation application, but the user does not logout of the OS.	
0		
Smart Paddle™	A paddle with a release knob on each side which allows the	
	paddle to move from one side of the image detector to the othe	
	The system identifies a Smart Paddle installed on the	
	compression device.	
Solaris™	The company Sun [™] Microsystems' version of the UNIX	
	Operating System.	
TEC	Tissue Exposure Control mode, an enhanced Manual Exposure	
	Control (MEC) mode.	
TechMate TM	The Hologic SecurViewRT technologist review workstation	
	accessory, as packaged with the Selenia to reduce space in the	
	examination room.	
technique	Combination of x-ray parameters (kV, mA, etc.) for a specified	
	view in a procedure.	
UPS	Uninterruptible Power Supply.	
View	The combination of a single x-ray image and a specified set of	
	conditions under which the image was acquired. The View is	
	not part of DICOM nomenclature, but in the context of DR, is	
	approximately synonymous with a DICOM image object.	
W	Tungsten	
Wildcard Character	A keyboard character that represents one or many characters,	
	for example the * or ?.	



11.0 International Symbols

This section describes the International Symbols on this system.

×	Type B Applied Part
	Protective Earth terminal
0	"OFF" (power)
	"ON" (power)
Ō	"OFF" for part of the equipment
⊙	"ON" for part of the equipment
\bigcirc	Power Standby
	Discard electrical and electronic equipment separately from standard waste. Send decommissioned material to Hologic or contact your service representative.
4	Dangerous Voltage
•	Manufacturer
М	Date of Manufacture
	Caution—Radiation



12.0 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.	
<u>,</u>	Caution:	The procedures that you must follow accurately to prevent the damage to equipment, loss of data, or damage to files in software applications.	
	Note	Notes indicate additional information.	

Chapter 1 General Information

Digital Mammography System

1.0 Warnings and Precautions

Read and understand this manual before you use the system. *Always* follow all the instructions in this manual.

This system is for use by qualified Operators only. The Operators must have experience in the principles of mammography before following the instructions contained in this manual.

¥

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!	To avoid the risk of electric shock, this equipment must only be connected to a supply mains with protective earth.
WARNING!	Only trained service engineers authorized by Hologic can open any of the panels. This system contains lethal voltages.
WARNING!	Electrical equipment used near flammable anesthetics can cause an explosion.
WARNING!	After power failure, remove the patient from the system before you apply power.
WARNING!	To keep the isolation quality for the system, attach only approved accessories or options to the system. Only the authorized personnel can make changes to the connections.

Selenia Instructions for Use

Chapter 1—General Information Warnings and Precautions



	WARNING!	Keep a 1.5 meter safe distance between the patient and any non-patient devices.
		Non-patient system components (like the
		Workflow Manager, the diagnostic review
		workstation, or the hard copy printer) must not
		be installed in the Patient Area.
	WARNINCI	Per North American electrical safety
	WARNING:	requirements, you must use a Hospital Grade
		receptacle to provide a correct Ground.
		The user or a service engineer must correct problems
<u>·</u>	warning:	before the system is used.
	Marnina	The user must arrange for preventive maintenance by
	warning:	an authorized Service Engineer.
	Marnina	This system can be dangerous to the patient and the
	warning:	user. Always follow the safety precautions for x-ray
		exposures.
	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:	Only qualified users can use this system.
2		
	Warning:	Control the access to the equipment according to
	5	local regulations for radiation protection.
2		
	Warning:	Keep your full body behind the radiation shield
	0	during the exposure.



Warning:	This device contains dangerous material. Send decommissioned material to Hologic or contact your service representative.
Warning:	The equipment has motors. You must be careful when you adjust the equipment for patient use. Observe equipment and patient at all times during setup. If a chair is necessary, use an adjustable chair set above its minimum height.
Warning:	Put both footswitches away from the patient and C-Arm area to prevent any accidental footswitch use. When the patient has a wheelchair, put the footswitches away from the area.
Warning:	Never leave the patient during the procedure if in contact with the system.
Warning:	Keep the hands of the patient away from all buttons and switches at all times.
Warning:	You increase the patient dose to high levels if you increase the AEC exposure adjustment setting. You increase the image noise or decrease image quality if you decrease the AEC exposure adjustment setting.
Warning:	If a paddle touches possible infectious materials, contact your Infection Control Representative for decontamination instructions.
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.

Digital Mammography System

Selenia Instructions for Use

Chapter 1—General Information Warnings and Precautions



v v	Varning:	The Face Shield does not protect from radiation.
	aution:	Do not use any heat source (like a heating pad) on the image receptor.
<u> </u>	aution:	Never turn off the Acquisition Workstation or Value Console Circuit Breaker except in emergency. The circuit breaker can turn off the Uninterruptible Power Supply (UPS) and risk data loss.
<u> </u>	aution:	To minimize possible damage from thermal shock to the Digital Image Receptor, follow the recommended procedure to turn off the equipment.
<u> </u>	aution:	Do not put any magnetic media near or on devices that create any magnetic fields, because stored data can be lost.
<u>i</u> c	aution:	The system is a medical device and not a normal computer. Do not make changes to the hardware or software that are not authorized. Install this device behind a firewall for network security. The computer virus protection or network security for this medical device is not provided (for example, a computer firewall). The network security and anti-virus provisions are the responsibility of the user.
<u>i</u> c	aution:	To prevent errors and possible data loss, only use approved accessories with this equipment.



2.0 Radiation Safety

2.1 Exposure Duration

Table 1: Factors That Limit Exposure Duration

Normal Factors	Abnormal Factors	
Manual mAs Timer	Release of the X-ray button before the exposure ends.	
	•	Exposure longer than the preset "back-up time"
	•	The independent safety hardware back-up timer.
	•	Detection of a generator fault

2.2 Operator Radiation Shield

The radiation shield meets the requirements of 21CFR 1020 and IEC 60601-2-45.2001. The shield has a rating of a 0.5 mm Pb (lead) equivalent and is permanently installed.



Verify the shield integrity every day before use.

2.3 Patient Face Shield

Note

The Patient Face Shield keeps the face of the patient out of the radiation field.

2.4 Exam Room Door Indicators

- The system provides for an external Power-On indicator.
- The system provides for an external X-ray Exposure In Progress indicator.

3.0 Mechanical Safety

The equipment meets the requirements of IEC 60601-1, UL 60601, and CSA 22.2 No. 601.1 by these safety features:

- The C-arm rotation braking is ensured upon loss of power.
- Compression release is disabled when a localization paddle is installed.
- Motorized compression force has a limit of 200 N (45 lb).

4.0 Emergency Off Switches

The Emergency Off switches remove the power from the Gantry. Do not normally use the Emergency Off switches to turn off the system. See *Emergency Off Switches* on page 13.



5.0 Interlocks

- Display of 58 Newtons (13 lb) or greater of compression force disables the C-arm vertical drive and rotation.
- Installation of a Localization Paddle disables the automatic compression release functions.
- Release of an **X-ray** button before the end of the exposure ends the exposure.
- Misalignment of the Light Field Mirror or the Filter prevents an x-ray exposure.
- Installation of an 18 x 24 cm shifting paddle prevents the x-ray exposure if the detected paddle position does not match the View.

6.0 Compliance

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

6.1 Compliance Requirements

The manufacturer is responsible for the effects of safety, reliability, and performance of this equipment with the following provisions:

- The electrical installation of the room complies with the appropriate requirements.
- The equipment is used in accordance with instructions for use.
- Assembly operations, extensions, re-adjustments, modifications, or repairs are performed by authorized persons only.
- The installed network and communications equipment must comply with an IEC Standard, and the complete system (network/communications equipment and Selenia Mammography System) must be installed to comply with IEC 60601-1 and IEC 60601-1-1.



Selenia Instructions for Use Chapter 1—General Information Compliance



	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, the ME Equipment or ME System should be observed to verify normal operation in the configuration in which it is used.
<u>,</u>	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.
<u> </u>	Caution:	Changes or modifications not expressly approved by Hologic could void your authority to operate the equipment.

6.2 Compliance Statements

The manufacturer states that this device is manufactured/conforms to:

- EN ISO 13485:2007
- FDA, 21 CFR [Parts 820, 900 and 1020]
- IEC 60601-1:1988 +A1+A2:1995 +A13:1996 Medical Electrical Equipment—General requirements for safety
- IEC 60601-1-1:2000-12 Medical Electrical Equipment—Collateral Standard: Safety requirements for medical electrical systems
- IEC 60601-1-2:2001 Medical Electrical Equipment—Collateral Standard: Electromagnetic compatibility for medical electric systems
- IEC 60601-1-3:1994 Medical Electrical Equipment—Collateral Standard: Requirements for radiation protection in diagnostic x-ray equipment
- IEC 60601-1-4:1996 +A1:1999 Medical Electrical Equipment—Collateral Standard: Programmable electrical medical systems
- IEC 60601-1-6 Medical Electrical Equipment Collateral Standard: Usability
- IEC 60601-2-28:1993-03 Medical Electrical Equipment—Particular requirements for the safety of x-ray source assemblies and x-ray tube assemblies for medical diagnosis
- IEC 60601-2-32:1994 Medical Electrical Equipment—Particular requirements for the safety of associated equipment of x-ray equipment
- IEC 60601-2-45:2001 Medical Electrical Equipment—Particular requirements for the safety of mammographic x-ray equipment and mammographic stereotactic devices
- IEC 62304:2006 Medical Device Software Software life cycle processes



- UL 60601-1: Medical Electrical Equipment, Part 1—General Requirements for Safety
- CSA: Medical Electrical Equipment Part 1: C22.2 No. 601.1–M90–General Requirements for Safety
- ISO 14971

7.0 Label Locations



Figure 1: Selenia Label Locations







Figure 2: Selenia Value Console Label Locations



Chapter 2 System Components and Controls

1.0 System Description

The Selenia is available with an Acquisition Workstation or a Value Console. The Acquisition Workstation can also include a technologist review workstation accessory (the Hologic SecurViewRT TechMateTM) packaged with the Selenia to reduce space requirements.



Figure 3: Selenia System Description

Figure Legend

1. 2.	Selenia Acquisition Workstation. Value Console	Contain the image processing electronics and provide the user interface.
3.	SecurViewRT TechMate	Provides a Technologist review workstation
		accessory.
4.	C-arm	The C-arm provides a platform for the x-ray
		tube, compression systems, and the digital
		image receptor. A pivot mechanism connects it
		to the Gantry.
5.	Gantry	The Gantry contains the electrical and
		mechanical subsystems for the Selenia. C-arm
		and Gantry comprise the Tubestand.
6.	Dual-function Footswitches	Allow hands-free C-arm vertical travel and
		compression movements.



2.0 Film Printer

Selenia accepts film printers. Printed films display the patient information, exposure techniques, projection, and facility information.

3.0 System Power Controls

3.1 Input Power Circuit Breakers

Location of Circuit Breaker	Description	
II	 At the lower right corner of the back panel Removes the power for service or in an emergency Up is turned On. Down is turned Off. When turned off, a system restart is required. Restart the system following <i>Procedure for Startup</i> on page 23. 	
Figure 5: Acquisition Workstation Circuit Breaker	 On the back of the Acquisition Workstation behind the shield. The Value Console Circuit Breaker (not shown) is on the back of the Value Console. Use only in an emergency. If the circuit breaker was used, wait until the beeps stop, then turn on the circuit breaker. The system automatically restarts. If the UPS does not respond automatically, reset the UPS. See <i>How to Turn On or Reset the Uninterruptible Power Supply (UPS)</i> on page 31. 	


Selenia Instructions for Use Chapter 2—System Components and Controls System Power Controls

3.2 Emergency Off Switches

- Press any Emergency Off Switch to remove Gantry power.
- Rotate the switch one-quarter turn to reset.
- Restart the system after any Emergency Off Switch is pressed. See the procedure in *Restart After the Emergency Off Switch was Activated* on page 30.



Figure 6: Emergency Off Switch



Figure 9: Gantry Emergency Off Switches

Figure 8: Value Console Emergency Off Switch

Selenia Instructions for Use Chapter 2—System Components and Controls Acquisition Workstation Controls and Display



4.0 Acquisition Workstation Controls and Display



4.1 Acquisition Workstation



4.2 Keypads

The Keypads provide **Power On**, **X-ray**, and **Compression Release** functions.



Figure 12: Keypad Controls

Figure Legend

- 1. **Power On** button
- 2. Power On indicator light
- 3. X-ray Indicator lights on both panels
- 4. **X-ray** buttons, one on each side of the keyboard. Press both at the same time to begin an x-ray exposure and hold until the tone stops.
- 5. Compression Release button



4.3 Keyboard and Trackball or Mouse



Figure 13: Selenia Keyboard

Figure Legend

- 1. Selenia Function keys
- 2. Trackball (Acquisition Workstation)
- 3. Future Use
- 4. Standard keyboard keys
- 5. Standard arrow keys
- 6. Standard number keypad
- 7. Mouse (Value Console)

4.4 Keyboard and Screen Controls

Use the trackball or the function keys to change Exposure techniques.



Figure 14: Selenia Function Keys on the Keyboard

Table 2: Seleni	a Function Keys
-----------------	-----------------

Key	Name	Function
Mode	Exposure	Changes AEC, TEC, and Manual exposure modes.
	Mode	
Filter	Filter	Changes the Filters.
		The Filter options depend on the x-ray tube. The Molybdenum
		system has Mo and Rh. The Tungsten system has Rh and Ag.
KV Î	kV Up	Increases kV.
kv↓	kV Down	Decreases kV.
mAst	mAs Up	Increases mAs.



$1 \mathcal{U} \mathcal{U} \mathcal{L}$

Key	Name	Function
mAs	mAs Down	Decreases mAs.
Grid In/out	Grid	Changes the Grid Position (In or Out).
Spot	Focal Spot	Changes the Focal Spot size (Small or Large).
Breast Dens.	Breast Density	Changes the TEC Breast Density settings.
Aec Pos	AEC position	Moves the AEC Sensor position.
New Entry	New Entry	Opens the New Patient Entry dialog box.
Close Exam	Close	Closes the Procedure. (Disabled when MPPS is installed).
	Procedure	

4.5 Bar Code Scanner

The Bar Code Scanner speeds the entry of data from records with bar codes.

4.6 The Selenia Display

Features	Display
 Totoku Display On-screen Display Adjustment dial (item 1) Power button (item 2). Green indicator lights when Display is turned on. Has the ability to tilt. 	Figure 15: Totoku Display



Features	Display
Features Barco Display • • Meets the DICOM requirements. • Press the recessed dial (on the right side) to display a menu. • Press and hold the recessed dial to turn the Display On/Off.	Display
Green indicator lights when Display is turned on.Has the ability to tilt.	Figure 16: Barco Display
• The Value Console uses only the Barco display.	1.8 101 2 2 Aprily

5.0 TechMate Controls and Displays



Figure Legend

- 1. TechMate Display
- 2. TechMate Keyboard
- 3. TechMate Trackball
- 4. **Power** Button for TechMate
- 5. TechMate UPS Reset

Figure 17: TechMate Controls and Indicators



Selenia Instructions for Use Chapter 2—System Components and Controls Tubestand Controls and Displays

6.0 Tubestand Controls and Displays



Figure 18: Tubestand

6.1 Gantry Controls and Displays

Gantry Controls

- 1. The C-arm Rotation Angle Displays on both sides of the Gantry show the C-arm angle.
- 2. The Emergency Off Switches on each side of the Gantry remove power from the Tubestand.
- 3. Gantry Rotation Switches (if installed) on each side of the Gantry move the C-arm to a programmed position.
- 4. The Input Power Circuit Breaker at the rear of the Gantry provides the overload protection.
- 5. Dual-Function Footswitches activate C-arm and Compression Device movement.





C-arm Controls and Displays 6.2 1. Tubehead Controls 1 2. Compression Device 3. Rear of the C-arm Side Rails 4. Rear of the Image Receptor 3 4 The Tubehead Controls on either side of the x-ray tube side covers provide the Collimator Override, C-arm Rotation, and Light Field functions. The Compression Device contains the AEC Sensor control and position LEDs, and displays compressed breast thickness and force. Handwheels allow manual compression control. The rear edges of the C-arm Side Rail have recessed push-button controls for motorized compression and C-arm movement. An additional C-arm Rotation button is available on the rear of the Image Receptor.



6.2.1 Compression Device Controls and Displays

Figure Legend

- 1. Compression Handwheels
- 2. AEC Sensor Position Controls
- 3. AEC Sensor Position Display
- 4. Compression Force Display
- 5. Compression Thickness Display



6.3 Dual Function Footswitches

- 1. Press the footswitch to activate.
- 2. Release the footswitch to stop motorized movement.

Figure Legend

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





Warning:

Put both footswitches away from the patient and C-Arm area to prevent any accidental footswitch use. When the patient has a wheelchair, put the footswitches away from the area.





1.0 Procedure for Startup

Digital Mammography System

Table 3: System Startup Procedures

Step	Description	
 Reset the Emergency Off Switches. Two on the Gantry One on the Acquisition Workstation (or Value Console) 	Figure 22: Emergency Off Switch	
2. Make sure that the circuit breakers are turned On.	Figure 23: Gantry Circuit BreakersFigure 24: Acquisition Workstation Circuit Breaker	
3. Perform Pre-Startup Checks.	 Look for open or loose panels, missing hardware, and indications of damage. Inspect the radiation shield for chips, cracks, breaks, 	
	and for tight attachments.	
	3. Inspect the paddles for small cracks.	
	4. Remove any obstructions to the Operator view and C-arm movement.	



Step	Description	
 4. Turn On the Selenia. Turn on the TechMate. (If the Acquisition Workstation or TechMate does not start, reset the UPS for that unit. See <i>How to Turn</i> On or Reset the Uninterruptible Power Supply (UPS) on page 31.) 	Image: state of the state	
	1. Press the Power On button (item 1). The green LED (item 2) illuminates.	
	2. Press the Power On button (item 3) and the TechMate turns on.	
	3. Allow the time for initialization and diagnostic tests to complete.	
5. Log on to the Operating System.	When the Logon dialog box appears:	
	 Enter your user name (case sensitive) for the Operating System. 	
	2. Select OK or press the Enter key.	
	3. Enter your Operating System password. (No characters appear in the field.)	
	4. Select OK or press the Enter key.	
6. Log in to the Application.	When the Login to System dialog box appears:	
	1. Select your ID from the drop-down list.	
	2. Enter your password (case sensitive). (Asterisks appear in the field.)	
	3. Select OK or press the Enter key.	

Table 3: System Startup Procedures



Note

Note

If the system remains on over the night, reboot the system daily to resync system communications.

If the system was turned off for 30 minutes or longer, the detector temperature needs time to adjust. Allow a minimum of 1 hour before you acquire images on a patient. The Launch dialog box disappears when the wait time is finished.

If it has been less than 30 minutes, click Dismiss in the Launch dialog box.





If during the internal checks, the system detects a fault condition, a message appears and startup is suspended until the problem is remedied.

2.0 Controls and Functional Tests

Perform these functional tests monthly.

Note

The C-arm movement is disabled when compression force of 58 N (13 lb.) or greater is applied.

Function	Control	Test
Counterclockwise C-arm Rotation	C-arm and Gantry Rotation Switches	 Press and hold the C-arm button and confirm the following actions: C-arm rotates Angle display changes Rotation stops when you release the button. Press and hold a Center button and confirm C-arm moves to zero degree position.



Function	Control	Test
Clockwise C-arm Rotation	C-arm and Gantry Rotation Switches	 Press the button and confirm the following actions: C-arm rotates Angle display changes Rotation stops when you release the button. Press and hold a Center button and confirm C-arm moves to zero degree position.
Compression Down		 Press the button. Confirm the compression brake engages and the light field lamp illuminates. Confirm the Compression Down movement stops: When you release the button. At the compression down force limit. At the lower compression travel limit







Function	Control	Test
		 When you release the button. When the C-arm reaches the lower travel limit.
Light Field Lamp		 Press the button. Confirm the light turns on for a short time. Confirm the light field lamp turns on when a Compression Down button is activated.
Collimator Override		 Press the light field button then the collimator button. Confirm the Collimator moves to the next field size. Repeat the steps 1 and 2 until you move through all Collimator field sizes.

Table 4: Monthly Control Function Tests



Function	Control	Test
Smart Paddle	TOCK SWIT	 Install a Smart Paddle. Rotate and hold the knob on the side of the paddle to move the paddle into a detent positions. Release the knob to lock the paddle in position. Turn on the light field lamp. Confirm the collimator position matches the paddle position. Repeat this procedure for the other two paddle positions. Apply compression and confirm the paddle does not unlock.
AEC Position	Aec Pos	 Select an AEC mode. Change the AEC position with the compression device arrow button. Confirm the indicator positions on the sides of the device and Acquisition Workstation display match the current position. Move past position 7 or 1. Confirm the Auto position indicator on the compression device illuminates and the display indicates Auto. Select the AEC arrows on the display and repeat steps 3 to 5. Use the keyboard function key and repeat steps 3 to 5.



3.0 Monthly System Tests

- Test all Emergency Stop Switches. Follow the on-screen reset procedure.
- Change the AEC Sensor Position switch, and make sure the compression device indicator and the Acquisition Workstation display change.
- Test all buttons on all keypads and footswitches. Include the compression release on the Acquisition Workstation.
- During Compression Device tests make sure the thickness and force displays change.
- During C-arm rotation tests, confirm that the LED Angle displays change.

4.0 How to Restart the System

4.1 Restart From Sleep Mode

1. To activate the system, move the trackball or mouse.



Figure 26: Operating System Login Screen

- 2. In the Login dialog box, enter reboot for the user name and password.
- 3. Wait for the system to restart.
- 4. Log on to the Operating System, then Log on to the Application software.
- 5. If the system was off for less than 30 minutes, select **Dismiss** in the **Launch** dialog box.

4.2 Restart After the Emergency Off Switch was Activated

- 1. Turn the Emergency Off switch by one-quarter turn clockwise to reset the switch.
- 2. When the Power Panel dialog box displays, "**Communication with the generator is lost**," examine the Emergency Off Switches and the circuit breaker:



Caution:

Do not close this dialog box by using the X in the upper corner.

- If the Gantry power is turned on, select the **Cancel** button.
- If the Gantry power is turned off, and the Emergency Off Switches and the circuit breaker are set correctly, select the **OK** button.
- 3. When the "Link established with generator" dialog box appears, select OK.



4.3 How to Turn On or Reset the Uninterruptible Power Supply (UPS)



Figure 27: The Acquisition Workstation UPS Switch Lever

- 1. Find the Acquisition Workstation UPS switch lever on the lower, right of the back panel.
- 2. Press the switch lever down and hold for two seconds or until the UPS beeps.
- 3. Start the Acquisition Workstation normally.



Figure 28: Value Console UPS Switch

- 1. Find the Value Console UPS switch, on the front of the UPS.
- 2. Press and hold the UPS switch for two seconds or until the UPS beeps.
- 3. Start the Console normally.

4.4 How to Turn Off the UPS

- 1. Make sure the Acquisition Workstation and TechMate or the Value Console are off. The UPS system begins to chirp repeatedly and can now be shut down.
- 2. Find the UPS switch. See the previous figures.
- 3. For the Acquisition Workstation, press and hold the switch lever down for two seconds or until the UPS beeps. For the Value Console, press and hold the UPS switch for two seconds or until the UPS beeps.
- 4. For the TechMate, perform steps 1 and 2 in UPS Operation for TechMate on page 32.
- 5. When the power is restored, press the UPS switch one time to turn on the UPS.



4.5 UPS Operation for TechMate

To reset the TechMate UPS:



Figure 29: Reset the TechMate UPS

- 1. Find the UPS access hole in the middle of the left side of the front cover. This hole aligns with the **Power** button on the UPS.
- 2. Put the eraser-end of a pencil into the hole to press the **UPS** button until the UPS beeps.
- 3. Press the button again and listen for a second beep.
- 4. Start the TechMate normally.

To turn off the TechMate UPS, follow steps 1 and 2.



5.0 How to Log Out

There are two methods to log out of the application. The system will remain on and another user can log in to use it.

	Method 2	
1. Select the Sign Out button. 1. Select Fill 2. Select Log	 Select File>Exit. Select Log out of the computer?. 	
Hologic Sign Out Figure 30: The Sign Out Button	 Exit from acq. station Do you want to: Power off the computer? Restart the computer? Log out of the computer? Sign in as a different user? 	
2. Select the Exit button.		
3. Select Yes.	Yes No	
3. Select Ye	26.	

Table 5: The Log Out Methods



6.0 How to Shut Down the System

- 1. Select File>Exit.
- 2. Select **Power off the computer?** from the Exit from acq. station dialog box. See the *figure Exit from Acquisition Workstation Dialog Box* on page 33.
- 3. Select Yes.

	Note	If a message appears and indicates that there are spool jobs, select log off or wait until the jobs are finished. (The spool continues again later.) Hologic recommends that you allow the jobs to complete before shut down.
	Caution:	Do not use a circuit breaker or Emergency Off switch as a routine method to turn off the Selenia.
<u>,</u>	Caution:	Never turn off the Acquisition Workstation or Value Console Circuit Breaker except in emergency. The circuit breaker can turn off the Uninterruptible Power Supply (UPS) and risk data loss.



Chapter 4 The Selenia Accessories

1.0 Introduction

Available accessories depend on your system configuration. Selenia performs screening and diagnostic applications. The Selenia S system performs only screening applications.

Accessory		Selenia	Selenia S
Routine Screening Paddles	18 x 24 cm	*	*
	18 x 24 cm FAST	*	*
	24 x 30 cm	*	*
	24 x 30 FAST	*	*
	Small Breast	*	*
Contact and Spot Compression	10 cm Contact	*	
Paddles	15 cm Contact	*	
	Ultrasound	*	
	7.5 cm Spot Contact	*	See Note
	Frameless Spot	*	
Magnification Paddles	7.5 cm Spot	*	
	10 cm	*	
	15 cm	*	
	10 cm Mag Localization	*	
	10 cm Mag Perforated Localization	*	
Localization Paddles	10 cm Perforated	*	
	15 cm Perforated	*	
	10 cm Rectangular Open	*	
	15 cm Rectangular Open	*	
Patient Face Shield		*	*

Table 6: Available Accessories



Table 6: Available Accessories

Accessory	Selenia	Selenia S
Magnification Platform	*	
Localization Crosshair Device	*	
Magnification Crosshair Device	*	



On the Selenia S system, only use the 7.5 cm Spot Contact Paddle for compression thickness calibration.

2.0 Patient Face Shield

Note

Remember to examine the Face Shield condition before use each day.

2.1 How to Install the Face Shield on the Tubehead

- 1. Put the tab ends on the open end of the Face Shield into the slots on the tubehead mount.
- 2. Slide the Face Shield into the tubehead mount until the Face Shield locks in position.

2.2 How to Remove the Face Shield

- 1. Pull the rear sides of the shield away from the tubehead.
- 2. Slide the shield off the mount.



Figure 32: Installation of the Face Shield



Warning:

The Face Shield does not protect from radiation.

3.0 Compression Paddles

There are 19 compression paddles for Screening and Diagnostic Procedures. 12 of the 19 paddles are Smart Paddles and have mechanisms that allow manual lateral movement (both left and right). The x-ray collimation moves with the Shifting Compression Paddle position. The other seven paddles include five magnification paddles and two large, 24 x 30 cm paddles.

3.1 How to Install the Compression Paddles

- 1. Align the mounting pins on the rear frame of the paddle with the mounting holes in the Compression Device.
- 2. Push the compression paddle into the Compression Device until the paddle stops.
- 3. Carefully pull the paddle out approximately 3 mm until the paddle clicks into position.

3.2 How to Remove a Compression Paddle

- 1. Hold the paddle by the metal sides of the frame.
- 2. Pull the paddle in a straight line from the mounting holes.



Figure 33: Mount the Compression Paddle



3.3 Maintenance and Cleaning

Clean the paddles after each use. See *Maintenance and Cleaning* on page 103 for cleaning instructions.

3.4 Smart Paddle System

A release knob on each side of the Smart Paddle allows it to move from one side of the image detector to the other. This enables the Technologist to optimize the breast position and imaging within the active area of the Image Receptor, especially when Lateral Views of a small breast are required.

To shift a Smart paddle:

- 1. Select a View on the Acquisition Workstation.
- 2. Rotate the release knob on the side of the paddle from the lock to the shift position.
- 3. While you hold the knob in the shift position, move the paddle to match the View. The paddle locks into position on a detent.
- 4. Release the knob to lock it in position.
- 5. Confirm the paddle is locked into position.
- 6. Activate the light field lamp and confirm the collimator matches the paddle position.



Figure 34: Shifting Compression Paddle

3.5 FAST Paddle Use

The Fully Automatic Self-adjusting Tilt (FAST) Paddle 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 paddle use with this type of breast provides these features:

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

The FAST paddle automatically tilts when the compression is applied. The paddle is in a flat position until a compression force of approximately 20 pounds (88 Newtons) is applied. The paddle then tilts until its maximum angle is reached at a force of approximately 30 pounds (132 Newtons).

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

3.6 How to Realign the Paddle Front Edge

If the front edge of the paddle with 30 lb (132 N) of compression force applied is not in alignment with the front edge of the Image Receptor, follow this procedure.

- 1. Release the compression and remove the paddle.
- 2. Loosen the hardware that holds the paddle to the frame. On a standard screening paddle, or a frameless spot paddle, turn the paddle upside down and loosen the hardware one complete turn. On a FAST paddle, use a hex wrench to loosen the paddle.
- 3. Turn the paddle over (if necessary) and loosen the two inside screws. See the *figure Alignment of Adjustment Screws* on page 39.
- 4. Install the paddle and apply 30 lb (132 N) of compression force.
- 5. Move the paddle to the correct position. (Do not release the compression.)
- Tighten both inside screws that you loosened in step 3.
- 7. Release the compression and remove the paddle.
- 8. Turn the standard screening or the frameless spot paddle upside down and tighten the hardware that fastens the plastic paddle to the frame.
- 9. Install the paddle and apply 30 lb (132 N) of compression force.
- 10. Confirm that the front edge of the paddle is aligned to the front edge of the Image Receptor.



Figure 35: Alignment of the Paddle



Figure 36: Alignment of Adjustment Screws

Digital Mammography System



Identification of Compression Paddles 3.7

Routine Screening Paddles 3.7.1



3.7.2





Ultrasound Paddle

15 cm Contact Paddle



7.5 cm Spot Contact Paddle Frameless Spot Paddle



C

Paddle

15 cm Magnification

3.7.3 Magnification Paddles



7.5 cm Spot Magnification Paddle



10 cm Magnification Localization Paddle

10 cm Magnification Paddle



10 cm Magnification Perforated Localization Paddle

Use the Magnification paddles when the Magnification Platform is installed.

3.7.4 Localization Paddles





10 cm Rectangular Open Localization Paddle



15 cm Perforated Localization Paddle

10 cm Perforated Localization Paddle



15 cm Rectangular Open Localization Paddle

3.7.5 About the Localization Paddles

The Installation method is the same for all Paddles. The system locks out Auto Compression Release when a Localization Paddle is installed. The AEC Sensor may require repositioning.



4.0 Magnification Platform

When the Magnification Platform is installed, the HTC grid automatically retracts and the default values for Magnification x-ray exposure techniques are set.

4.1 How to Install the Magnification Platform

- 1. Remove the Face Shield (see *Patient Face Shield* on page 36).
- 2. Remove the compression paddle.
- 3. Move the Compression Device above the slots in the C-arm.
- 4. Hold the Magnification Platform by the support brackets and align the hooks of the bracket with the mounting slots on the C-arm.
- 5. Put the hooks into the C-arm slots.
- 6. Push the frame down so that the clips lock the Platform in position.

When the platform is installed, you can use only the Magnification paddles. (See *Magnification Paddles* on page 41.)

4.2 How to Remove the Magnification Platform

- 1. Remove the Magnification paddle.
- 2. Press the locking clips on the Magnification Platform down to release the hooks.
- 3. Lift the Magnification Platform up and out by the support bracket.
- 4. Reinstall the Face Shield.



Figure 37: Magnification Platform Installation



5.0 The Localization Crosshair Device

The Localization Crosshair device, used with a localization paddle enables the location of a specified spot on the breast.

5.1 How to Install the Crosshair Device



Figure 38: Installation of the Crosshair Device

- 1. Remove the Face Shield from the tubehead.
- 2. Move the Compression Device below the C-arm mounting slots.
- 3. Hold the Crosshair device by the support brackets and slide the hooks into the C-arm slots.
- 4. Push the device down to the locked position.
- 5. Install the Localization paddle on the Compression Device.

5.2 How to Remove the Crosshair Device

- 1. Rotate the Crosshair device to the left or right.
- 2. Press the two locking levers inside each mounting arm.
- 3. Lift the device up and from the C-arm. Be careful. The device is top heavy.
- 4. Install the Face Shield.



Figure 39: Crosshair Locking Levers

Note



5.3 Localization Crosshair Device to the Localization Paddle Alignment



Before you perform the following adjustment, make sure the Localization paddle is aligned to the edge of the image receptor. See *How to Realign the Paddle Front Edge* on page 39.

- 1. Install a rectangular localization paddle.
- 2. Loosen the adjustment lock screw on the bottom of the Crosshair Device. See Number 1 in the image to the right.
- 3. Put a piece of white paper on the breast tray so that you can see the shadows of the crosshairs.
- 4. Move the Localization paddle approximately 6 cm above the image receptor.
- 5. Turn on the light field.
- 6. Move the Crosshair Device until the rectangle of light aligns with the opening in the localization paddle.
- 7. Tighten the adjustment screw.



Figure 40: Adjustment Lock Screw

6.0 The Magnification Crosshair Device



Figure 41: Installation of the Magnification Crosshair Device

- 1. Remove the face shield from the tubehead.
- 2. Align the Magnification Crosshair Device with the grooves of the face shield rails.
- 3. Push forward until the device locks into position.
- 4. To remove the Magnification Crosshair Device, pull and slide the assembly toward you.



Chapter 5 The User Interface

Digital Mammography System

1.0 Introduction to the User Interface

The process of image acquisition, enhancement, and output is done through three screens:

- Select a Patient screen
- Patient Views screen
- Preview Images screen

When you select an option on a screen, other screens may appear for information entry or selection of additional options. This manual covers the screens that appear while you are logged in as a "tech".

Select a Patient

• Select a Patient

Patient Views





Preview Image



2.0 A Screening Exam Sequence Suggestion

Table 7: Workflow Suggestions to Select a Patient and Acquire an Image

		Action	What it looks like
1.	Sta Sys Sta	rt up the system and log in to Operating stem and Application. See <i>Procedure for</i> <i>rtup</i> on page 23.	
			Figure Legend
			1. Power On button
			2. Power On indicator light
2. 5	Selec	t the patient.	
	•	Query the MWL	Worklist Local Exams
	a.	Select Worklist tab in upper left of screen.	Patient Name Local Exams Patient, One Search
	b.	Use the Bar Code Scanner to input query data (the procedure opens), or type the query data.	Terretoria Fastina (* 16 serch): Patient Cont Fastina Co
	c.	Select Query.	Patient
	d.	Select the patient from the list, then select the Enter key.	Name: DOB: ID:
	e.	Select the patient from the list.	Gender:
	•	Search the local database	New Eats Close
	a.	Select the Local Exams tab in the upper left of the screen.	
	b.	Select the Patient Name tab (or ID tab) then enter the data.	
	c.	Select the Search button. See <i>More</i> <i>Information about Searches</i> on page 53 for information about wildcard searches.	
	d.	Select the patient from the list.	

	Action	What it looks like
•	Add a New Patient	
a.	Select New from the Patient area of the screen.	
b.	Enter the required information.	
c.	Select the Accept button.	
3. Selec schedu •	et the Procedure, when more than one is lled. Select from the scheduled procedures list.	The Edit Admin Info
a.	Find the procedure in the list.	Downie da Permise bland Manangrady Bolokidal Procedure 12 DM Diagootis Manangrady Bolokidal Procedure
b.	Click the Select button.	Outputs
•	Add a Procedure	Procedure
a.	Select New in the Procedure area.	Procedure: Screening-Bilateral Mammog
b.	Select the procedure from the list.	Procedure #. AN08262003A
c.	Select the Accept button.	New Edit Add
4. Edit	the Patient Information, if necessary	citi Dalant
a.	Select the patient from the patient list.	Edit Patient
b.	Select Edit from the Patient area.	*First Name: Sandy
c.	Make the changes in the Edit Patient screen.	*Patient ID: 123-44-5678 *DOB: (MMDDYYYY) 01 412 *Age of Patient: 056Y
d.	Select the Accept button.	"NOTE Fields with an attentik '* are mandatory and require input Clear Cancel Accept
5. Insta Accesso and ali	Ill the accessories. See <i>The Selenia</i> pries on page 35 for installation, removal, gnment procedures for each device.	 Face Shield Compression Paddle Magnification Platform Crosshair

Table 7: Workflow Suggestions to Select a Patient and Acquire an Image



Action		What it looks like
6. Select the view.		
•	Use the default view sequence.	The tai Admis tais Patient, One Serving literal Varianzatio
a.	Acquire the selected view.	Md Varie Restance Varies Open Section Description x ccc Loc LMD Filled Section x ccc Loc LMD Filled Section
b.	Acquire an image for each of the remaining views.	
•	Select another view from the displayed views.	Comp Fores Mode FTLL Cold Mode AUTINALITY Comp Fores Mode FTLL Cold Mode AUTINALITY Comp Fores Mode Status Cold Mode AUTINALITY Cold M
a.	Select any view displayed on the Patient View screen.	
b.	Acquire the selected view.	Add View Remays Unexposed
с.	Repeat for the remaining views.	Add View
•	Add a view.	
a.	Click Add View in the Patient View screen.	
b.	In the dialog box, select a view from the list of Standard Mammography Views. To add many views, press and hold the Control key and select the views.	
с.	Select the OK button.	
7. Selec	ct the Exposure Techniques.	Coop Fores Mode RLLL Coll Mode: ALTOMATIC Coop Falces Mode MARTAL Coll Size 36:20m Coop Falces Mode MARTAL Coll Size 36:20m Diago Size 21:20m ALTOREY Mode 27 10.0 0 R Control
8. Posit	tion the system and Patient.	
a.	Adjust the vertical height of the Breast Platform.	
b.	Set the C-arm angle.	
C.	Put the arm and hand of the patient in a safe location.	
d.	Compress the breast.	

Table 7: Workflow Suggestions to Select a Patient and Acquire an Image
	Action	What it looks like
9. Acqu	uire the image.	1 0
a.	Make sure the message area of the screen displays "Ready" (item 1) and there is no "X" through the x-ray indicator (item 2).	Generator Statur Ready Ganarator Error
b.	Press and hold the x-ray buttons.	Based Generator
c.	Inspect the image on the Preview Screen then select the Accept or Reject button.	kVp: 26 organ: mGy mAs: 39.5 entrance: mGy 3.8 cm: 10 lbs. filter: Rh Reject Accept
10. Clo	se the procedure.	
a. b.	Select the Close Procedure button. Select the Yes button.	Close Procedure Andrea Sign Out Patient

Table 7: Workflow Suggestions to Select a Patient and Acquire an Image





Chapter 6 The Select a Patient Screen

1.0 Introduction

The Select a Patient screen appears when the system application loads. This screen allows you to locate a patient, and access non-routine Acquisition Workstation options with no patient selected.



Figure 42: The Select a Patient Screen

Figure Legend

- 1. Menu Bar Menu Bar on page 64
- 2. Search for a Patient *Select a Patient* on page 52
- 3. Patient Query *How to Search the Acquisition Workstation Database (Local Exams)* on page 52
- 4. Today's Worklist More Information about Searches on page 53
- 5. Exposure Techniques Exposure Information on page 59
- 6. Add/Edit the Patient Information How to Add a New Patient on page 53
- 7. Add/Edit a Procedure New, Edit, Add a Procedure on page 56
- 8. Outputs Image Output on page 58
- 9. Special Modes Special Modes Buttons on page 78
- 10. Generator Status Generator Status on page 78
- 11. System Message System Messages on page 78



2.0 Select a Patient

A patient can be added to the system or already exist and be selected from a list. There are two places to search for an existing patient, the worklist which is a list of scheduled patients from a Modality Worklist Provider and the Local Exams which are the patients on the computer in the workstation.

2.1 How to Query the Modality Worklist Provider

1. Select the **Worklist** tab in the upper left area of the Select a Patient screen.



If you changed the date from the 'today' setting, select the **Today's Worklist** tab.

- 2. Enter the query data.
 - If you use the Bar Code Scanner to input the query data, the procedure opens.
 - If you use the keyboard to type the query data, select the **Query** button, then select the patient from the list to open the procedure.



Note

Note

This query does not search the local Acquisition Workstation database, Local Exams. An Asterisk is not necessary for this search.

Worklist Local Exams Patient Query Todays Worklist

If your search for one match with a Patient ID or Accession Number has results, the patient Procedure opens automatically. Selecting a patient adds them to the Local Exams database.

2.2 How to Search the Acquisition Workstation Database (Local Exams)

- 1. Select the Local Exams tab in the upper, left area of the Select a Patient screen.
- 2. Select the **Patient Name** or **ID** tab.
- 3. Enter the Patient Name or ID in the Search box. Or, enter part of a Patient Name or part of an ID number and use an asterisk for the missing characters.
- 4. Select the **Search** button or press the **Enter** key.



2.3 More Information about Searches

• When part of the patient name is entered, the system applies a wildcard character at the end of the name.



Note

You can use the asterisk (*) as a wildcard character to increase your search. For example, when you enter R*ph, Patients with the names Randolph and Rudolph will display. If you use a wildcard character, make sure that you include a minimum of one other character.

- You can use one or more of the search fields.
- You can limit the query to a date range.
- Most worklist providers require information in the Patient ID and Accession Number fields to match. Data in these fields is case sensitive.
- You must include the caret (^) character between the name parts (for example, DOE^J).

3.0 How to Add a New Patient

The Patient area of the screen allows you to Add a New Patient.

1. Select New in the Patient pane to display the New Patient Entry dialog box.

Pa	New Patient Entry	
*Last Name:	5 1 11	
*First Name:		
Middle Name:		
*Patient ID:		
*DOB:(MMDDYYYY		
*Age of Patient:		
*Gender:	F	
Referring Physician:		
Patient Location:		
*Accession Number:	Support and a support of the support	
*Procedure Description:	Mammography / Screening-Bilateral Man	nmography /
NOTE: Fields with an asterisk '' are mandatory	and require input.	
Clear	Cancel	Accept

Figure 43: New Patient Entry Form

- 2. Enter the Patient demographic information. Use the Tab key or the trackball to move through the fields. You must enter information in the Fields that have an asterisk.
 - The **Patient Name** fields must contain only letters or numbers.
 - The **Clear** button clears the information you entered.
 - The **Cancel** button closes the form without your changes.
- 3. Select the **Accept** button to enter the Patient in the Acquisition Workstation Local Database. The Patient Views screen displays.



Warning:

Verify the Patient Demographic Application information before you acquire an image.



4.0 How to Edit the Existing Patient Data

	Edit Patient		
*Last Name:	Smith		
*First Name:	Sandy		
Middle Name:			
*Patient ID:	123-44-5678		
*DOB: (MMDDYYYY)	01 /12 /1946		
*Age of Patient:	056Y		
*Gender:	F		
"NOTE: Fields with an asterisk '" are man	datory and require input.		
Clear	Cancel		Accept

Figure 44: The Edit Patient Dialog Box



This procedure only changes the patient information for images in the case study not yet taken. The Acquired images must be resent using Image/Spool Management and Resend, after the information is edited. See Image/Spool Management.



Caution:

Note

Note

Do not edit the patient information if you use a Modality Worklist. See *About Patient Information Edits with a Modality Worklist and PACS* on page 55.

To edit the Patient information:

- 1. Select Edit in the Patient Box to display the Edit Patient dialog box.
- 2. Make the changes to the fields that have wrong or missing information. Use the Tab key or the trackball to move through the fields.



The **Cancel** button closes the dialog box without any changes. The **Clear** button clears last name, first name, DOB, and age.

- 3. Select the **Accept** button.
- 4. Verify that the correct changes appear in the screen before you acquire a new image.



4.1 About Patient Information Edits with a Modality Worklist and PACS

- Make the patient information changes on the HIS/RIS *before* any information is sent to the Selenia.
- Make the patient information changes on the PACS if the images were sent to the Selenia.
- Any patient information changes on the Selenia must be a last option. Some RIS systems do not have the ability to change this information at the Selenia.
- The Modality Worklists transfer the Patient Information from a HIS/RIS system to the Selenia. Any change on the Selenia to this information can cause a mismatch of information when the Selenia images are sent to PACS. Work with your HIS/RIS administrator before you make any changes on the Selenia to the patient information.

4.2 How to Change the Patient Information After an Image Is Accepted

When an image is accepted, the image is immediately sent to the output. If the related information requires any change, then repreview, save, and resend the image as an additional record and delete the previous record.

Remove any wrong images from:

- Printer
- Diagnostic Review Workstation
- PACS

Correct the information at the AWS:

- 1. Select the Local Exams tab and search for the patient.
 - If patient information is wrong, select **Edit** in the Patient Pane.
 - If the accession number is wrong, select **Edit** in the Procedure pane.
- 2. Change the wrong patient information.
- 3. Select the **Accept** button.
- 4. Select the **Close Procedure** button.
- 5. Select Admin>Image Mgt.
- 6. Search for the patient.
- 7. Select the **Repreview** tab.
- 8. Select the first thumbnail image.
- 9. Select the **Repreview** button.
- 10. Select the **Save** button.
- 11. Repeat the steps 10 to 12 for the remaining images in the procedure. Eight thumbnail images display (if there were four in the original procedure.)
- 12. Select the last four thumbnail images (the corrected views).
- 13. Select **Resend** to all required outputs.
- 14. After you confirm that the four images were resent, have a manager delete the four images with the wrong labels.



5.0 New, Edit, Add a Procedure

5.1 How to Add a New Procedure

- 1. Open an existing Patient.
- 2. Select New in the Procedure pane.

Proced	lure	
Procedure: Procedure #:	Screening AN08262	Bilateral Mammog 003A
New	Edit	Add

Figure 45: New Procedure, Edit a Procedure, and Add a Procedure Buttons

- 3. Select **Yes** in the **Creating a New Procedure will close the current Procedure** dialog box. Any open procedure will close.
- 4. When the **New Procedure** dialog box appears, enter a new **Accession** Number.

	New Procedure Patient, Six				
*Accession Number: *Procedure Description:	Mammography 7 Screening-Bilateral Mar	nmography			
*NOTE: Fulls with an attentik ** are mandatory and require input					
Clear	Cancel	Accept			

Figure 46: New Procedure Dialog Box



Note

The **Cancel** button closes the dialog box without any changes. The **Clear** button clears the Accession number.

- 5. Select the **Procedure Description** from the drop down list.
- 6. Select the Accept button.

5.2 How to Edit a Procedure

- 1. Select the Edit button in the Procedure area of the screen.
- 2. Change the Accession Number. You cannot change the Procedure selection.
- 3. Select the **Accept** button.



5.3 How to Add a Procedure

The Add a Procedure function allows you to have many Procedures open at the same time.

1. Select the **Add** button in the Procedure pane.

	Add Procedure Patient, One	
*Accession Num	oer: 2000001	
*Procedure Descript:	on: Mammography Screening-Bilateral M	/ammography /
		<u> </u>
OTE: Fields with an asterisk '* are mandatory and require input.		

Figure 47: Add a New Procedure Dialog Box



Note

The added procedure must use the same Accession Number as the open procedure. If you do not plan to use the same Accession Number, select the **New** button and follow *How to Add a New Procedure* on page 56.

- 2. From the drop-down list, select the Procedure to add.
- 3. Select the **Accept** button.



6.0 Image Output



Figure 48: Select the Output

Before acquiring an image, verify that you selected the correct output. All patient images must be printed or committed to a PACS. Select the Output group to use for this procedure from the drop down list. To edit existing or create new output groups, see *Outputs* on page 68.

There is also a statement of the current number of Jobs in the Queue.



7.0 Exposure Information

7.1 The Exposure Pane

Comp Force Mode: Comp Release: Comp Thickness : Paddle II	FULL MANUAL 50.0mm	Coll Mode : Coll Size : Image Size: M FAST CENT	AUTOMATIC 24x29cm 24x29cm EP	Classifier Share Classifier Clas
Exposure More	Filter Mc Rh	27		

Figure 49: Exposure Techniques

The top, left side displays the status of these options:

- Compression Force Mode
- Compression Release
- Compression Thickness
- Collimation Mode
- Collimation Size
- Image Size
- Paddle ID (type and position)

Select the Exposure Techniques in the bottom area. Use the up/down arrows in the Exposure Mode area (the lower, far-left box) to select the Exposure Mode. The exposure default values for that mode appear to the right.



7.2 About Exposure Modes

Table 8: Exposure Modes

Exposure Modes (Item 1)	Operator Adjustments	Where the function is in the Techniques Pane
Auto-Filter	Select the AEC Exposure Adjustment (item 2); AEC Sensor (item 3) (Positions 1-7 or Auto).	1 2 3 Comp Force Mar IVLL Coll Made ALTOMATIC Comp Biddress Mar MARAL Coll Made ALTOMATIC Comp Trackers Storms Trackers EXERCIT: 24 MARAL CONTER The storm of the store
Auto-kV	Select AEC Exposure Adjustment; AEC Sensor.	Comp Flows Mode PULL Coll Mode: AUTOMATIC Comp Flows Mode MANIAL Coll Size: 20:0% Comp Flows Works Mode MANIAL Coll Size: 20:0% Public DL, 24:04/09CM FAST CENTER TATTORY The Cont Mode Mania
Auto-Time	Select kV; Filter; AEC Exposure Adjustment; AEC Sensor.	Comp Frees Mode: FIUL. Comp Endease Mode: MANNAL: Call Size: 2 hz/Rem Comp Endease Mode: MANNAL: Call Size: 2 hz/Rem Tradication MATCHINE: Software Field Size: 2 hz/Rem Tradication Manual Call Size: 2 hz/Rem Tradi
TEC	Select Breast Density (item 4), then select Accept or change the Exposure Techniques, which puts you in Override (Manual) Mode.	Compression Force Mode: PRE Collimato: AUTOMATIC Compression Roleway Mode: VASUAL Recepto: 2420 HTC GRD Termine Termine Pression Roleway Mode: PRE Collimato: AUTOMATIC Termine Pression Roleway Mode: PRE Collimato: AUTOMATIC Pression Roleway
Manual	Manually calculate and select all x-ray techniques and Exposure Factors (kV, Filter, and mAs).	Comp Flores Made: PULL Comp Flores Made: MULL Comp Flores Made: MANUAL Coll Size: 242590 Padder D: 21/25/15/201 FAST CENTER MANUAL Mo Rh Rh Rh Rh Rh Rh Rh Rh Rh Rh Rh Rh Rh



Note

When you change the mode, all Exposure Techniques reset to the default values for that mode.



7.2.1 About the Manual Exposure Mode

In the Manual mode the Operator sets all Exposure Techniques. The default values appear when you select the View, then you make any necessary adjustment.

- To change an Exposure technique with the trackball, scroll to the value with the up/down arrows, or select the option box.
- To change an Exposure technique with the keyboard, press the Function Key for the item at the top of the keyboard.

7.2.2 About the TEC Exposure Mode

The Tissue Exposure Control (TEC) Mode is an enhanced Manual Exposure Control mode.

1. Before you position and compress the breast, select the **Breast Density** type on the Acquisition Workstation screen or press the **Breast Density** key.



Figure 50: Breast Density Options

- 2. Compress the breast. The x-ray exposure techniques for kV, mAs, and filter update.
- 3. Select the **Accept** button in the Breast Density area (or press the **Enter** key). When "Ready" displays in the Generator Status message area, the system is ready to acquire an image.

Comp Force Mode: Comp Release : Comp Thickness : Paddle II	FULL MANUAL 50.0mm D : 24CMX29C	Coll Mode : Coll Size : Image Size: M FAST CENT	AUTOMATIC 24x29cm 24x29cm ER	Generator Status Standby Generator Error Reast Generator	\mathbf{X}	Dystem Messages	
Exposure Mode TEC	Triter Mc Rh	27	/ ↑ 1	20.0	VT	NORMAL ACCEPT	↑

Figure 51: Accept TEC Exposure Techniques

If you change one or more of the parameters (kV, mAs, or filter), the system enters the Override Mode. When the Override message displays, the system is ready to acquire an image.





Figure 52: The Override Mode

There are three methods to exit the Override Mode and return to TEC mode:

- Change the Breast Density setting.
- Release the compression then compress the breast.
- Select TEC mode a second time.

7.2.3 AEC Exposure Mode

Note

The Automatic Exposure Mode (AEC) uses the pre-exposure Scout pulse, and for some modes the compression thickness to calculate the exposure.

The AEC options are Auto-Filter, Auto-kV, and Auto-Time. See *About Exposure Modes* on page 60 for the exposure technique selections available with each mode.

7.2.4 AEC Exposure Adjustment Settings



AEC exposure adjustment is not available in all geographic regions.

You can use the AEC Exposure Adjustment controls to increase or decrease the target pixel value. Each step changes the target pixel value by approximately 15% and makes a related change in dose. See *figure AEC Exposure and Sensor Position Adjustments* on page 63.

Set this control in a range from +4 to -3 to increase or decrease the exposure. This adjustment remains until you change the adjustment or restart the system.







Figure 53: AEC Exposure and Sensor Position Adjustments

7.2.5 AEC Sensor Position Indicator Settings

Select one of the seven positions to indicate the area of interest for AEC calculations. Or, select Auto AEC where a larger area is tested.



Note

Always confirm the position of the AEC Sensor before an AEC exposure.

The AEC Sensor Position has eight available positions.

- The Positions 1 through 7 are manually selected, from the chest wall edge (position 1) to the nipple edge (position 7).
- Position 8 (Auto) automatically positions the sensor.

To change the AEC Position selection, use:

- The buttons on the compression device.
- The AEC Pos key on the keyboard.
- The AEC position arrows on the Acquisition Workstation display.

Set the AEC Sensor Position after you compress the breast. The displays on the side of the Compression Device and the lower right corner of the screen indicate the position setting.



7.2.6 AEC Alarm Messages

When the system cannot use the selected AEC mode to acquire an Image, one of the following messages displays.

Message	Reason and Corrective Action
Calculated Exposure	The AEC Sensor is over a thin or small area of breast tissue.
Time Less Than Min	Use the AEC Sensor in a position under the breast tissue.
	Alternately, use AutoTime and a lower kV.
Calculated Exposure	Very dense breast tissue, an implant, pacemaker, or other
Time Exceeds Max	anatomy changed the AEC sensor. Move the manual AEC
	sensor where there are no obstructions. Alternately, use
	AutoTime and a higher kV.

1. Select **OK** in the dialog box.

2. Select the **Reset Generator** button.

8.0 Menu Bar

The Menu Bar provides easy access to non-routine Acquisition Workstation options. These options are available while the Acquisition Workstation application runs unless a dialog box or Procedure is opened.

The table below shows the Menu Bar options when a patient procedure is *not* selected.

Menu	Options	Function
File	Exit	Exit from Acquisition Workstation
Edit	Standard Setup	Select the startup default values (Standard
		Setup Screens on page 66).
	View Order Editor	Change the view order (View Order Editor on
		page 67).
	User Setup	Add, edit, delete a user profile (Add or Edit a
		User on page 67).
	Outputs	Add, remove output devices (Outputs on
		page 68).
Admin	Image Management	Retrieve the images from the image spool
		(Image Management on page 70).
	Protect Patients	Protect or Unprotect patient records (Protect
		<i>Patients</i> on page 72).
	Manage Queues	Find a specified job in the queue (Manage
		<i>Queues</i> on page 72).
	Eject	Remove a disk from CD/DVD drive (Eject on
		page 74).

Table 10: Menu Bar Options and Functions

Menu	Options	Function
	Import	Copy images from a CD to the system (<i>Import</i>
		on page 74).
	Retrieve Priors	View previously-acquired images (Retrieve
		Priors on page 75).
	PPS Status*	Additional options (when MPPS Service is
		installed) for close of a procedure (MPPS
		Status on page 75).
	Calibrate	Access calibration procedures (Calibrations on
		page 75).
	Test Patterns	Access test pattern procedures (Test Patterns
		on page 76).
	DR Device Control	Access by Service personnel (DR Device
		Control on page 77).
	Available Disk Space	Displays status of hard disk drive space
		(Available Disk Space on page 77).
Stereotactic**	Enable or Disable Stereo	Engage or disengage the stereo mode.
	mode**	Available when the StereoLoc II is installed.
Info	About the Acquisition	(Info Menu on page 77)
	Workstation	

Table 10: Menu Bar Options and Functions

* This option appears when the MPPS Service is installed.

** This option appears when the Stereotactic Service is installed.



8.1 Edit Menu Function

8.1.1 Standard Setup Screens

Pre-compression Force (%)	50 7
Full Compression Force (%)	100 🔻
Compression Force Units	Ibs. 7
Compression Release Mode	
Compression Force Mode	FULL
Collimator	
Default Exposure Mode	MANUAL
Default Magnification Mode	MANUAL
Set Technique Defaults	Close
Move your mouse over a field to display help information here	

Figure 54: The Setup Screen

- 1. Select Edit>Standard Setup.
- 2. Use the trackball or the keyboard Tab key to move between fields.
- 3. Use the trackball or arrow keys to display and select options.
- 4. Select the **Close** button to exit the dialog box.



When Manual Collimation is selected, the function that automatically changes the Collimation when a paddle is changed or moved turns off.

8.1.2 Set Technique Defaults

Note

Select the **Set Technique Defaults** button to display the Technique Default Setup screen. You must set the default values for all modes.

- 1. Select Manual Exposure Mode and select the system startup defaults.
- 2. Change the mode to Auto-time and select the options.
- 3. Continue to change the mode and select the options for all remaining exposure modes including Magnification modes.



8.1.3 View Order Editor



Figure 55: View Order Editor

- 1. Select Edit>View Order Editor.
- 2. Select the procedure from the drop-down menus.
- 3. Select the Views for each Procedure:
- 4. Move the View icons to change the order.
- 5. Select **Close**. Your view order preferences load when you log in.

8.1.4 Add or Edit a User

A signed-in user can add or delete a user at the same or lower level. To access the Edit User options, select **Edit>User Setup**.



Figure 56: The Edit Users Dialog Box



Table 11: User	r Setup	Options
----------------	---------	---------

Option	Steps
Add a new user	1. Select New.
	2. Enter the requested information. The password must contain a minimum of 6 characters.
	3. Select the Accept button.
• Edit a user	1. Select your name from the list.
	2. Select Edit.
	3. Change the Initials, if necessary.
	4. If requested, enter the Password. The password must contain a minimum of 6 characters.
	5. Select the Accept button.
Delete a user	1. Select the User to delete.
	2. Select the Delete button.
	3. Select the Yes button to the confirmation prompt.
Close the dialog box	Select the OK button to close the Edit Users dialog box.

8.1.5 Outputs

To Add New Output Groups:

- 1. Select **Edit > Outputs**.
- 2. Select New.
- 3. Enter the name for this Output.
- 4. Select OK.
- 5. Select the new name in the **Edit Outputs** dialog box.
- 6. Select Edit.

- Edit output: L	aserPrinter2
archives cdrw physician_displays isser_carreras vdevagfs4500M_001	Devices currently selected: DS4500M Remove Item Configure
	Cancel Accept

Figure 57: Edit Output Device Dialog Box



Note

Outputs are first created during installation.

- 1. Select the tab for the type of device.
 - CDRW = CD-RW
 - Laser Camera = Printer
 - Physician Display = Diagnostic Review Workstation, Technologist Review Workstation, CAD, etc.
 - Archive = PACS
- 2. Select the device name.
- 3. Select the **Add** button to move it to the right column.
- 4. To add another output device to the new group, select the tab for the device type.
- 5. Select the device name.
- 6. Select the **Add** button.
- 7. Select the **Accept** button when finished with the group.
- 8. Select the **OK** button.

To Edit Outputs:

- 1. Select **Edit > Outputs**.
- 2. Select an Output name. If the Output keeps its current name, go to step 7.
- 3. Select the **Copy** button.
- 4. Enter a new **Output name**.
- 5. Select the **OK** button.
- 6. Select the new name in the Edit Outputs dialog box.
- 7. Select the **Edit** button.
- 8. Select the tab for the type of device to add.
- 9. Select the device to add.
- 10. Select the **Add** button to move it to the right column.
- 11. Select the name of any output in the **Devices currently selected** column that is not needed in the group.
- 12. Select the **Remove Item** button.
- 13. Select the **Accept** button.
- 14. Select the **OK** button.



8.2 Admin Menu

8.2.1 Image Management

1. Select **Admin>Image/Spool Management**. The Choose a New Patient dialog box appears.

-		Choose A New	Patient		
Patient Nam	e Patient ID				
Enter Patient	Name (* to search):				
Patient^Two	D*				Search
	Search Results: 1 entries				Select
	Name		ID Se	x	DOB
	Patient, Two	00581382	F	05/06/1932	
noxt					
					Close

Figure 58: Choose a New Patient Dialog Box

- 2. Select the **Patient Name** tab or the **Patient ID** tab.
- 3. Enter the search information in the Enter Patient Name or ID box.
- 4. Select the **Search** button.
- 5. Select a patient from the list.
- 6. Select the thumbnail image of the image to resend or repreview.
 - You can select more than one to resend.
 - If there are more than four images, use the arrows on the side to scroll.
- 7. Select the Tab for **Repreview** or **Resend**.
- 8. Continue with directions in *The Resend Options* on page 71 or *The Repreview Option* on page 72.



Note

If the commit function is enabled and archive has committed the image to storage, the line "Commit: Accepted" appears in the information about the image in the upper right pane. See the *figure Image Management Resend Options* on page 71.



8.2.2 The Resend Options



Figure 59: Image Management Resend Options

- 1. Select the **Resend** button.
 - To resend all images on the hard disk drive for this patient, select Resend All Images.
 - To resend selected images, select the thumbnails of the images to resend, then select the **Resend Selected** button.
- 2. Select the Output from the drop-down list.
- 3. If the selected Output has a printer, select the **Print Image Processing Options**.
- 4. Select the **Resend** button.
- 5. Select the **OK** button.



You cannot change any Patient or image object information from the resend function. This includes image, LUT modifications and DICOM information such as Patient name spelling, ID, and comments. To send new information you must make a new file with the Repreview function, save it, and Resend the new file.



8.2.3 The Repreview Option

- 1. Select the thumbnail image to review.
- 2. Select the **Repreview** tab, then select the **Repreview** button.
- 3. When you finish reviewing, select the **Cancel** button.
- 4. To create a new file to resend, make any required changes to the markers, comments, or use the **Edit View** function, then click **Save**. If you select a FOR PRESENTATION image, you cannot change the markers and ROI is not enabled.
- 5. Make any necessary changes in the **Contrast** or **Density**. These changes do not save when you save the image.
- 6. Select the **Save** button.



Make sure that you send the newly-saved image. The original is a possible selection.

8.2.4 **Protect Patients**

Note

Use the Protect function to prevent automatic removal of patient records from the hard drive.

- 1. Select Admin>Protect Patients.
- 2. Search for the patient to protect.
- 3. Select the patient to protect from the search results.
- 4. Select the **Protect** button at the bottom of the screen.
- 5. When the confirmation dialog box appears, select the **Yes** button.
- 6. Select the **Close** button.

To create more space on the hard disk drive, unprotect patients who do not need protection.

To unprotect a patient record:

- 1. Select the patient.
- 2. Select the Unprotect button.

8.2.5 Manage Queues

Note

Use Manage Queues to find problem jobs or a specified job in the queues.



Warning:

When you select Delete Job, the job is permanently deleted. If you did not send the job, resend it to a storage device later.



Chapter 6—The Select a Patient Screen Menu Bar





Figure 60: Manage Queues

Figure Legend

- 1. View the list of available queues.
- 2. Enable or disable a job spooler.
- 3. Update the list of Queued Jobs.
- 4. View the list of jobs in the queue.
- 5. Remove the inactive jobs from the queue. Resend stopped jobs.

Table 12: How to Use Manage Oueu	2 12	Table 12: How to Use	Manage	Oueues
----------------------------------	------	----------------------	--------	--------

Option	Steps
• To find the status of a job:	 In the Devices list, select the queue to view. Select the OK button.
• To delete a job:	 Select the queue in the Devices list. Select the Suspend <name> Spooler button.</name> Select a job from the Queued Jobs list, or Ctrl+click to select many jobs. Verify that the selected job is the job to delete. There is no "undo". Select the Delete From Queue button.
	6. Select the OK button.
• To resend a stopped job:	 Select the queue in the Devices list. Select a job marked 'Stopped' from the Queued Jobs list, or Ctrl+click to select many jobs. Verify that the selected job is the job to resend. Select the Resend Stopped Job button. Select the OK button.



8.2.6 Eject

Note

Select **Admin>Eject** to open the CD-RW drive drawer when there is a CD in the CD-RW drive.



When the CD-RW drive is empty, press the button on the CD-RW drive to open the drawer.

When there is a disk in the drive, the drive drawer opens only from the menu. To close the drive drawer, press the CD-RW drive button.

8.2.7 Import

To import the information from a CD:

- 1. Open the CD-RW drive drawer.
 - Press the CD-RW drive button to open the tray if there is no CD in the drive.
 - Select Admin>Eject to open the drive if there is a CD in the drive.
- 2. Put the CD that contains the images to import on the tray.
- 3. Press the CD-RW drive button to close the tray.
- 4. Select Admin>Import.
- 5. In the dialog box, find the file (or files) to import.
 - Select a "+" to open a list of folders and/or files.
 - Select the empty box to select the file or folder.



Tip:

Note

To deselect a file, click the checkmark. When you deselect a folder, the files below the folder are not deselected. You must clear each file.

6. Select the **Import** button. The file is copied from the disk and is available in the **Select a Patient** screen.



Do Not press the **Eject** button on the drive while the system reads from or writes to the disk.

If you try to eject or read the CD while the import process finishes, you can cause the drive to stop.



8.2.8 Retrieve Priors

To find the Images in the Local Database, select the Local Exams tab and search for the Patient.

To Retrieve Priors from the PACS system:

- 1. Select Admin > Retrieve Priors.
- 2. Enter the patient name in the Name field or ID in the ID field (wildcards allowed).
- 3. To limit the results to a date range, select the checkbox **Search by Date Range**, then enter the Date Range for the Prior Images. The date format must match the configuration setting, (mm/dd/yyyy or dd/mm/yyyy).
- 4. Select the **Query** button or press the **Enter** key. If the button is not enabled, you need more characters in either the **Name** field or **ID** field.
- 5. Select the patient name (or names) to import.
 - You can select many patients.
 - You can deselect patients.
 - You can use the title bars to sort by field.
- 6. When the **Import** button activates, select **Retrieve and Import**. The images import into the Acquisition Workstation Local Database.
- 7. Select the **OK** button.
- 8. Select the Close button.

After the import completes, select the Local Exams tab to display the Patient Images. If there are more than 50 records in your query request results, adjust your search.

8.2.9 MPPS Status

When this optional feature is installed, a menu item is added to sites with the MPPS Service Class Provider. Workflow changes with three Close Procedure options instead of one option.

8.2.10 Calibrations

Caution:

When Calibration is needed, the Select a Patient screen remains in Standby and the **Calibration Needed Alarm** message appears in the **System Message** box.

When you turn on the system, a message appears if the system needs any calibration. You cannot acquire the patient exposures until you perform the required calibration. All the review, resend, and administrative functions continue to work.



It is important to follow the directions on the screen when you calibrate a system.

Selenia Instructions for Use

Chapter 6—The Select a Patient Screen Menu Bar

Note





Clean the detector and acrylic block before you perform a Calibration.

If you are in the Calibration Mode screen by accident, select the **End Calibration** button. If you started the Calibration, but you did not accept images, you can reject the Image and exit the Calibration.



When you accept one image, do not end the calibration before you acquire, view, and accept all the images.

8.2.11 Test Patterns

Note

Note



See DICOM Printer Quality Control in the Selenia Quality Control Manual for more information about the SMPTE test pattern.

- 1. Select Admin>Test Patterns.
- 2. Select the Preview size (item 1, from the image below).
- 3. Select the **Test Pattern** from the **Pattern** field (item 2, from the image below).
- 4. From the **Output** area of the screen (item 3, from the image below):
 - Select the Output **Device**.
 - Select the Film Size.
 - Select the **Print True Size** check box.

		Pattern: SMPTE	7
Full Screen Maximized Window			
Output			
Size: 18CMx24CM Paddle: 3328x2560	🏹 📝 Print True Size (if available)		
Devices: 8900	7		
Preview	w Send Clean Up Close		
Previe	w Send Clean Up Close		

Figure 61: Test Pattern Dialog Box

- 5. Select from the following options (item 4, from the image above):
 - To view the test pattern **Full Screen**, select the **Preview** button. To return to the dialog box, click any area on the Full Screen.
 - To send the Test Pattern to the selected output, select the **Send** button.
 - To remove any files from previous "sends" before you send the current images, select the **Clean Up** button.
 - To exit the Test Pattern option, select **Close**.



8.2.12 DR Device Control

This function is for use by service personnel.

8.2.13 Available Disk Space

This option displays the remaining number of images the hard disk drive and CD can hold.

The system automatically deletes some images/patients at specified levels to get back hard disk drive storage space for new procedures. Automatic image removal does not include the following:

- Images in a queue to an output device.
- Images for protected patients. (Make sure that you unprotect patients when you do not need their files.)
- Images that are not successfully committed to an archive device. (Archive devices send commitments after Images go to long-term storage.)



Caution:

If the disk does not have enough available space, the system does not acquire an image. A message displays the capacity of the disk. Delete unneeded images to restore the disk space.

8.3 Info Menu

The Info menu displays the information about the Acquisition Workstation.

8.3.1 Gantry ID

To find the Gantry ID (serial number):

- 1. Select the Info tab.
- 2. Select About the Acquisition Station.



Figure 62: Location of the Gantry ID (serial number)



9.0 Special Modes Buttons

9.1 Non-imaging Mode Button

The Non-imaging mode is available to all users and allows the x-ray exposure without the acquisition of images. This mode is not available when you have a patient selected. When you select the Non-imaging mode, the Image Receptor deactivates.

To access this mode, select the **Non-Imaging Mode** button.

Non-Imaging Mode	

Figure 63: Non-Imaging Mode Button



Caution:

Protect the Image Receptor from excessive radiation by covering it with lead while using the Non-imaging mode.

To exit the Non-Imaging mode, select the End Non-Imaging Mode button.

9.2 The Review Button

This button is used during the Reject /Unreject process. See *How to Accept a Rejected Image* on page 99.

9.3 The Simulate Capture Button

This function is for use by service personnel.

10.0 Generator Status

This area displays the current status of the system. See the *figure The Select a Patient Screen* on page 51.

11.0 System Messages

The System Messages area displays the system status.



12.0 Alarms



Figure 64: The Alarm Icon

Alarms are generally output problems that occur due to jobs that fail at the output device. The Alarm Icon shows the number of alarms. Select the icon to view the details. Erase the alarm to clear it.





Digital Mammography System

1.0 Patient Views

When you select a patient, the Views for the scheduled Procedure display on the Patient View screen in your preset order. The program automatically highlights the first View in the sequence. To choose a different View, select that View.



Figure 65: Patient View Screen

Торіс	Information
Work with Views (item 1)	Work with Views on page 82
Add/Edit the Patient Information	How to Add a New Patient on page 53
Add/Edit a Procedure (item 2)	New, Edit, Add a Procedure on page 56
Select the Image Output	Image Output on page 58
Jobs in the Queue (item 3)	Displays the number of jobs in the outputs.
Change Exposure Techniques	Exposure Information on page 59
Acquire the View	Acquire Images on page 83

Work with Views 2.0



Figure 66: Buttons Used on the View Screen

Table	14:	View	Options
1 110 10	- - .	1 10 00	opnono

Option	Actions	
Add View (item 1)	1. Select the Add View button.	
	 Select an additional View from the list of Standard Mammography Views. To add multiple views, hold down the Control key and select the View. 	
	3. Select the OK button.	
	4. Select the new view icon before you acquire the image.	
Remove Unexposed (item 2)	Unused view icons are automatically reclaimed, but you can delete the views with the Remove Unexposed Views button.	
	 Select the All button to remove all unused icons, or the Selected button then indicate which views to delete. 	
	2. Select the Remove Unexposed button.	
Implant Present (item 3)	Select the box if there is an implant present in the breast. The box should remain checked for all views in the procedure.	
Close Procedure (item 4)	1. Select the Close procedur e button.	
	 Select the Yes button to close the procedure or the No button stay on the screen. 	



3.0 Acquire Images

The system is ready to acquire images when the Generator Status indicates "Ready" (item 1 in the figure below), and the X through the x-ray symbol (item 2, in the figure below) in the Generator Status box is not displayed.



Figure 67: Ready for Exposure Indicators


Chapter 8 The Preview Image Screen

1.0 Introduction

The Preview Image screen appears when you acquire an image. The right side of the screen has image tools and buttons to Accept or Reject the image.



Figure 68: The Preview Image Screen

Figure Legend

- 1. Acquired Image Display Area
- 2. Tools

2.0 The Preview Image

The Preview Image is a non-diagnostic, processed image for anatomy and position confirmation. The Patient Name, ID, and the View appear on a bar on the top of the Preview.

2.1 **Progressive Preview**

The first preview image appears for a quick check of the breast position. This image updates with an image of better quality to check for motion or image artifacts.



2.2 Film Label Area

The film label area which is the small rectangle in the upper, left corner is blank in the Preview.

A Digital Marker next to the label displays the Technologist initials and the marker for the selected View. You can move or change the marker.

2.3 Hanging Options

The Images in this manual are displayed in the Dorsal orientation (the Operator faces the patient).



Figure 69: Hanging Options

This option can affect how images display on your diagnostic workstation. Confirm the effect of the options with the vendor.



3.0 Tools for Image Enhancement and Annotation

Figure Legend

- 1. Comments
- 2. Markers
- 3. Edit View
- 4. Implant Present
- 5. Zoom/Pan Thumbnail
- 6. Crosshair
- 7. Measurement
- 8. Window/Level
- 9. ROI size Drop-down
- 10. Set Center/Width
- 11. Display AEC Regions
- 12. Exposure Index
- 13. Exposure Techniques
- 14. Reprocess View
- 15. Magnification
- 16. Full Zoom/Pan
- 17. Quick Zoom/Pan
- 18. ROI



Figure 70: The Preview Screen Tools



3.1 Comments

The comment text is inserted in the DICOM header of the Acquired Image. The diagnostic review workstation displays the comments.

Patient Informatio	n
Technologist:	Andrea
Patient:	Test^Patient
Exam Time:	20030516 084538
Study:	Screening-Bilateral Mammography
Additional C	omment
Additional C	omment.

Figure 71: Image Comments

- 1. Select the **Comments** button.
- 2. Select the Additional Comment checkbox.
- 3. Select the field below the checkbox and enter your comment.
- 4. Select the **OK** button to save the comment.



Note

To remove all comments, select the **Clear Comments** button. To close the dialog box without a change to the comments, select the **Cancel** button.

3.2 The Edit View Button and Markers

Select the **Edit View** button to update the DICOM header and enable the image to hang correctly at the Diagnostic Review Workstation. The tabs and buttons allow you to change just the Marker.

	RMLO	RML	RLM
RRL	RRM	RRS	RRI
	Clear All M Edit Vi	larkers ew	

Figure 72: The Markers in the Preview Pane



Selenia Instructions for Use

Chapter 8—The Preview Image Screen Tools for Image Enhancement and Annotation

3.2.1 How to Use the Marker function

- 1. If the marker is wrong, select the Clear All Markers button
- 2. Select the Tab for the set of markers you need.
- 3. Select the marker you need. You can select a maximum of three markers from the available set for each image.
- 4. Move the marker to the correct position.

If you change a Marker, you change the label on the image, but not the DICOM header. You will not see the change on the Patient View page.

3.2.2 How to Use the Edit View button

Before You Accept the Image

- 1. Select the Edit View button.
- 2. Select the required view from the displayed list.
- 3. Select **OK**. The correct image displays on the preview screen and the marker and the direction the image hangs are correct.

After You Accept the Image

If the image is Accepted before you notice the wrong view, the incorrect image was sent to the selected output. Make the corrections, then resend the corrected image with the Image or Spool Management menu. See *Image Management* on page 70.

- 1. Before you make any changes:
 - If you printed the mislabeled image, find and discard that image.
 - If the image was sent to a Diagnostic Review Workstation, delete the mislabeled image.
 - If the image was sent to the PACS, tell the PACS administrator to delete the mislabeled image.
- 2. Repreview the Image. (See *The Repreview Option* on page 72.)
- 3. Select the **Edit View** button on the Preview screen.
- 4. Select the required view from the displayed list.
- 5. Select Save. A new thumbnail image appears. Resend this image.

3.3 Implant Present

Select the checkbox on the Patient View screen when an implant exists. When the checkbox is selected, the system marks future images with an Implant Present label. The checkbox on this screen enables you to change the selection before you accept the image.

Chapter 8—The Preview Image Screen Tools for Image Enhancement and Annotation



3.4 Quick Zoom/Pan

Quick Zoom/Pan is active after the image is first displayed, before image processing completes.

- 1. Select the **Quick Zoom/Pan** button.
- 2. To pan through the image, drag with the right trackball button.
- 3. To turn off the function, select the **Quick Zoom/Pan** button.

3.5 Full Zoom/Pan



Figure 73: Full Zoom/Pan Image



The Full Zoom/Pan function becomes available after image processing is complete. The box in the thumbnail image on right side of the screen shows where the magnification view is in the complete image. You can move this box to any area in the image.

- 1. Select the Full Zoom/Pan button.
- 2. To move through the image, drag with the right trackball button.
- 3. To turn off the function, select the Full Zoom/Pan button again.



Selenia Instructions for Use

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3.6 ROI



Figure 74: ROI Size Drop-down Menu

Method 1: Draw a custom box.	Method 2: Use a pre-defined box
1. Select the ROI button.	1. Select the ROI button.
2. From the drop-down menu, select	2. From the drop-down menu, select
User Draw.	the size.
3. Click in the image and drag to draw	3. Click the area of interest in the
the size you need.	image.



3.7 Crosshairs



Figure 75: Preview with Crosshairs

- 1. Install a localization paddle.
- 2. Acquire the image.
- 3. If necessary, select the **Zoom/Pan** button and move through the image.



- 4. Select the **Crosshair** button
- 5. Select the area of interest in the image.
- 6. Drag the crosshairs to the needed position, or select another area.
- 7. If necessary, use the **Zoom/Pan** functions to follow the crosshair lines to the grid from the localization paddle. The crosshair lines remain when the **Zoom/Pan** function is enabled and disabled.



Selenia Instructions for Use

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3.8 Measurement



Figure 76: A Measurement on the Preview Screen

- 1. Select the **Measurement** button
- 2. Click the first point then drag to the second point. The distance displays when you release the button.
- 3. Select the "+" at either end of the line to clear. There lines are not saved with the image.



3.9 Magnification

Note

Use the Magnification function **I** to view part of the image at 2X the displayed resolution.

- 1. Select the **Magnification** button.
- 2. Move the box that appears at the area of interest.
- 3. To turn off the magnification function, select the **Magnification** button a second time.



You can use the magnification function on a Full Zoom image.



Figure 77: Preview with Magnification



Selenia Instructions for Use

Chapter 8—The Preview Image Screen Tools for Image Enhancement and Annotation

3.10 Window/Level

Note



- 1. Select the **Window/Level** button _____
- 2. Select in the image and hold while you move the trackball.
 - Roll Up or Down to change the brightness (Level).
 - Roll Right or Left to change the contrast (Window).
- 3. To return to the original settings, double click the image.
- 4. Select the **Window/Level** button to turn off the feature.



This operation does not change the final, processed Image.

Center:	2000	Width:	2500	Set

Figure 78: The Window/Level Settings

To enter a known **Window/Level** value:

- 1. Select the field (Center or Width) to change.
- 2. Enter a new number.
- 3. Select the **Set** button.

To return to the original setting, double click the image.



3.11 Display AEC Regions

To display the AEC area selected by the system, select the **Display AEC Regions** checkbox.

- Manual sensors show on the image as a white rectangle with rounded corners.
- The two Auto Sensor areas show on the image as 1cm by 1cm white squares.

To hide the AEC areas, deselect the **Display AEC Regions** checkbox.



Figure 79: AEC Regions



3.12 Exposure Index

Use the Exposure Index as a general guide to compare the selected Exposure Techniques to the quality of the acquired image.

- "Image Status: OK" shows that the image processing was applied.
- "Image Status: Raw" shows that image processing was not applied.
- A number after the image status shows a problem with the image processing. Reject the image. Contact Service Support.



Figure Legend

- 1. Exposure Index-pixel count
- Low exposure area—evaluate the image for excess noise. Re-acquire, if necessary.
- 3. Satisfactory, but below ideal exposure area
- 4. Ideal exposure area
- 5. High exposure area the image is more than sufficient. You can reduce the radiographic technique for the next exposures.
- 6. The Exposure Index line related to imaging of the ACR phantom
- For Manual and TEC modes, the Exposure Index calculates an area approximately one centimeter from the chest wall on the centerline of the image.
- For all AEC modes, the Exposure Index calculates for the AEC areas used.

3.13 Exposure and Dose/Entrance Surface Exposure Information



Figure 81: Exposure Information

The exposure information is displayed in the Tools area. An estimate of the average glandular organ dose and the entrance surface exposure are displayed.



4.0 Accept or Reject

Select Accept or Reject to close the Preview screen. The sequence continues by highlighting the next view.

4.1 Accept

To accept the image, select the **Accept** button.

- The full resolution image with all related attributes is marked as accepted in the pool.
- The system transmits the image to the selected output devices.

4.2 Reject

Note

To reject the image:

1. Select the **Reject** button.



Reject Analysis uses this information (required for Quality Control).

Image Rejection Information		
	Tech: Andrea - Rejection List	
	Patient Motion	
	O Detector Underexposure (excessively noisy images)	
PHX: ID	O Improper Detector Exposure (Saturation)	
	⊖ Artifacts	
	O Incorrect Patient ID	
	O X-ray Equipment Failure	
	🔿 Software Failure	
	🔿 Blank Image	
	⊖ Wire Localization	
	○ Q.C.	
	O Aborted AEC Exposure	
Patient: Patient, One		
Exam time: 12/11/08 05:45 PM		
	Return to preview Reject	

Figure 82: The Reject Reasons List

2. Select the reason for the reject.



3. Select the **Reject** button. The system moves the image to the Reject Bin. An asterisk displays next to the View label of a rejected image.



Figure 83: A Rejected Image

5.0 Additional information about Accepted/Rejected Images

After you reject an image, the Preview closes and you can:

- Re-acquire the rejected View.
- Select a different View.
- Close the Procedure.

5.1 The Image Repetition Information Dialog Box

When you accept a second image for the same View, the Image Repetition Information dialog box appears.

- 1. Select the reason for the repeat.
- 2. Close the dialog box.

5.2 How to Accept a Rejected Image

To accept a rejected image:

- 1. Reject the image that was taken as the replacement for the rejected image.
- 2. In the Patient View screen, select the **Review** button (below the Output box).
- 3. In the **Review** dialog box, select the tab for the View.
- 4. Select the **Rejected Image** thumbnail image to accept.
- 5. Select the **Comments** button to add or delete comments.
- 6. Select the **OK** button to close the **Comments** dialog box.





Figure 84: Set Accepted Button on Review Screen

- 7. Select the **Set Accepted** button.
- 8. Select the **OK** button to close the **Review** dialog box.



Figure 85: Accept Rejected Image

When you accept a rejected image, the Send Image to Output dialog box displays. Select Yes to send the images to the selected output group.



Note

If you select the **No** button, you can resend the images with the Image/Spool Management function.



Additional information about Accepted/Rejected Images

5.3 How to Reject an Accepted Image

To reject an Accepted Image:

- 1. In the **Patient View** screen, select the **Review** button.
- 2. In the dialog box, select the tab for the View to reject.
- 3. Select the Accepted Image.



Figure 86: Set Rejected Button

- 4. Select the **Set Rejected** button.
- 5. Select a reason for the reject. See the *figure The Reject Reasons List* on page 98.
- 6. Select the **Reject** button.
- 7. Select the **OK** button. The displayed thumbnail image disappears from the Patient View screen.



Note

After you reject an Image with this method, the image is **not** removed from the output queues. To delete the image from the queues, see the table in *Manage Queues* on page 72. The Rejected Image is added to the Reject Analysis Bin for tracking purposes.

Chapter 9 Maintenance and Cleaning

1.0 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.



Digital Mammography System

Caution:

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

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

1.1 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 and water with one part commercially available chlorine bleach (normally 5.25% chlorine and 94.75% water) and nine parts water
- 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 which touch the patient.





1.2 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 most safe and effective method.

2.0 Care and Cleaning—Acquisition Workstation and Value Console

2.1 How to Clean the Preview Display

Avoid touching the display screen.

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 ingredients described above and used according to the directions of the manufacturer can be used.



2.2 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.

3.0 **Preventive Maintenance**

			Reco	mmended	Frequency		
Maintenance Task Description	Each Use	Daily	Weekly	Bi- weekly	Monthly	Quarterly	Semi- annually
Clean & disinfect paddle	х						
Clean & disinfect breast	x						
platform							
Visually inspect all paddles for	х						
damage							
Recommended/ required start		x			х		
of operation verifications							
All daily recommended/		x					х
required calibration							
Diagnostic Review Workstation			x				
Quality Control *							
Detector Flat Field Calibration *			x				
Artifact Evaluation *			x				
Phantom Image *			x				
Signal to Noise / Contrast to			х				
Noise Measurements *							
Compression Thickness				х			
Indicator *							
DICOM Printer Quality Control			x				
View boxes and Viewing			х		х		
Conditions							
Repeat/Reject Analysis*						x	
Compression *							х
Visual Checklist *					х		
Emergency Stops					х		
Clean array dust filter					х		

Table 15: User Preventive Maintenance

* Refer to Quality Control Manual



Digital Mammography System

- 1.0 Dimensional Information
- 1.1 Tubestand (Gantry with C-arm)



Figure 87: Tubestand Dimensions

- A. Height with C-arm Travel
- B. Height
- C. Width
- D Depth Weight

213.0 cm (84.07 inches) ±1.0 cm (0.4 inches) at highest C-arm travel

190.0 cm (74.8 inches)

65.0 cm (25.6 inches) ± 1.0 cm (0.4 inches) (C-arm at 0° position)

110.0 cm (43.3 inches)±1.0 cm (0.4 inches) 300 kg (661 lb) ±20 kg with Image Receptor

1.2 Acquisition Workstation



Figure 88: Acquisition Workstation Dimensions

- A Height
- B Width
- C Depth
- D Swivel Arm Radius Weight

203 cm (79.9 inches) (maximum)

107 cm (42.1 inches)

76 cm (30.0 inches) 60.5 cm (23.81 inches) 120 kg (264.5 lb) (maximum)



1.3 Selenia Value Console



Figure 89: Value Console Dimensions

Α	Height	189.4 cm (74.6 inches)
В	Width	101.2 cm (39.8 inches)
С	Depth	60.5 cm (23.8 inches)

2.0 Operating Environment

Weight

2.1 General Operating Conditions

Temperature Range Maximum rate of temperature change Relative Humidity Range BTU output 20 °C (68° F) to 30 °C (86° F) <10 °C/hr 10% to 80% non-condensing Typical Range 1700–2500 BTU/hr

163.3 kg (360 lb) (maximum)

3.0 Storage Environment

3.1 Tubestand

Temperature Range-25 °C (-13° F) to +60 °C (140° F)HumidityZero to 95% humidity - non-condensing (not packaged for
outdoor storage)

3.2 Image Receptor

Storage Temperature Range Maximum rate of temperature change Storage Humidity Range 10 °C (50° F) to 40 °C (104° F) <15 °C/hr 10% to 80% humidity — non-condensing (not packaged for outdoor storage)



4.0 Electrical Input

4.1 Tubestand

Mains Voltage

Mains Impedance

Maximum Power Consumption Standby Current Maximum Line Current 200/208/220/230/240 VAC nominal, true sinusoidal, (tap selectable at installation) ±10%, Single Phase, 50/60 Hz ± 5%, permanently wired Maximum line impedance not to exceed 0.20 ohms for 220/230/240 VAC, 0.16 ohms for 200/208 VAC 6.5 kVA for 5 second duration 2.0 A (maximum) 35 A for 5 seconds (momentary rating per NEC 660.2)

25 A minimum, refer to NEC 660.6

Recommended Branch Circuit Breaker Rating

4.2 Acquisition Workstation

Input Line Voltage

Input Current

Frequency Number of Phases 100/120/200/208/220/230/240 VAC nominal, true sinusoidal, (tap selectable at installation), ± 10% 8.0 A maximum @ 100/120 VAC 3.5 A maximum @ 200/208/220/230/240 VAC 50/60 Hz ±5% Single

5.0 Acquisition Workstation Technical Information

Computer Memory Disk capacity Storage Media Display Adapter Card Display

Network Interface Remote Diagnostics Graphical User Interface 2 GB RAM minimum > 60 GB Image Storage CD-RW Disks 1600 x 1200 matrix minimum 8 bit gray scale display 1600 x 1200 matrix minimum 450 cd/m2 nominal brightness

10/100 Base-T EthernetInternetX-ray exposure controlConfigurable mammographic WorkflowPatient demographicsBrightness and contrast controlMagnification screenPixel value readoutQC test toolsSystem Status MonitoringError reportingUnattended archiving and printingRated for a 0.5 mm Pb (lead) equivalence

Radiation Shield Pb equivalence



Tubestand Technical Information 6.0

6.1 C-arm

Motorized C-arm Rotation Speed	Variable speed (18° per second maximum). Rotation speed is service selectable: 50% to 100% in 5% increments. Motor Control provides soft start and dynamic braking.
C-arm Rotation Range	+195° +2° /-0° to 0° ±0.5° to -150° +0° /-2° (detent at 0°, ±90°). Rotation angle is displayed on both sides of Gantry.
Vertical Travel	74.5 cm (29.5 inches) total travel.
Source-to-Image Distance (SID)	66.0 cm (26.0 inches) ± 1.0 cm (0.4 inches)
Source-to-Image Receptor Support Device Distance	64.0 cm (25.2 inches) ± 1.0 cm (0.4 inches)
Magnification Ratio	<i>1.8 x for objects 22.5 mm above the magnification platform breast support surface</i>

Compression 6.2

Manual Compression Force	Limited to a maximum of 300 N +0/-89 N (67.4 lb +0/-20 lb) from 0° to +/-90° C-arm rotation. Not less than 169 N (38 lb) for a C-arm angle range greater than +150° and an angle less than -150°.
Motorized Compression	Functions in three operating modes, Pre-compression, Full-Range, Dual Compression, user selectable through software.
Pre-Compression Force	67 N +0/-22.3 N to 133.5 N ±22.3 N (15 lb +0/-5 lb to 30 lb ±5 lb)
Full-Range Compression Force	89 N ±22.3 N to 178 N ±22.3 N (20 lb ± 5.0 lb to 40.0 lb ± 5.0 lb)
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	Up/Down controls on both sides of C-arm and on 2-position footswitch (Motorized). Handwheel on both sides of Compression Device (Manual).
Compression Release	Manual or Automatic. Motorized Release mode controlled by push-buttons on both sides of the C-arm. User selectable automatic release mode raises Compression Device upon exposure termination. All release functions are disabled if a Localization paddle is detected.
Automatic Compression Release	Moves the compression device upward a predetermined distance (10 cm). For Magnification Mode this may be less.



compression device and image receptor alignment requirement.

Compression Down Motion Variable Speed	Service Selectable between approximately 10% through 100% of full speed.
Compression Force Display	Two LED Displays on the Compression Device show the compression force through the range of 10 lb to 67.4 lb (44.5 N to 300 N) in 1 lb (4.4 N) increments.
Compression Force Display Accuracy	±4.5 lb (±20 N)
Compression Thickness Display	Two LED Displays on Compression Device measures between 0 and 15 cm above image receptor in 0.1 cm increments. The display is visible from both sides of the patient.
Compression Thickness Accuracy	±0.5 cm for thicknesses between 0.5 cm and 15 cm
Compression Paddles	Compression paddles are transparent. The paddles are composed of polycarbonate. The paddles provide a parallel plane to the image receptor and do not deflect by more than 1 cm difference from any surface providing compression under 25 lb (111.1 N) of compression force (except for F.A.S.T. paddles.). The paddles are adjustable to provide the focal spot,

6.3 X-ray Tube: Molybdenum

Focal Spot (NEMA / IEC)	Large (0.3 mm) Nominal Small (0.1 mm) Nominal
Tube Voltage	20 kV to 39 kV
Tube Current	Large Focal Spot = 100 mA between 25 and 32 kV Small Focal Spot = 30 mA between 25 and 32 kV
Anode Rotation	180 Hz (9600 RPM minimum)
Anode Angle	Bi-angular: Large focal spot at 16°, Small focal spot at 10°. X-ray tube angle at 6° to provide 22° (Large FS) and 16° (Small FS) anode to Image Receptor plane angle.
Anode Material	Molybdenum
X-ray Window	Beryllium 0.8 mm thickness (maximum)
X-ray Tube Housing	
Over Temperature Protection Sensor	Internally connected in series with the stator common lead.
Maninum Tananatuna Tuka Hausina	55 °C (151° T)

Maximum Temperature, Tube Housing55 °C (151° F)Surface41 °C (105.8° F)Surface56 fety ClassIEC 60601-1, Class I,IEC 60601-2-28

6.3.1



Inherent Tube Filtration	0.0 mm Al equivalent
Added Filtration	Two-position filter changer mechanism to carry a 30-micron (0.03 mm) Molybdenum foil filter and a 30-micron (0.03 mm) Rhodium foil Filter. Type of filter is user selectable.
Beam Quality HVL for Mo/Mo Operation	At a given kilovolts, the measured HVL with the compression paddle in the x-ray beam is equal to or greater than the value of $kV/100+0.03$ (in units of mm of aluminum) but less than the value of $kV/100+0.12$ (in units of mm of aluminum).
Beam Quality HVL for Mo/Rh operation	At a given kilovolts, the measured HVL with the compression paddle in the x-ray beam is equal to or greater than the value of $kV/100+0.03 \text{ mm}$ Al (in units of mm of aluminum) but less than the value of $kV/100+0.19 \text{ mm}$ Al (in units of mm of aluminum).
Radiation Output	Equal to or greater than 800 mR/second for at least 3 seconds. Output is measured through the Compression Paddle, 4.5 cm above the breast support, 4.0 cm from the chest wall edge, using exposure techniques of Mo/Mo target/filter, large focal spot, 28 kV.

6.3.2 X-ray Beam Filtration and Output

6.4 X-ray Tube: Tungsten

Tube Voltage	22 kV to 39 kV
Tube Current	Large Focal Spot = 100 mA between 25 and 32 kV Small Focal Spot = 30 mA between 25 and 32 kV
Anode Rotation	180 Hz (9500 RPM minimum)
Anode Angle	Bi-angular: Large focal spot at 16°, Small focal spot at 10°. X-ray tube angle at 6° to provide 22° (Large FS) and 16° (Small FS) anode to Image Receptor plane angle.
Anode Material	Tungsten
X-ray Window	Beryllium 0.8 mm thickness (maximum)

6.4.1 X-ray Tube Housing

Over Temperature Protection Sensor	Internally connected in series with the stator common lead.
Maximum Temperature, Tube Housing Surface	55 °C (151° F)
Maximum Temperature, Tube Head Cover Surface	41 °C (105.8° F)
Safety Class	IEC 60601-1, Class I, Type B, IEC 60601-2-28



6.4.2 X-ray Beam Filtration and Output

Inherent Tube Filtration	0.0 mm Al equivalent
Added Filtration	Two-position filter changer mechanism to carry a 60-micron (0.06 mm) Rhodium foil filter and a 60-micron (0.06 mm) Silver foil Filter. Type of filter is user selectable.
Beam Quality HVL for W/Rh Operation	At a given kilovolts, the measured HVL with the compression paddle in the x-ray beam is less than the value of $kV/100 + 0.03$ (in units of mm of aluminum).
Beam Quality HVL for W/Ag operation	At a given kilovolts, the measured HVL with the compression paddle in the x-ray beam is less than the value of $kV/100 + 0.03$ (in units of mm of aluminum).
Radiation Output	Equal to or greater than 230 mR/second for at least 3 seconds. Output is measured through the Compression Paddle, 4.5 cm above the breast support, 4.0 cm from the chest wall edge, using exposure techniques of W/Rh target/filter, large focal spot, 28 kV.

6.5 X-ray Collimation

Available Collimated X-ray Fields:

Table 16: Collimation Settings

	0	
Predefin	ed Collimation Settings	
	24 x 29 cm	
	18 x 24 cm	
	15 x 15 cm	
	10 x 10 cm	
	7.0 x 8.5 cm	

Light Field Indication 6.6

Light Field Lamp

Illuminates for 30 seconds, ±5 seconds, upon pressing a Light Field switch on either side of the x-ray tubehead or by pressing a Compression Down switch. Extinguishes automatically upon

Light Field Illuminance

Light Field-to-X-ray Field Congruency

exposure initiation. A shatter shield is provided.

160 lux (minimum)—meets 21 CFR 1020.31 requirements. *Lamp is adjustable to provide alignment of the light field to the* x-ray field. Within 2% of SID

6.7 X-ray Generator

Ripple	2% or less (typical), maximum 4%
Topology	$Pulse\ width\ modulated\ High\ Frequency,\ active\ servo\ controlled$

Appendix A—System Specifications Tubestand Technical Information



6.7.1 kV /mA Range

Table 17: LFS mA setting as a function of kV

Large Focal Spot	
kV	mA
20*	75 mA
21*	80 mA
22	85 mA
23	90 mA
24	95 mA
25-32	100 mA
33	85 mA
34-35	80 mA
36-37	75 mA
38-39	70 mA
*Molybdenum Tubes only	

Table 18: SFS mA setting as a function of kV

Small Focal Spot	
kV	mA
20	20 mA
21	22 mA
22	24 mA
23	26 mA
24	28 mA
25-32	30 mA
33-34	28 mA
35-37	26 mA
38-39	24 mA



Note

All above values are for mAs values \geq 40 mAs. For mAs values < 40 mAs, the mA value is adjusted by the mA Factor listed in the table below as a function of mAs.

Table 19: mA Factor as a function of mAs

mAs	mA Factor
4-6	10%
8-10	20%
12-22	30%
24-38	60%
<u>≥</u> 40	100%



	Large Focal Spot Manual 1	nAs Range:
	TABLE 1 (default):	3 mAs through 400 mAs, 22 steps: 3, 4, 5, 6.4, 8, 10, 12.5, 16, 20, 25, 32, 40, 50, 64, 80, 100, 125, 160, 200, 250, 320, 400.
\bigwedge	Note	Lower limit is 4 mAs when grid is in field.
	TABLE 2 (user selected):	3 mAs through 400 mAs, 55 steps: 3, 4, 5, 6, 7, 8, 9, 10, 12, 14, 16, 18, 20, 22, 24, 26, 28, 30, 32.5, 35, 37.5, 40, 42.5, 45, 47.5, 50, 55, 60, 65, 70, 75, 80, 85, 90, 95, 100, 110, 120, 130, 140, 150, 160, 170, 180, 190, 200, 220, 240, 260, 280, 300, 325, 350, 375, 400.
	Note	Lower limit is 4 mAs when grid is in field.
	Note	In AEC modes, in Large Focal spot, the mA setting adjusts downward, if necessary, to achieve the calculated mAs. mA can be adjusted as low as 10 mA.
	Small Focal Spot Mag Mar	nual mAs Range:
	TABLE 1:	3 mAs through 100 mAs, 16 steps: 3, 4, 5, 6.4, 8, 10, 12.5, 16, 20, 25, 32, 40, 50, 64, 80, 100
	TABLE 2:	3 mAs through 120 mAs, 38 steps: 3, 4, 5, 6, 7, 8, 9, 10, 12, 14, 16, 18, 20, 22, 24, 26, 28, 30, 32.5, 35, 37.5, 40, 42.5, 45, 47.5, 50, 55, 60, 65, 70, 75, 80, 85, 90, 95, 100, 110, 120
6.7.2	Accuracy, Reproducibility	y, and Linearity
	Reproducibility	< 0.05 coefficient of variation for 10 consecutive exposures (21 CFR).
	Linearity	< 0.10 for adjacent mAs selections per the following: (X1-X2) is less than or equal to 0.10 (X1+X2) where X1 and X2 are average mR/mAs values for consecutive exposures (21 CFR).
	mAs Accuracy	$\pm 5\%$ or ± 2 mAs, whichever is greater, from indicated, measured from the ground side of the tube circuit.
	kV Accuracy	Within 1 kV of the indicated kV



7.0 Imaging System Technical Information

7.1 Image Receptor

Fluid ingress	No fluid from incidental spillage on the top surface of the Image Receptor seeps inside
Deflection	Does not exceed 1.0 mm at maximum compression
Active Imaging Area	24 cm x 29 cm Nominal. The active image area is marked on the digital image receptor/breast platform cover.
Image Size, Screening and Diagnostic Exams	18 x 24 cm nominal; locations: center, left, right 24 x 29 cm nominal; center location only
Image Size, Diagnostic-Spot Compression Image Size, Diagnostic-Magnification Image Pixel Size	18 x 24 cm nominal; locations: center, left, right 18 x 24 cm nominal; center location only 70 μm
Digital Image Receptor MTF -Nyquist frequency	> 0.40
DQE at 0 c/mm	50% or greater at 7.0 mR -0.0/+0.7 mR x-ray exposure
DQE at Nyquist frequency	15% or greater at 7 mR -0.0/+0.7 mR x-ray exposure
Saturation	X-ray exposure level at which image pixels are saturated is not less than 1000 mR
Dynamic Range	Linear response over at least 400:1 in x-ray exposure
Output Image	14 bit image data
Scatter Rejection	Lorad HTC™ high transmission cellular grid
Chest Wall Access	The distance from the outside edge of the Image Receptor enclosure to the Active Image Area along the chest wall is less than 5 mm.
Lateral Wall Access	The distance from the outside edge of the detector enclosure to the active detector area along the edges perpendicular to the chest wall is less than 40 mm.
ACR Phantom Score at MGD = 2mGy	At least 5 fibers, 4 specks, 4 mass
Image Preview Time	The time between completion of an x-ray exposure and availability of the Preview image: less than 20 seconds.
Transmission Limit	Within Federal Regulatory limit for screen-film mammography systems (21CFR 1020): 0.1 mR/h.

8.0 Tissue Exposure Control (TEC) Mode

The Breast Density Default Setting

Breast Density Panel Mammography Unit Status Polling Selectable via the Generator default screen on the Acquisition Workstation. Breast density defaults to the default setting at the beginning of each new study and when the exposure mode is first changed to TEC.

Update rate once every 2 seconds maximum

Between image capture cycles, Acquisition Workstation requests and displays current technique at a rate not slower than once every 2 seconds. In TEC mode, the rate increases to no slower than once per second, but not faster than once per 0.75 seconds. System response time for a single Generator Technique command is 1 second or less.

9.0 Automatic Exposure Control (AEC)

AEC Sensor	 Positionable in seven locations, centered laterally in the image receptor support device. Position #1 is located 1 cm from the chest wall. Positions are spaced in 1.7 cm increments. The AEC region can be selected automatically by processing the pre-exposure image data to find the corresponding dense portion of the breast. When the AEC region auto-selection is enabled, the detector area that is scanned is dependent on the default x-ray collimation that is linked to the compression paddle ID. Change of collimation by the operator is ignored for AEC region auto-selection. LEDs for each AEC sensor position on the compression gantry indicate the selected position. The selected AEC region can be viewed after selecting the check box. The size and available position of the manual AEC regions are indicated at the x-ray input surface of the compression paddle.
AEC Auto-Filter Mode	The filter and kV are selected using the recommended table based exposure technique based on compressed breast thickness. The exposure is terminated at an mAs value as determined by the AEC system to yield a pixel count to which the unit has been calibrated.
AEC Auto-kV Mode	The Filter is not selectable. The filter defaults to Mo if the system is configured with Moly Tube. The filter defaults to Rh if the system is configured with a Tungsten Tube. Starting kV is determined by the Gantry. mAs is determined by the Digital Detector.

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Appendix A—System Specifications Hardcopy Film Printing Devices



AEC Auto-Time Mode	The kV and filter are both user selectable. The exposure is terminated at a mAs value as determined by the AEC system to yield a pixel count to which the unit has been calibrated.
Exposure Termination	An exposure is aborted or prevented if: The predicted exposure time exceeds the x-ray tube limits. The predicted exposure time exceeds the safety backup timer setting. The predicted exposure time is less than 400 msec when the grid is in the x-ray field or less than 30 msec when the grid is out of the x-ray field. The user releases the exposure button prematurely. There is an error within the Selenia FFDM system.
AEC Reproducibility	AEC exposure reproducibility meets MQSA requirements as follows: For a 4 cm thick average breast phantom, coefficients of variation in mAs and digital values shall be less than 0.05 for four consecutive exposures of this phantom under each of the three AEC modes.
Exposure Adjustment	There is a "user-adjustable" scale factor that can be used to increase/decrease the final x-ray exposure (mAs) by 15% for each step from the default setting. A total of eight settings are available on the Acquisition Workstation, which allows the user to change the exposure from -55% of the default value to 160% of the default value.



AEC exposure adjustment is not available in all geographic regions.

10.0 Hardcopy Film Printing Devices

This section lists the relevant technical characteristics for the hardcopy film printing device.

10.1 Image Engine

Note

Film sizes	20 cm x 25 cm (8 inches x 10 inches) 25 cm x 30 cm (10 inches x 12 inches), optional
Pixel size	<u><</u> 70 μm
Grey level resolution	≥8 bits
Maximum Film Optical Density	<u>></u> 3.5



10.2 Interface

Connectivity	DICOM 3.0 compatible
Data port	Ethernet
Transfer rate	Minimum: 10 Mbits/sec
Look up Tables (LUT)	Field programmable; Linear

10.3 Printed Film

Print to fit

Patient Information

If a 24 x 29 cm image is printed on a 20 x 25 cm (8 x 10 inches) film, the printer prints the image to fit onto a 20 x 25 cm (8 x 10 inches) film without cropping the image or without dropping rows and columns of data Film meets ACR/MQSA requirements for patient

demographics. Printed Information Is User Configurable.

11.0 TechMate

СРИ

Operating System Computer Memory Disk capacity Display User Interface Dual Intel® Xeon® Processors Microsoft Windows® 2 GB RAM minimum 400 GB SATA minimum 3 Mega Pixel, 21 inch LCD, 2048 x 1536 resolution Keyboard, Trackball
Appendix B The Mobile Selenia

Digital Mammography System

This appendix provides information about Selenia systems installed in a mobile environment.

1.0 General Information

1.1 The Vertical Position Override Switch (VPOS)

The Mobile Selenia has a Vertical Position Override Switch (VPOS) on the back of the Gantry. This switch can start the Selenia if the C-arm is lower than its normal operating position after rough-road travel.



Figure 90: Location of Vertical Position Override Switch (VPOS)

1.1.1 Background

Switches located in the Gantry limit the C-arm vertical travel to a pre-defined range. When the C-arm reaches its highest or lowest point of normal travel, the switches engage to stop further motion. If the C-arm should travel past the Limit switches (in the event of switch failure or other electrical malfunction), a second switch engages that removes power from the system. A Field Engineer would normally be contacted to repair and restart the Selenia.

In a mobile environment (during transport), it is possible for the C-arm to 'creep' downward and beyond the two Limit switches. If this occurs, the Selenia would be disabled—the lower limit switch would engage, preventing the system from normal start-up. This is not a failure condition that would require a Field Engineer's attention. The Vertical Position Override Switch is used to allow the operator to start the system and raise the C-arm to within its normal working limits.



1.1.2 When to use the VPOS

This condition shows on the first startup after transport. The Acquisition Workstation starts normally, but the Gantry does not startup. After Logon, an alert message appears that communication to the Generator is lost.

1.1.3 How to Use the Vertical Position Override Switch to Restart the Gantry

- 1. Press and hold the Vertical Position Override Switch—the Gantry starts. Continue to hold the switch pressed.
- 2. After approximately 5 seconds, lift the C-arm approximately 7.5 cm (3 inches) with the normal system controls. Release the VPOS after you lift the C-arm.
- 3. At the Acquisition Workstation, accept the prompt to restart the Generator. The Gantry turns off for a few seconds, then automatically restarts.
- 4. The system is ready for normal use.

2.0 Safety Conditions and Other Precautions



Warning:

The Selenia radiation shield is not approved for mobile use and is not provided. The coach manufacturer must provide radiation shielding that meets requirements.



Mobile Specifications



Digital Mammography System



sinusoidal VAC power is supplied per the Dimensions power input specifications and loading characteristics. Intermittently, the power source must provide 65 Amps at 208 VAC for 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. The temperature and humidity inside the vehicle must be

When shore power is unavailable, mobile power sources

that provide equivalent performance may be employed. (see Mobile Specifications.) Proper system function and performance can only be ensured if continuous true

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.

For permanent archive, the Selenia must be attached to a PACS system or connected to a suitable hard-copy printer when in use.

3.0 Mobile Specifications

3.1 Shock and Vibration Limits

Vibration Limit

Shock Limit

Caution:

Caution:

Not greater than 0.35 G (2 Hz to 200 Hz), as measured at mounting point of system to coach.

Not greater than 1.0 G (¹/₂ sine pulse), as measured at mounting point of system to coach. An "air ride" coach suspension is recommended.



3.2 Coach Environment

3.2.1 Operating Environment

Temperature Range	20 °C (68° F) to 30 °C (86° F)
Maximum Rate of Temperature Change	<10 ºC/hr
Relative Humidity Range	10% to 80% non-condensing
BTU output	Typical Range 1700–2500 BTU/hr

3.2.2 Non-operating/Transit Environment

Temperature Range	15 °C (59° F) through 35 °C (95° F)
Maximum Rate of Temperature Change	<15 °C/hr
Relative Humidity Range	10% to 80% non-condensing

3.3 Electrical Input

3.3.1 Tubestand

200/208/220/230/240 VAC nominal, true sinusoidal, Mains Voltage (tap selectable at installation) ±10%, Single Phase, 50/60 Hz \pm 5%, permanently wired Mains Impedance Maximum line impedance not to exceed 0.20 ohms for 220/230/240 VAC, 0.16 ohms for 200/208 VAC Maximum Power Consumption 6.5 kVA for 5 second duration Standby Current 2.0 A (maximum) Maximum Line Current 35 A for 5 seconds (momentary rating per NEC 660.2) Recommended Branch Circuit Breaker 25 A minimum, refer to NEC 660.6 Rating

3.3.2 Acquisition Workstation

Input Line Voltage	100/120/200/208/220/230/240 VAC nominal, true sinusoidal,
	(tap selectable at installation), $\pm 10\%$
Input Current	8.0 A maximum @ 100/120 VAC
	3.5 A maximum @ 200/208/220/230/240 VAC
Frequency	50/60 Hz ±5%
Number of Phases	Single

4.0 How to Prepare the System for Transport

Perform these steps before you transport the Mobile System:

- 1. Rotate the C-arm to 0 degrees (CC position).
- 2. Lower the C-arm to its lowest position.
- 3. Remove all power from the system.



4. Remove all system accessories (like the Face Shield, and Compression Paddles) before transport. Store all accessories correctly to make sure the components are not damaged.

5.0 How to Test the System Integrity After Transport

Instructions: Photocopy this checklist for use after the system is relocated.

Name_____ Date____

System Serial Number_____

Table 20: Integrity Checklist		
Evaluated	Control Functions	Reference
Mechanical System Tests		
	Compression Up/Down	Controls and Functional Tests on page 25
	Compression Release	Controls and Functional Tests on page 25
	C-arm Rotation	Controls and Functional Tests on page 25
	C-arm Up/down	Controls and Functional Tests on page 25
	Collimator Override	Controls and Functional Tests on page 25
	Light Field Lamp	Controls and Functional Tests on page 25
	Smart Paddle System	Controls and Functional Tests on page 25
	Emergency Off Switches	Emergency Off Switches on page 13
Quality Control Tests		
	Artifact Evaluation	Selenia QC Manual
	SNR/CNR Measurement	Selenia QC Manual
	Phantom Image Evaluation	Selenia QC Manual
	Compression Thickness	Selenia QC Manual

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