



R2[™] DMax

User Manual Software Versions 8.6–8.7

PN MAN-01042 Rev 001



Screen-Film Mammography

R2[™] DMax User Manual

Software Versions 8.6-8.7

PN MAN-01042 Rev 001



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For support in Europe, South America, or Asia, contact your local dealer or distributor.

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Part 1: Introduction

- ► 1.1. Resources Available
- 1.2. How to Use this Manual
- ► 1.3. Warnings and Cautions

This manual provides instructions for the operation and maintenance of the R2 DMax system in the clinical environment. R2 DMax is used in radiology clinics to scan and digitize mammography films. The two options available for the R2 DMax system are:

- DigitalNow[™], used to produce archivable DICOM For Presentation mammography images from the scanned films.
- ImageChecker[®], computer-aided detection (CAD) software used to assist in the evaluation of regions of the breast on routine screening and diagnostic mammograms.

1.1. Resources Available

In addition to this manual, the following resources are available to assist you when working with the R2 DMax system.

- **R2 Member Center**: This website provides quick access to electronic (pdf) versions of Hologic manuals and training materials. You can find the R2 Member Center by visiting the Hologic website (www.hologic.com). See the R2U flyer that accompanies this product for instructions on accessing the R2 Member Center.
- **Manuals**: The manuals for the R2 DMax system and its applications are listed below:
 - Understanding R2 ImageChecker
 - R2 DMax User Manual
 - R2 DMax/DM/DX/LS Service Manual
 - R2 DMax/DM/DX/LS Release Notes
 - R2 DMax/DM/DX/LS DICOM Conformance Statement

You can obtain additional copies of printed manuals through your Hologic Account Manager. The DICOM Conformance Statement and Cybersecurity Best Practices documents are available at www.hologic.com.

- **Training**: The Hologic Applications team is available to train your staff, should you feel they need additional training. To purchase additional personalized instruction, contact your Hologic Account Manager.
- Technical Support and Service: For support in North America contact:

Toll Free:	+1.866.243.2533 (+1.866.CHECKED)
Email:	r2support@hologic.com
Hours:	Monday – Friday, 6:00 AM – 5:00 PM, PT (GMT –8:00)
Website:	www.hologic.com

For support in Europe, South America, or Asia, contact your local dealer or distributor.

1.2. How to Use this Manual

This manual was written to ensure safe and proper use of the system. Before use, read this manual carefully in order to realize the full capabilities of the system. If something is unclear during daily use or if a problem occurs, please refer to this manual.

Manual Organization

This manual is organized as follows:

- Part 1: Introduction provides a listing of support resources available from Hologic, information about using this manual, and safety information.
- Part 2: Description provides background information on the system, component and feature descriptions, and an overview of operation.
- Part 3: Studies provides instructions for working with films, scanning them, displaying study results, verifying and labeling images, and searching for a study.
- Part 4: Alerts provides a listing of alert conditions with instructions for resolving the alerts and diagnosing and solving common problems.
- Part 5: Controls provides instructions for using the system configuration and maintenance options that appear on the Controls screen.
- Part 6: Maintenance provides instructions for cleaning the system and running the weekly tests.
- ▶ **Part 7**: **Using DigitalNow and the R2 Patient ID System** provides information about archiving images with the R2 DigitalNow[™] feature and the R2 Patient ID system.

Conventions Used in this Manual

This manual uses the following conventions to provide technical and safety information of special interest.

A Note: Background information provided to clarify a particular step or procedure.

Important: An instruction provided to ensure correct results and optimal performance.

- A Caution: An instruction that, if not followed, can result in damage to the system.
- **Warning!** An instruction that, if not followed, can result in a hazardous condition.

Also note the following conventions:

- *Italic* and **boldface** typefaces are used for emphasis. Examples: 'Press *carefully* to engage,' '**Do not** press down.'
- When this manual directs you to touch a button on the touch-screen monitor, the name of the button is shown in boldface type; for example, 'Touch **Start**'.
- If you are reading this manual online, click on the **blue hyperlinked text** to jump to the referenced section.

1.3. Warnings and Cautions

Note: For radiological warnings and precautions associated with R2 ImageChecker, see the manual, Understanding R2 ImageChecker.

Warnings: System Operation



- Use the system only outside the patient environment. The system is rated for use only in an office environment.
- Remove all potentially obstructive jewelry and clothing before loading films in the processing unit, to minimize the possibility of injury due to moving parts, or damage to the system.



• The hand-held barcode reader is a Class II laser device that emits laser light. Do not stare into beam.



- The symbol next to the power connector indicates a potential shock hazard. Ensure that the system is connected to a power receptacle that is properly grounded and provides voltage and current within the specifications of the system to reduce the likelihood of electrical shock or fire hazard.
- There is a small risk of electrostatic discharge (static electricity). It is mostly a nuisance and not likely to cause harm. However, a shock can cause accidental injury, if, for example, you suddenly withdraw your arm and hit it against something. To minimize possibility of shock, maintain greater than 40% relative humidity in the room where the unit is kept, use shoes with natural-material soles, place equipment on nonpolymer floors such as concrete, wood, or floors treated with static dissipative, and/or routinely touch metal with a metal object, such as a key, to painlessly discharge the charge from your body.
- Do not place liquid containers on the device. In the event of a spill, shut down power to all components prior to cleaning to minimize the possibility of electrical shock. Do not operate the device if internal components are exposed to liquid. Contact your service representative.

Warnings: Maintenance



- Before cleaning the computer, always shut down the system according to the shutdown procedures in this user manual and disconnect the power cord to prevent the possibility of electrical shock. Never use alcohol, benzene, thinner, or other flammable cleaning agents.
- Never use a two-prong plug or extension cord to connect primary power to the system. Use of a two-prong adapter disconnects the utility ground and creates a severe shock hazard.
- The R2 scanners are heavy. It is recommended that two people participate when moving equipment.

Cautions: System Operation



- Operators should review this user manual and receive training before using the system.
- To ensure optimal system performance, perform the system verification tests and scanner maintenance procedures as instructed in this user manual.
- To ensure proper system operation, use only separator sheets and barcodes provided by Hologic.
- To properly identify patients in the archive system, be sure to enter the patient information and the barcode from the separator sheet into the patient identification system correctly.
- For sites with multiple processing systems: When entering data into the R2 Patient ID web page, check the name of the processing unit on the web page to ensure that you are entering the patient information into the processing unit that you will use to scan the case.
- For the display images to correspond correctly to the film position at the display unit, be sure to orient and order the films correctly when scanning, as described in this user manual.
- Ensure that the case retrieved from the archive to be used for comparison is from the same patient as the current case under review. It is recommended that you use the patient flash for this confirmation.
- Use only standard mammographic film: 18×24 cm or 24×30 cm.
- Do not attempt to scan films that have labels or tape with edges within 1 mm of the edge of the film, or labels that are not pressed flat, or have curled corners, as they may jam.
- Do not attempt to scan bent, damaged, or curled films, as they may jam. Be sure to remove paper clips or other foreign materials from the films.
- Always shut down the system according to the procedures provided in this user manual. Failure to shut the system down properly can cause loss of data or damage to the computer operating system.
- This equipment has been tested and found to comply with the limits for a Class A digital device, pursuant to Part 15 of the FCC Rules. These limits are designed to provide reasonable protection against harmful interference when the equipment is operated in a commercial environment. This equipment generates, uses, and can radiate radio frequency energy and, if not installed and used in accordance with the instruction manual, may cause harmful interference to radio communications. Operation of this equipment in a residential area is likely to cause harmful interference at her or his own expense.

▲ **Note:** The device design and mode of operation are consistent with current standard mammography clinical practices, as governed by the requirements of the Mammography Quality Standards Act (MQSA) of 1992. Users are cautioned to comply with the MQSA or the appropriate national standards, when implementing R2 systems in clinical protocols.

Cautions: Installation and Maintenance

This product contains no user-serviceable parts. To prevent damage to the system:



- When necessary, shut down the system according to the procedures recommended in this user manual.
- Do not attempt to install or repair the R2 system. Only trained personnel, authorized by Hologic, are qualified to install or repair the system.

• Maintain equipment in a well-ventilated, air-conditioned environment.

Part 2: Description

- ▶ 2.1. Overview
- > 2.2. System Components
- ▶ 2.3. Overview of Operation
- 2.4. Licensed Features
- ▶ 2.5. Other Features
- ► 2.6. System Inputs and Outputs

Part 2 provides an overview of the system, a description of system components, an overview of system operation and features, and information about CAD processing.

2.1. Overview

The R2 DMax system is a full-featured mammography film-scanning platform designed to:

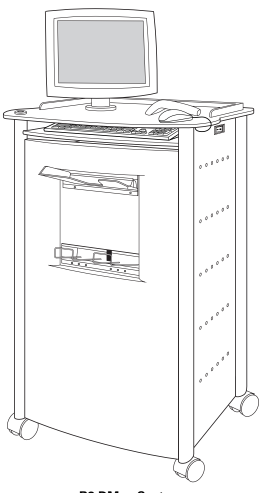
- Scan mammography X-ray films and convert them into digital image files.
- Allow users to review the scanned images, and to reorient, label, and reprocess the images as needed.
- Interface with a patient database to retrieve patient information, which can then be integrated with the study images.
- Use Hologic's DigitalNow[™] feature to transmit the digital mammography image files to an output device such as a PACS for archiving.
- Use Hologic's ImageChecker[®] CAD algorithm to analyze the images in order to mark regions of interest on routine screening and diagnostic mammograms, and transmit the ImageChecker results to an output device such as the R2 CheckMate[™] Ultra display unit where they can be viewed.

 \triangle **Note:** This manual uses the following terms to describe mammography images and media:

- Case: a series of X-ray films associated with a particular patient.
- Stack: a group of cases including a separator sheet on top of each case.
- Study: a series of digitized images associated with a particular patient.

The R2 DMax system consists of a touch-screen monitor, a handheld barcode scanner, a keyboard, a film scanner, and a computer that performs the DigitalNow and ImageChecker processing, all housed in a movable cart.

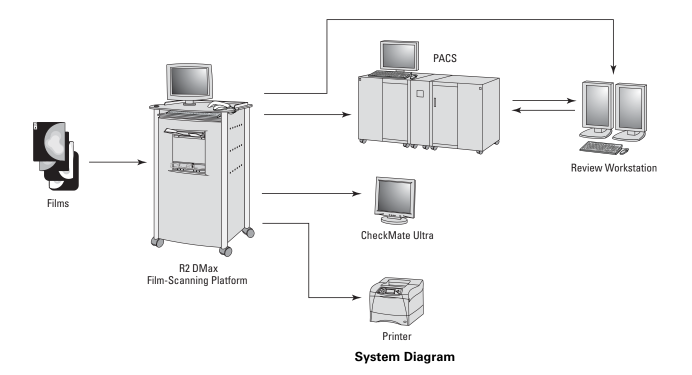
When using the system, the technologist loads a stack of cases into the film scanner. Each case can include from 1 to 24 X-ray mammography films. The technologist inserts a barcoded R2 separator sheet before each case. When the technologist touches the Start button on the touch screen, the system begins scanning the films, converting each film into a digital image, and starting a new case each time it encounters a separator sheet. After each case is scanned, the technologist has the option to review the case and verify that the films were scanned correctly. Then the system sends the images or CAD results to output devices such as display units, PACS or diagnostic review workstations.



R2 DMax System

If the system is configured with the DigitalNow license, the R2 processing unit can send images to virtually any DICOM-conformant workstation and/or archive, allowing images to be displayed, stored, and managed universally.

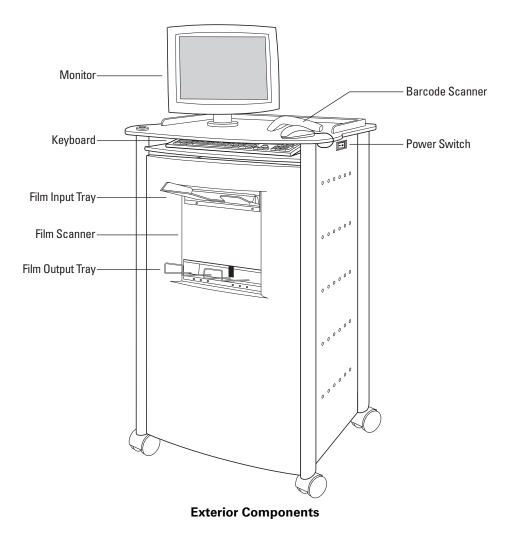
If the system is configured with the ImageChecker CAD license, it analyzes the scanned images with a proprietary algorithm (i.e., a set of criteria) that identifies regions of interest, which can include clusters of bright spots (suggestive of calcification clusters), and dense regions with or without radiating lines (suggestive of masses or architectural distortions). The system generates CAD marks identifying the regions of interest. Images with CAD results can be sent to an R2 display unit, PACS, review workstation, or printer. For more information about the CAD marks and the ImageChecker algorithm, see *Understanding ImageChecker 8.5-8.7*.



2.2. System Components

This section describes the major system components.

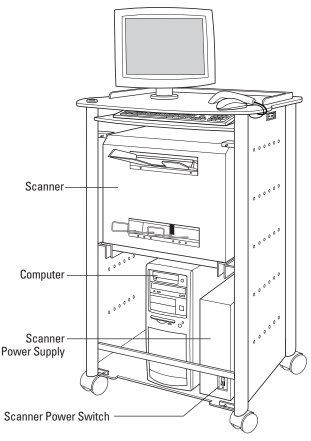
- **Touch-Screen Monitor**: Use to control and monitor the system (start scanning, select studies, view system status, etc.).
- **Scanner**: Use to convert the mammogram films to digital images. You stack the films and place them in the input tray. After the scanner digitizes the images, it stacks the films in the same order in the output tray. The DMax system scans each film in about 25 seconds.
- **Barcode Scanner**: Use to select a study to display by scanning a barcoded patient ID or the barcode on a separator sheet.
- **Keyboard**: Use to type search criteria when searching for studies, and to enter patient information.
- **Mouse**: Use to select screen objects as an alternative to the touch screen.



- **Separator Sheets**: The separator sheet provides a unique R2 ID number (a 10digit barcode) that the system reads and uses to identify each case and associate the scanned images with the patient's films. Each case fed into the R2 system has its own separator sheet.
- **Test Sheets and Films**: Use for the weekly tests to ensure optimal system performance. For more information, see '6.4. Running the Weekly Tests'.
- Lead Markers: Read by the R2 system to identify standard views automatically. For more information on lead markers, see 'R2-Supported Lead Markers' on page 24.

The following components are located behind the front cover. You'll only need to access these components for certain maintenance procedures.

- Computer: Processes the scanned films and images.
- Scanner Power Supply: Provides power to the scanner.



Interior Components

2.3. Overview of Operation

The R2 GreenLight[™] user interface is designed to guide you as you use the system. The system's touch-screen interface provides quick access to all system functions. The user interface features three main screens – **Studies**, **Alerts**, and **Controls** – plus a common area with film scanning and search options at the right. You can go to each screen by touching one of the tabs at the bottom of the display. Onscreen messages appear when appropriate and provide helpful instructions.

Studies Screen Overview

When scanning films, you will use the **Studies** screen, which is shown below:

Reg ImageChecker®	R2 Patient ID	Search 🔎
R2 ID: 0000022609 Receive Time: Oct 10 15:26 Status: Processing Images R CC L CC R MLO L MLO Patient Information Receive Time For ImageChecker® M1000 SYSTEM R23075cl.img REST FILM FOR ImageChecker® M1000 SYSTEM R23075cl.img REST FILM FOR ImageChecker® M1000 SYSTEM R23075cl.img R250 SYSTEM CIRCLE, LOS ALTOS, CA 94022 Patient ID: 123456test		Auto Screening Uni Right Uni Left Screening Imp Any
Name: Tester^Teresa Accession:		Status Ready
Studies 🔥 Alerts	Controls	Start

The following items appear on the Studies screen:

- **Study icons** each represent one study. Scroll the study icon list to view additional studies. To display the results of any completed study, simply touch the icon for the study. The appearance of the study icon indicates the status of the case; the various icons that may appear are shown in the table on page 14.
- **Study Information** appears for each case of films as they are scanned and processed. The Study Information includes the R2 ID, Receive Time (the time scanning began for the study), and current Status, as well as thumbnail images of the films and patient information discussed below.
- Thumbnail **Images** appear as they are scanned and can be viewed at any time after the system scans the films.
- **Patient Information** includes the patient flash from the scanned film and other data imported from a patient database.

The Study icon at the top represents the study currently being processed or the study most recently processed. The Study icons below represent studies processed earlier.

The white triangle indicates the study icon linked to the study information currently on-screen.

- The **R2 Patient ID** button appears only if your system is configured to use the R2 Patient ID system. Click the button to access the R2 Patient ID database. For more information see '7.3. Using the R2 Patient ID System'.
- The **Search** field allows you to type patient data or a barcode number and quickly find the patient record. You can also activate the Search field with the handheld barcode scanner.
- The **Scanning Protocol** list allows you to select a scanning protocol that matches the number and types of films in your cases. Includes R2 standard scanning protocols and any custom protocols created on site.
- The **Start button** is used to begin scanning and processing a stack of films. When scanning begins, the button changes to Stop.
- The **Status** field, which appears above the Start button, indicates the scanner status, for example, '**Ready**', '**Scanning**', etc.

Scanning and Processing

To scan a case, you must first create a stack of films. The case at the top of the stack is scanned first. You select a scanning protocol based on the type of cases you will be scanning (see '3.1. Working with Films' for more information).

To scan films you simply:

- Arrange the films with a separator sheet on top of each case.
- Organize films and separator sheets in a stack.
- Place the stack in the film input tray.
- Touch the **Start** button.

The scanner picks up the first separator sheet followed by the first film in the stack. A new study icon appears at the top of the list, and a white circle on the icon flashes to indicate that the system is scanning or processing the films. Each separator sheet includes a barcode that the system uses to organize the results. When the system scans the barcode on the separator sheet, the barcode number appears on the Studies screen as the 'R2 ID'. As the films are scanned, the system displays thumbnail images, the patient flash, and any patient information linked to the R2 ID.

When the system finishes processing the case, the Status field changes to 'Completed'. A white arrow on the study icon flashes to indicate that the system is sending study results to the display unit, PACS/workstation, or printer on your network.

▲ **Note:** Before the study results are sent to the display or archive device, you may be required to 'verify' that the system scanned the films correctly. This option is configured on the Controls Connectivity screen. For more information, see '**5.2. Connectivity Options**'.

Study Icon	Status		
(blinking circle)	Case films are being scanned or processed.		
(D blinking arrow)	Case results are being sent to a display or archive device.		
(blinking question mark)	Scanned images for the case require verification.		
(blinking exclamation mark)	Case films are being scanned or processed, but a non-fatal fault condition has occurred.		
() blinking arrow)	Case results are being transmitted, but a non- fatal fault condition has occurred.		
(1) (blinking question mark)	Scanned images for the case require verification, and a non-fatal fault condition has occurred.		
	Case is complete, results have been sent.		
	Case is complete, but a fault condition has occurred that may require user intervention.		
Alert	Case has failed. A fault condition has occurred that requires user intervention.		

The appearance of the study icon indicates the status of the case:

For more about using the Studies screen, see 'Part 3: Studies'.

System Alerts

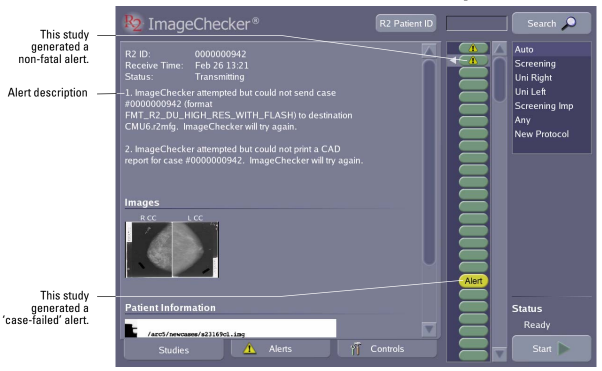
If a problem occurs during scanning or processing a case, or while transmitting case results, the system issues an alert. There are three general types of alerts:

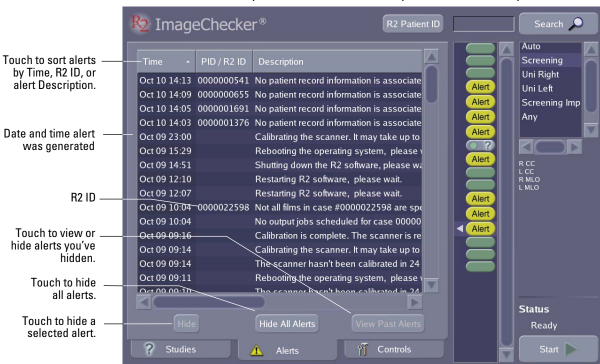
Alert Icon	Status
	A non-fatal fault condition has occurred. This alert appears on the case study icons (see previous table).
	The software has processed the images successfully, but a fault condition has occurred that may require user intervention.
Alert	Case has failed, results have not been sent.

A description of the alert condition appears for the selected study in the area of the screen below the study's R2 ID.

The outcome of any alert condition depends upon the nature of the problem the system has encountered. In the example shown below, the system is experiencing problems as it attempts to send case results to a CheckMate Ultra display unit and to a printer. If after repeated attempts it cannot transmit the case results, the system may issue a 'case-failed' alert and change the study icon accordingly.

The system produces alerts for a variety of reasons. When necessary, the alert description provides instructions for resolving the problem. In some cases you will want to refer to 'Part 4: Alerts' for more information about a particular alert.





When an alert is issued, the yellow alert icon (Δ) appears on the **Alerts** tab at the bottom of the screen. If you touch the **Alerts** tab, you can see recent system alerts.

When reviewing alerts, you can use the Alerts screen to sort alerts by time issued, R2 ID, or by description. You can also hide one or more alerts.

Controls Screen

The **Controls** screen is used to configure system settings to meet your particular needs, run maintenance procedures, and display system information.

	R2 Imag	eChecker®		R2 Patient ID		Search 🔎
Touch a screen name to display its contents.	About Maintenance Off Outputs Patient ID Performance Printing Scanning Service	Scanning Protoco Auto Screening Uni Right Uni Left Screening Imp Any Rcc 24 manual	New Edit Remove	R CC R MLO L MLO R CC ID L CC ID L CC ID L MLO ID L MLO ID	Alert Alert Alert Alert Alert Alert Alert Alert	Auto Screening Uni Right Uni Left Screening Imp Any
	? Studie	s <u>À</u> A	Move Up Move Down letts	ŶĴ Controls	Ĭ	Status Ready Start

For details about the settings, utilities, and information on the **Controls** screen, see **'Part 5: Controls'**.

2.4. Licensed Features

The system features are determined by the licensed options selected by each site, allowing you to invest only in the features you want, as well as add features at a later date. The following features can be obtained by purchasing the appropriate license:

- DigitalNow DICOM Export
- ImageChecker Screen-Film CAD

These features are described further in the following sections.

DigitalNow DICOM Export

With DigitalNow, R2 film-scanning platforms send digitized film images to a DICOM-conformant archiving system for future review. Sites using full-field digital mammography (FFDM) systems can scan prior analog studies and compare those priors with the current digital study on a softcopy review workstation. Sites planning to move to digital in the future can plan ahead and send the digitized images to an archive system now, and retrieve them for comparison later to future digital studies.

R2 systems configured with DigitalNow utilize the separator sheet barcode ('R2 ID') for identifying the patient associated with the digitized film images. This allows for the proper identification, storage, and retrieval of digitized film images in the archive system.

The system can be configured to produce 50- or 100-micron output images for any output device. For digitized film images, the processing units perform dynamic window leveling on each image, as well as blacken image areas that lie outside the film edges.

ImageChecker Screen-Film CAD

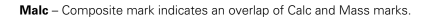
The ImageChecker license provides Hologic's CAD algorithm, which is used to identify and mark regions of interest on routine screening and diagnostic mammograms to bring them to the attention of the radiologist after the initial reading has been completed.

ImageChecker provides three types of RightOn[™] CAD marks (Mass, Calc, and Malc) that can appear in the results. Each mark identifies a region of interest for the radiologist to review.



Calc – Marks regions suggestive of calcifications.

Mass - Marks regions suggestive of masses or architectural distortions.



The CAD marks can be augmented using the EmphaSize[™] and PeerView[™] features. For more information, see the manual, *Understanding ImageChecker 8.5–8.7*.

2.5. Other Features

In addition to the licensed features, the R2 DMax system includes the following standard features, which are described further below:

- Automatic Film Marker Identification
- Patient Database Connectivity
- Reimbursement Barcode Report

Automatic Film Marker Identification

If your four, standard-view films were processed using R2-supported lead markers, and the markers are well-placed, you can scan the films using Auto film detection mode. This means your cases can consist of any number of films (up to four), and you can place them in the scanner in any orientation (e.g., rotated or emulsion side up or down) and in any order.

This timesaving feature enables sites to scan as many cases as possible in as short a time as possible. The lead markers must be a supported type, they must not be attached with putty or other opaque substances, and they must be placed properly (as per MQSA standards). R2-suported lead markers are available from Hologic, Livingston Products, Siemens, Techno-Aide, or All-Craft Wellman. For the supported marker types, see 'R2-Supported Lead Markers' on page 24.

Patient Database Connectivity

When scanning films for digital archiving, sites need the ability to retrieve patient information in order to add the required identification information to the resulting Digital Mammography X-Ray images. Sites can retrieve patient data by querying a PACS or RIS (Radiology Information System). As an alternative, sites can enter patient data manually by using the R2 Patient ID feature.

The R2 software retrieves patient information from the patient database – such as the patient name, medical record number, and birth date – and links this information to the patient images. Patient information can be printed on the CAD Results Report (see page 44), and on the Reimbursement Barcode Report (see page 19).

Each case fed into the R2 film-scanning system must have its own *separator sheet* with an R2 ID barcode, which the system uses to associate the patient films with the images in the system. To link the patient database information to R2 images, you use a keyboard or barcode scanner to enter the R2 ID from the separator sheet.

Reimbursement Barcode Report

The Reimbursement Barcode Report is a listing of all separator sheet barcode numbers scanned in a specific time period. You can schedule it to run monthly, weekly, or on demand for a specified date range.

The report lists the date, the cases processed on that date, and the time the case was processed. This can be correlated to the patient using the R2 barcode number written in the patient file, thereby completing the reimbursement audit trail.

The report lists only cases that were successfully processed. For an example printout, see 'Printing a Reimbursement Barcode Report' on page 70.

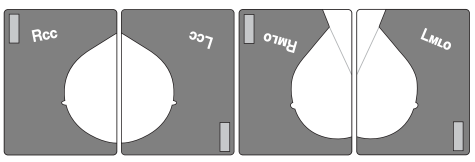
2.6. System Inputs and Outputs

The R2 processing unit supports up to 24 images per case. All DigitalNow scanned film images can be viewed at a digital review workstation and/or archived on a PACS. Other devices, such as the CheckMate Ultra display unit, can display up to four views per case, if the views are supported by the display unit.

Supported Views

The R2 processing unit allows you to scan and digitize all standard mammography views, and then label them appropriately. The four 'standard views' processed automatically are:

- RCC Right Cranio-Caudal
- RMLO Right Medio-Lateral Oblique
- LCC Left Cranio-Caudal
- LMLO Left Medio-Lateral Oblique



The Four Standard Views

The following table lists all supported views and the output devices that can accept these views.

Supported Views		DICOM Workstation,	Display Unit, Printer
Standard View	View Label	PACS	(or CAD SC)
Cranio-Caudal	СС	•	•
Medio-Lateral Oblique	MLO	•	•
Equivalent View			
Medio-Lateral	ML	•	•
Exaggerated Cranio-Caudal	XCC	•	•
Cranio-Caudal Exaggerated Laterally	XCCL	•	•
Cranio-Caudal Exaggerated Medially	XCCM	•	•
Reversed Equivalent View			
Latero-Medial	LM	•	_
Latero-Medial Oblique	LMO	•	
Cranio-Caudal From Below	FB	•	
Superolateral to Inferomedial Oblique	SIO	•	
	-		

• = Output supported to this device — = Output not supported to this device (or format)

In the preceding table, 'Display Unit' refers to a CheckMate Ultra display unit or Mammolux motorized viewer. These devices (and the R2 postscript printer) can display (or print) no more than four views per case.

If your system has been configured with the ImageChecker license, the R2 processing unit can produce ImageChecker CAD results for all supported views. However, ImageChecker results for certain views may not be available with some output devices and formats.

View Modifiers

View modifiers can be added to describe any supported view. The following table lists the supported DICOM view modifiers.

View Modifier	Label	View Modified
Axillary Tail	AT	MLO
Cleavage	CV	CC
Partial View*	(none)	Any
Rolled Lateral	RL	Any
Rolled Medial	RM	Any
Rolled Inferior	RI	Any
Rolled Superior	RS	Any
Magnification	M	Any
Spot Compression	S	Any
Implant Displaced**	ID	Any
Implant Present*	(none)	Any
Tangential	TAN	Any

*The Partial View and Implant Present modifier labels are NOT added to the view description, but are present in the DICOM header of the DigitalNow image.

** Implant-displaced views with a maximum of 2.5 cm (1 in) of implant imaged.

▲ **Important!** Certain view modifiers prevent the system from performing CAD processing. For more information, see the manual, Understanding ImageChecker 8.5–8.7.

Results Output Options

The R2 processing unit can send results with any of the following formats to a PACS or review workstation.

• **DigitalNow** is a traditional DICOM Digital Mammography X-Ray For Presentation image where the image is comprised of Natural Pixel Values, the values actually created by the film scanner. (DigitalNow images do not include CAD results.) DigitalNow results can be created as 50- or 100-micron image files. The file sizes can vary depend upon the film size and resolution.

Film Size	50µ Image	100µ Image
18 × 24 cm	35 MB	8.6 MB
24 × 30 cm	50 MB	14.4 MB

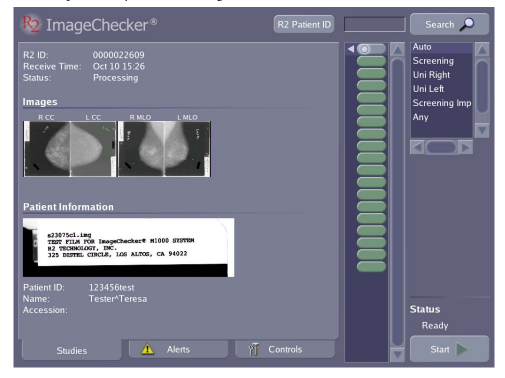
- **R2 Auto** is the format used to display the four screening views with CAD results on a CheckMate Ultra display unit or Mammolux motorized viewer. With this format, the CAD results appear over a montage of low-resolution images. Normally, the four screening views appear, although it is also possible to produce CAD results from films that show Equivalent Views.
- **Mammography CAD SC** is a Secondary Capture image that provides the same lowresolution images and CAD results as appear in the R2 Auto format. This format is useful for PACS that do not support CAD SR.
- **Postscript** files, suitable for printing on a postscript printer, provide the same low-resolution images and CAD results as appear in the R2 Auto format. For an example, see the CAD Results Report on page 44.
- Mammography CAD SR 8.x results are derived from the latest version of the ImageChecker algorithm. This is the DICOM standard format for CAD results. Mammography CAD SR objects are generally useful only when viewed with the images on an advanced diagnostic review workstation.
- Mammography CAD SR 5.x provides backward compatibility with previous releases of ImageChecker 5.x products. The 5.x format provides RightOn Calc and Mass marks but not EmphaSize or PeerView.

Part 3: Studies

- ► 3.1. Working with Films
- ▶ 3.2. Preparing Cases to Be Scanned
- ▶ 3.3. Scanning with the Auto Protocol
- ► 3.4. Scanning with a Manual Protocol
- ► 3.5. Scanning with the Any Protocol
- ► 3.6. Adding Cases to the Stack
- ► 3.7. Stopping Scanning and Removing a Case
- ► 3.8. Viewing Study Information
- ► 3.9. Verifying Scanned Cases
- 3.10. Correcting Scanned Images
- ► 3.11. Printing CAD Results
- ► 3.12. Searching for a Study
- ► 3.13. Using Queue Manager

Part 3 provides instructions for working with films, scanning cases, working with scanned images, displaying study results, and searching for a study.

For most operations you will be using the Studies screen, which is shown below:



3.1. Working with Films

To use the system successfully, please read and understand the film requirements and modes used to scan the films.

Film Requirements

All films must meet the following requirements:

- Original films only no copies
- No bent, damaged, or curled films
- 18×24 cm or 24×30 cm standard full-field views
- No sticky residue, wax or grease pencil marks on films
- No labels over the edges of films, or curled labels
- MQSA or other required standards

Because the scanner uses suction cups to lift each film into the scanner, anything that interferes with the suction cups, such as a patient label placed improperly or a wax pen mark, may cause the scanner to drop films.

R2-Supported Lead Markers

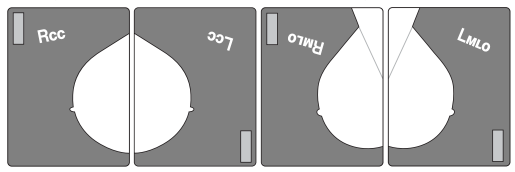
The R2 film-scanning platform can scan any film-based mammograms that meet the requirements listed above. However, when you use R2-supported lead markers with the four standard views, there is an additional degree of convenience: you can load the films in any orientation and in any order. (For more information, see 'Scanning Protocols' on page 26.)

R2-supported lead markers are available from Livingston Products, Siemens, Techno-Aide, All-Craft Wellman, and Hologic. An illustration of the supported type from each company is shown below.



Lead Marker Placement

Correct placement of the lead markers is essential for optimal results. The following examples show lead markers that are easy for the system to read, and that are not touching the patient ID label, the breast tissue, or the film edge.



Correct Auto Protocol Lead Marker Placement

The following lead markers are incorrectly placed, as explained below:



Incorrect Auto Protocol Lead Marker Placement

Separator Sheets

The R2 DMax system is shipped with 500 separator sheets. Run through all sheets on a rotating basis to ensure that patient barcode numbers are not reused before the images are deleted from the system. Do not discard the separator sheets; they are reusable.

The separator sheets are sturdy and able to withstand a great deal of use. However, be sure to store them flat, do not write on them with anything greasy or waxy, do not put tape on the edges, and keep them clean. Most importantly, do not obscure the barcode or the barcode number.

To order additional separator sheets, contact your Hologic Account Manager or Technical Support.

Important! It is critical that ImageChecker users keep the separator sheet with the films until after the radiologist has read the films and CAD results.

Scanning Protocols

The system includes six preconfigured scanning protocols: Auto, Screening, Uni Right, Uni Left, Screening Imp, and Any. You can also create your own site-specific scanning protocols, see '5.7. Scanning Protocols'.

The default (i.e., preselected) scanning protocol is Auto. You can change the default scanning protocol using the Controls Scanning screen (as described on page 73).

Auto Protocol

When using the Auto protocol, the system recognizes the lead markers and uses them to properly identify, orient, and process films with the four standard views.

You can load the films for each case in any order (e.g., RMLO, LCC, RCC, LMLO) and in any orientation (e.g., rotated 180°, emulsion side up or down). The system automatically scans all films in the stack. Each case in the stack can have any number of films up to four. The system automatically adjusts if there are fewer than four films. However, in Auto mode, the system will not process duplicate views within the same case (e.g., two RCC films).

After you touch the Start button, the system processes all cases in the stack (up to 100 films and separator sheets in all).

Manual Protocols

R2 provides four preconfigured Manual scanning protocols: Screening, Uni Right, Uni Left and Screening Imp. Each protocol requires you to load films with specific views in a predefined sequence. For the views used with each predefined scanning protocol, see '3.4. Scanning with a Manual Protocol'. You can also define custom Manual scanning protocols for your site, or alter the included protocols as needed.

When using a Manual scanning protocol, the system does not recognize the lead markers. Instead, it relies on you loading the films:

- With the views specified by the selected scanning protocol.
- In the correct order defined by the selected protocol.
- In the correct orientation emulsion (dull) side down, landscape for small films, portrait for large films.
- With the patient flash in the correct orientation.

The system also assumes that each study includes the same film views (as defined by the selected protocol). If you are using a Manual protocol (because, for example, you don't use R2-supported lead markers) and the cases to be scanned include different numbers of films or different views, create stacks of like cases and scan them separately using the appropriate scanning protocol.

When using a Manual scanning protocol, as with the Auto protocol, the system automatically scans all cases in the stack. When scanning is complete, the system switches back to the default scanning protocol, as configured on the Controls Scanning screen.

Any Protocol

This special scanning protocol allows you to scan any number of films (up to 24 per case) in any order and orientation. When using the Any protocol, you must identify all views manually after the case is scanned.

Choosing a Scanning Protocol

The scanning protocol you choose depends on the views present in the case, and the condition of the films. In general, you will want to select a scanning protocol that suits about 90% of the cases in the stack, which will minimize the adjustments that you have to make after scanning. However, sometimes it may be faster simply to scan with the Any protocol and correct the images afterwards.

▲ Important! Many scanning errors, including scanning films in the wrong order or scanning a film in the wrong orientation, can be overcome using the GreenLight Image Correction feature. When in doubt, scan the films as best you can and fix any problems after scanning. However, note that if multiple patients are scanned under one separator sheet, this error cannot be corrected via Greenlight; the cases must be rescanned to ensure that each patient has a different separator sheet.

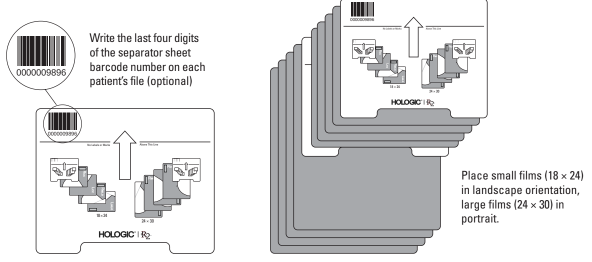
The following table provides basic guidelines for selecting a scanning protocol.

If your films	Then see
 Are the four standard screening views (RCC, LCC, RMLO, LMLO) and Were taken using an R2-supported lead marker (those shown on page 24) and Have well-placed lead markers 	3.3. Scanning with the Auto Protocol
 In all cases include the same number and type of views, and: Have unsupported, missing, or bad lead markers, or Have incorrectly placed lead markers (e.g., over a breast image, under an ID label, or only partly on film, as shown on page 25), or Have non-standard views or views with modifiers, or Have more than four views per case 	3.4. Scanning with a Manual Protocol
 Include views that do not conform to any available scanning protocols 	3.5. Scanning with the Any Protocol

3.2. Preparing Cases to Be Scanned

Use this procedure to create a stack of cases to be scanned.

1 Place a separator sheet on top of each film case.



Creating a Stack of Cases

Important! It is critical that ImageChecker users keep the separator sheet with the films until after the radiologist has read the films and CAD results.

- 2 Create a neat stack of films and separator sheets. Observe the following guidelines.
 - Put small-film cases at the top of the stack, mixed-film cases in the middle, and large-film cases on the bottom.
 - If the number of films in the case does not match any of the defined scanning protocols, use the Any protocol.
 - Use the correct number of films per case, as specified by the selected scanning protocol.
 - If you plan to scan with the Auto protocol, you can use from one to four films, as long as they have supported lead markers and are from the four standard views with no duplicate views. You can arrange the films in any order and in any orientation. See '3.3. Scanning with the Auto Protocol'. Load only the four standard views with no duplicates.
 - If you plan to scan with a Manual protocol, you must arrange the films exactly as defined by the desired scanning protocol. See '3.4. Scanning with a Manual Protocol'.
 - If you plan to use the Any protocol to scan a special case (one that doesn't conform to any available scanning protocol), then see '3.5. Scanning with the Any Protocol'.

▲ **Note:** If you make a mistake when preparing a case for scanning, you can correct it after scanning is complete via the Image Correction screen. See '**3.10. Correcting Scanned Images**' for more information.

3.3. Scanning with the Auto Protocol

Use this procedure to scan cases using R2's Automatic Film Marker Identification (AFMI) feature. The Auto protocol accepts up to four films per case. You can scan with the Auto protocol if all cases in the stack include only the four standard mammography views (RCC, LCC, RMLO, LMLO), and were processed using a supported lead markers (those shown on page 24) that were well-placed.

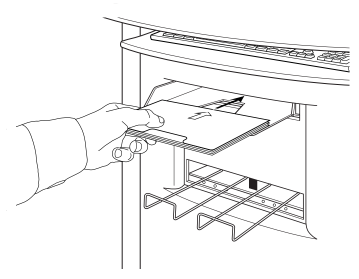
- For cases that do not meet these criteria, but conform to a different scanning protocol, see '3.4. Scanning with a Manual Protocol'.
- For cases which do not confirm to any defined scanning protocol, see '3.5. Scanning with the Any Protocol'.

▲ **Important!** The Auto protocol does not support duplicate views (for example, two RCC films or two LMLO films). Use a Manual protocol or the Any protocol for cases with duplicate views.

To scan with the Auto protocol

- 1 Prepare the cases to be scanned as described in '3.2. Preparing Cases to Be Scanned'. You can arrange the films of the four standard views in any order and in any orientation.
- **2** Load the stack into the input tray. Insert the stack with:
 - The arrow on the top separator sheet pointing into the scanner.
 - The leading edges of the films and separator sheets aligned flush.

You can load up to 100 sheets – including films and separator sheets – up to 20 four-film cases.



Loading Films

- 3 Verify that Auto is selected in the Scanning Protocol list on the Studies screen.
 ▲ Note: You cannot change the scanning protocol if a case is currently being scanned.
- **4** Touch **Start**. The system status changes from 'Ready' to 'Scanning', and the **Start** button becomes a **Stop** button.



- **Processing Films**: After the system scans the first film, it displays a new study icon at the top of the study list, and a white circle on the icon flashes to indicate that the films are being processed. If you touch this icon, you can monitor the progress of the study as the films are processed. There are three possible outcomes:
- **Success**: If the system processes the case successfully and verification of cases is not required, then the system automatically sends the study results to the output device (display unit, printer and/or PACS archive). An arrow on the icon flashes to indicate that the system is sending the results. After the system sends the results, the icon appears solid green.



• Verification Required: If verification of cases is required, then you must review the scanned image thumbnails and patient information and confirm that the results are acceptable. If no Alert has been issued, then the Done button appears on the Studies screen. When you touch Done, the system sends the results to the output device (display unit, printer and/or PACS archive).



• Alerts: If the system replaces the study icon with an Alert icon, you may be required to take corrective action. In many situations instead of rescanning the case, you can often correct the alert and reprocess the images via the Image Correction screen.

For more information on the study icons, see the table on page 14. Also see '3.10. Correcting Scanned Images', and 'Part 4: Alerts'.

5 When the cases have been scanned successfully, put the films and corresponding separator sheets together in the patient folders for hanging.

Note: If the system has been configured to use a different scanning protocol by default (on the Controls Scanning screen), then after scanning a stack of films, the system returns to the default protocol.

3.4. Scanning with a Manual Protocol

When you use a Manual scanning protocol, each case in the stack must include the correct number and type of film views, as defined in the protocol. You may also need to use a Manual protocol if films:

- Have unsupported, missing, or bad lead markers, or
- Have incorrectly placed lead markers (e.g., over a breast image, under an ID label, or only partly on film, as shown on page 25), or
- Have non-standard views or views with modifiers, or
- Have more than four views per case

If one of the cases in the stack has any of the characteristics listed above and does not have the number of film views specified in any existing Manual protocol, use procedure '3.5. Scanning with the Any Protocol' for that case.

The system ships with four predefined Manual scanning protocols.

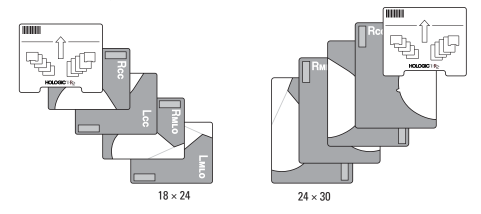
Predefined Manual Scanning Protocols	Views Inc	luded
Screening	RCC LCC	RMLO LMLO
Uni Right	RCC RMLO	RXCC
Uni Left	LCC LMLO	LXCC
Screening Imp	RCC LCC RMLO LMLO	RCC ID LCC ID RMLO ID LMLO ID

Your system may have additional user-created scanning protocols specific to your site. For more information, see '5.7. Scanning Protocols'.

For a list of supported views and modifiers, see 'Supported Views' on page 20.

To scan with a Manual protocol

- 1 Prepare the cases to be scanned as described in '3.2. Preparing Cases to Be Scanned'.
- **2** Arrange the films and their separator sheets in the order shown on the screen below the selected scanning protocol, and with the orientation shown below.



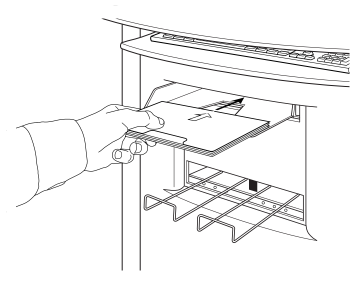
Orientation

- Emulsion (dull) side down
- Small: landscape
- Large: portrait.

Important! When using a Manual scanning protocol, use the same film views for all cases in the stack, and arrange the films in the order defined by the selected protocol.

- **3** Load the stack into the input tray. Insert the stack with:
 - The arrow on the top separator sheet pointing into the scanner.
 - The leading edges of the films and separator sheets aligned flush.

You can load up to 100 sheets – including films and separator sheets – up to 20 four-film cases.



Loading Films



4 Touch the desired scanning protocol. In the example shown at left, Screening is selected. The views defined for the selected protocol are displayed below the protocol list.

Note: You cannot change the scanning protocol while a case is being scanned.

- **5** Touch **Start**. The system status changes from 'Ready' to 'Scanning', and the **Start** button changes to **Stop**.
- 6 A new study icon appears at the top of the study list, and a white circle on the icon flashes to indicate that the films are being processed. If you touch this icon, you can monitor the progress of the study as the films are processed. There are three possible outcomes:
 - **Success**: If the system processes the case successfully and verification of cases is not required, then the system automatically sends the study results to the output device (display unit, printer and/or PACS archive). An arrow on the icon flashes to indicate that the system is sending the results. After the system sends the results, the icon appears solid green.
 - Verification Required: If verification of cases is required, then you must review the scanned image thumbnails and patient information and confirm that the results are acceptable. If no Alert has been issued, then the Done button appears on the Studies screen. When you touch Done, the system sends the results to the output device (display unit, printer and/or PACS archive).
 - Alerts: If the study icon is replaced with an Alert icon, you may be required to take corrective action. In many situations instead of rescanning the case, you can often correct the alert and reprocess the images via the Image Correction screen.

For more information, see '3.10. Correcting Scanned Images'. For information on alert messages, see 'Part 4: Alerts'.

7 When the cases have been scanned successfully, the films can be returned to the patients' film folders. If necessary, keep the separator sheet with the case for CAD display.

▲ **Note:** If the system has been configured to use a different scanning protocol by default (on the Controls Scanning screen), then after scanning a stack of films, the system returns to the default protocol.

3.5. Scanning with the Any Protocol

Use the Any protocol to scan cases that do not conform to an existing scanning protocol, for example, if the case has missing films, extra films, or views not included in any existing scanning protocol.

The Any protocol contains no predefined views. This allows you to scan any number of films (up to 24) with any type of view and later identify the views manually once the case has been scanned.

A neat and ordered stack can save time spent adjusting the image orientation after scanning. Keep the following in mind:

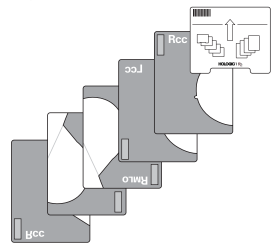
- Remove films from the stack that would normally be discarded rather than saved (for example, technical retakes).
- Orienting the films correctly in the stack saves time adjusting the orientation later via the Image Correction screen.

► To scan with the Any protocol

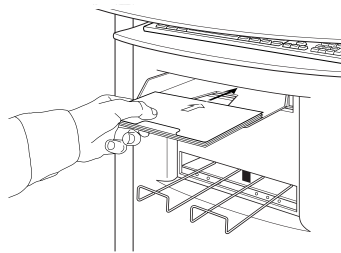
- 1 Prepare the case to be scanned as described in '3.2. Preparing Cases to Be Scanned'.
- **2** Place the separator sheet on top of the stack for each case.

You can arrange the films in any order, although you may wish to use a sequence defined by your site (for example, the four standard views first, followed by additional or equivalent views). As needed, turn films a quarter turn to match the orientation shown on the separator sheet.

If you load films in a random order as shown below, you will have to reorient the images later via the Image Correction screen.



- **3** Load the stack into the input tray. Insert the stack with:
 - The arrow on the top separator sheet pointing into the scanner.
 - The leading edges of the films and separator sheets aligned flush.



You can load up to 100 sheets – including films and separator sheets – up to 20 four-film cases.

Loading Films

- Auto Screening Uni Right Uni Left Screening Imp Any
- 4 Select the Any protocol by touching Any.
 ▲ Note: You cannot change the scanning protocol while a case is being scanned.
- **5** Touch **Start**. The system status changes from 'Ready' to 'Scanning', and the **Start** button changes to **Stop**.

A new study icon appears at the top of the study list. As soon as the first film is scanned, the icon displays the yellow alert symbol. The case status shows 'Failed' with an explanation that the image contains an unknown view. After the entire case has been scanned, you must identify all image views via the Image Correction screen and reprocess the case. To do so:

- 6 Touch an image to display the Image Correction screen. See '3.10. Correcting Scanned Images' for further information on reorienting and labeling images. Identify each image and correct the orientation as needed.
- 7 When you have correctly reoriented and labeled all images, touchReprocess on the Image Correction screen to reprocess the images.

The system returns to the Studies screen, and a white circle on the study icon flashes to indicate that the system is processing the views. When processing is complete, a white arrow on the icon flashes to indicate that the results are being transmitted to a display unit, printer, and/or archive system. After the system transmits the study, the icon appears solid green.

8 When the cases have been scanned successfully, the films can be returned to the patients' film folders. If necessary, keep the separator sheet with the case for CAD display.

Note: After scanning the case, the system returns to the default protocol configured on the Controls Scanning screen.

3.6. Adding Cases to the Stack

You can add one or more cases anywhere in the stack. If you're not in a hurry for the case, add it to the bottom. If you're in a hurry, add it to the top. (Cases at the top are processed first.)

▲ **Note:** This procedure can produce an alert indicating there were too few films for the previous case. However, it is often faster to remove the films and produce the alert than to wait for the films to be scanned and processed.

Important: Verify that cases you add to the stack conform to the selected scanning protocol, or select the desired protocol before scanning the case.

To add cases to the stack

- 1 Prepare the cases to be scanned as described in '3.2. Preparing Cases to Be Scanned'.
- **2** If the system is currently scanning, touch **Stop**. The scanner will stop after it finishes the current film.

Important: When you touch **Stop**, the system always resets to the default scanning protocol. If the system generates an alert, touch **OK** in the pop-up window.

- **3** Remove the films and separator sheets from the feeder. Rebuild the stack in the order you want to run the cases, making sure the films and separator sheets are in the correct order.
 - If you want to run the new cases first, add them at the top of the stack.
 - If the system did not scan all the films from the last case, take the scanned films from the output tray and put them together with the films that were not scanned.
- 4 Insert the new stack of cases into the feeder.
- 5 If necessary, select the scanning protocol you want to use.
- 6 Touch Start to resume scanning.

3.7. Stopping Scanning and Removing a Case

You cannot 'cancel' a study, but if you don't want to continue scanning the current set of films, you can stop the system and remove any remaining films from the input tray (as described below).

▲ **Note:** This procedure can cause the system to produce an alert indicating there were too few films. However, it is often faster to remove the films and produce the alert than to wait for the films to be scanned and processed.

▲ Important: If you changed the scanning protocol from its default setting before you started scanning (e.g., if the default scanning protocol on the Controls Scanning screen is Auto and you touched Screening before touching Start), the system resets the scanning protocol to its default when you touch Stop in step 1 of the procedure below. If you want to resume scanning with the non-default protocol after removing one or more cases, you'll need to select that scanning protocol again before touching Start in step 6.

► To stop scanning

1 Touch **Stop**. The scanner will stop after it finishes scanning the current film.

 \triangle **Note:** If the system generates an alert, touch OK in the pop-up window.

- **2** Remove the entire stack from the input tray.
- **3** Remove the remaining films for the cancelled case.
- **4** As desired, remove any other case, that is, the separator sheet and the films associated with it.
- **5** Place the remaining cases back in the input tray.

Important! Be sure that the next item to be scanned is the separator sheet for the next case.

- 6 Select the desired scanning protocol, if not already selected.
- 7 Touch **Start** to resume scanning.

3.8. Viewing Study Information

As each study is processed, the system displays the study information on the Studies screen. The results include the image thumbnails, patient flash (if present), and patient information.

You can display study information in one of three ways:

- You can touch the study icon in the study list.
- You can type the patient identification, name, or R2 ID (separator sheet barcode number) and press (-Enter) (or touch **Search**).
- You can use the handheld barcode scanner to scan in the patient identification or R2 barcode number.

For more information about searching, see '3.12. Searching for a Study'.

No ImageChecker®	2 Patient ID	Search 🔎
R2 ID: 0000022609 Receive Time: Oct 10 15:26 Status: Processing Images RCC LCC RMO LMO Patient Information R20075cl.img TEST FILM FOR ImageChecker® M1000 SYSTEM R23075cl.img TEST FILM FOR ImageChecker® M1000 SYSTEM R23075cl.img R		Auto Screening Uni Right Uni Left Screening Imp Any
Name: Tester^Teresa Accession: Studies Alerts 🎢 Cor	Itrols	Status Ready Start

the top represents the study currently being processed or the study most recently processed. The Study icons below represent studies processed earlier.

The Study icon at

For any scanned case, the system displays the following:

- The **R2 ID**, which is the barcode number from the study's separator sheet.
- The **Receive Time**, which is the time film scanning began for the selected study.
- The study **Status**. Depending upon the case, the Status field can display a variety of messages (see table, next page).
- Thumbnail **Images** of films. These images appear as they are processed and can be viewed at any time after processing is complete. Touch any thumbnail image to open the Image Correction screen.
- **Patient Information** shows the patient flash from the scanned film (if present), followed by the Patient ID, Name, and Accession number from the patient information system. The patient flash image appears for each film as it is scanned.

After the system scans the last film for a study, it retains the patient flash image from that film as part of the study information.

Note the following:

- To prevent errors, always compare the patient information with the information appearing on the patient flash.
- The image thumbnails always appear in film scanning order. Once the films have been scanned, you cannot change the order.
- The thumbnails should always appear in anatomic orientation (e.g., with the RCC nipple pointing left). If they are not, reorient the images as described in '**3.10**. **Correcting Scanned Images**'.
- For ImageChecker users, CAD marks do not appear on the touch-screen monitor. The CAD marks appear only on the output devices (display units, review workstations, printouts, etc.).

The following table shows the Status conditions that the system can display for a case.

Case Status Message	Meaning
Processing	The system is scanning and processing the selected case.
Needs Verification	The system requires you to review and 'verify' the case images before it will send the study results to a display device or archive.
Transmitting	The system is transmitting the scanned images to a display device or archive.
Completed	The system has finished transmitting the study.
Failed	The system was unable to complete the processing or study transfer and has issued an alert.

If one or more images in the study could not be processed for any reason, a yellow 'X' appears on the image. However, all scanned images will still be sent to the PACS if your system is configured for archiving. See 'Part 4: Alerts' for more information on alert messages.



3.9. Verifying Scanned Cases

When using DigitalNow at some facilities, the system will require you to 'verify' scanned cases before you archive them. The verification step is configured by going to the Controls Connectivity screen and setting 'Verification before sending Output' to 'On' (see '5.2. Connectivity Options').

▲ **Important!** Hologic recommends that facilities using DigitalNow always verify the images before archiving them to reduce the chance of improperly oriented or identified images being sent to a PACS. Also, from time to time, be sure to check images received at the PACS to confirm that films continue to be properly scanned and digitized.

Note: The Verification option is only available for systems configured with the DigitalNow license. Systems that are configured with just the ImageChecker license do not provide the Verification option.



If your system has been configured to require verification, then after the system processes the cases it displays a blinking question mark on each study icon. The question mark also appears on the Studies screen tab at the lower left of the screen.

► To verify cases

1 Touch the study icon for the case you want to verify. The system displays the results for the selected study.



- **2** Review the case results. Check to see that the images are oriented correctly and that the patient information was retrieved successfully from the archive.
- **3** Touch the Done button or make corrections.
 - If the results are acceptable, touch the Done button. In response, the system transmits the results to the archive or other output device.
 - If the results are not acceptable, or if the system produced an alert for the case, you may need to take corrective action. In some situations, you may need to reorient or label images by using the Image Correction screen. For more information, see '3.10. Correcting Scanned Images'.
- 4 Repeat these steps for each case needing verification.

3.10. Correcting Scanned Images

After you scan a case, you use the Image Correction screen to reorient images, change film view labels and modifiers, or add a DICOM study description. For example, you must correct the scanned images if a case includes one or more non-standard views or view modifiers, or if the system is unable to identify a lead marker. In addition, you must use the Image Correction screen for all cases scanned with the Any protocol.

In many cases, the system issues an alert to let you know that it could not completely process the case. However, you can use the Image Correction screen even with cases that have been completely processed without alerts.

The Image Correction screen allows you to:

- Reorient scanned images (flip or rotate them)
- Change the film view labels and modifiers (e.g., RCC to LCC)
- Edit the DICOM study description for the case

Note: If films for two or more patients were scanned under a single separator sheet, you must rescan the case with a different separator sheet for each patient.

To correct scanned images

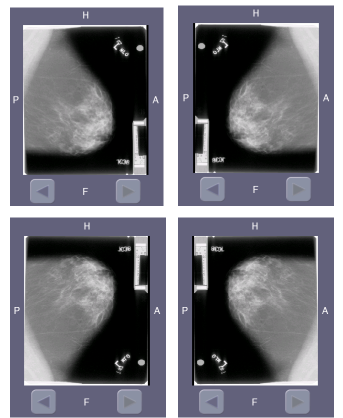
1 Open the Image Correction screen. Touch the study icon for the study you want to work with. Then touch one of the thumbnail images. The Image Correction screen opens as shown in the following example:

H P	Side Left Right Both	View CC XCC MLO XCCL ML XCCM LMO FB LM SIO	Modifiers M ID S TAN RL AT RM CV RI RS
F DICOM Study Description: Mammography Digiti	zed Film	Partial View Implant Present	Reprocess

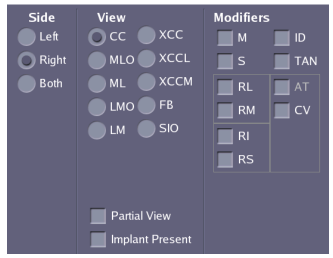
The image thumbnails for the case appear at the top of the screen. The image you selected appears enlarged at left. To navigate through the study images, touch the arrow buttons below the enlarged image, or touch any of the image thumbnails.

▲ **Note:** If the displayed images, their labels, and DICOM Study Description are all correct, then simply touch Cancel to return to the Studies screen. (Cancel does not cancel the study – it simply returns to the previous screen.)

- 2 Reorient image, as needed.
 - **a** Touch the enlarged thumbnail at the left of the screen. Each time you touch the image, it alternately flips 90° along the vertical and horizontal axes. Flipping the image a total of four times restores the original orientation.
 - **b** Check the image orientation with the indicators on the four sides of the image (Left, Right, Anterior, Posterior, Head, Foot). When the orientation is correct, go on to step **3**.



3 Modify the view label, as needed. To define or change the view label, build it from left to right: select a side, a view, and then modifiers (as needed).



- **a** Touch the proper Side selection: **Left**, **Right**, or **Both** (for cleavage view).
- **b** Touch the proper View selection. Note the two additional selections below the view area for Partial View and Implant Present.
- **c** Touch to select Modifiers as needed.

As you build the view label, invalid selections are eliminated. For example, if you choose Both for the side, the system automatically sets the view to CC and the modifier to CV. You can chain valid modifiers together as needed.

Some modifiers (for example, M and S) appear before the view designation; other modifiers (such as ID and TAN) appear after the view designation.

See 'Supported Views' on page 20 for a complete list of views and modifiers.

Note: You must select a view when using the TAN modifier, as TAN can be used with any view.

Note: ImageChecker will not process views modified with the Partial View or IP modifiers. For further information see Understanding ImageChecker 8.5-8.7.

- 4 Repeat steps 2 and 3 for each image.
- **5** Modify the DICOM study description, as needed. If desired, touch the DICOM Study Description field and use the keyboard to change the DICOM study description for the case. The default description is 'R2 Mammography Digitized Film'.

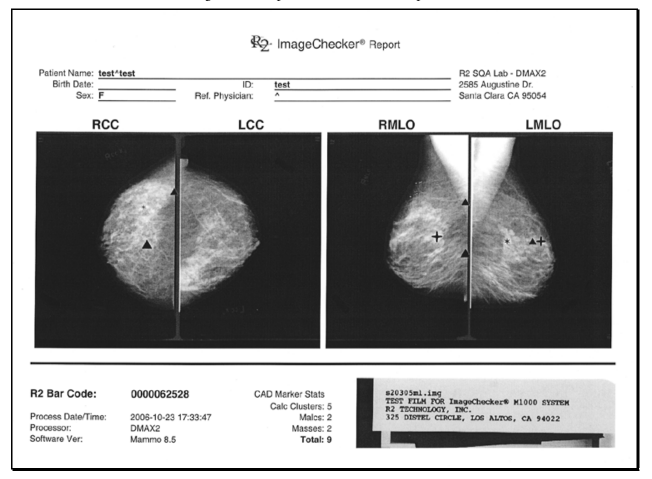
Note: All images in the case will contain the same DICOM Study Description; modifying it in one image changes it for all images in the case.

- 6 Reprocess the case. Once the images and their labels are correct, touch Reprocess. The system cancels the old case (removes the old study icon), reprocesses the case (all images), and creates a new study icon at the top of the study icon list. If the system does not issue an alert, it:
 - Automatically sends the results to the output devices (PACS, display unit, printer, etc.) if Verification is off.
 - Waits for you to click the Done button if Verification is on before sending the results.

3.11. Printing CAD Results

If you have configured the system to print CAD results for every study (as described in '5.6. Printing Options'), the CAD results are automatically printed when processing is complete.

The following is an example of a CAD Results Report:



If you want to create an additional paper printout of the CAD results, touch any of the thumbnail images for the case and reprocess it. Because the films are not rescanned, the results for the case will be identical to the original.

3.12. Searching for a Study

You can search for and display the information from any previously processed study using any of the following study data:

- R2 ID (from the separator sheet barcode)
- Patient ID
- Name

The search is limited to the 200 most recently processed studies (or fewer, depending on the available disk space in the processing unit's computer).

▶ To search for a study and display its results

1 Type the data for the study you want to find. You can just start typing; you don't need to touch the Search field first. You can also enter data by using the handheld scanner to scan a barcode from a patient record or separator sheet.

▲ **Note:** The system does not accept (or need) wildcard characters (e.g., *). The search function will find any data that partially or completely matches the text you enter. If you leave the search field blank, the search returns all studies on the system.

- 2 Touch Search. The search results appear:
 - If only one study matches the search criteria, the system displays the study information automatically.
 - If multiple matches occur, the system displays the Search Results window, as shown in the following example.

R2 ImageCh	necker®		R2	Patient ID		Search 🔎
	s	earch Results			×	Auto
R2 ID: 00 Receive Time: Se Status: Fa	R Yours	search matche	d multiple cas	es. Please pick	one:	Screening Jni Right Jni Left
2 films were found v #0000000660. For Manual.	Name	SID / R2 ID	Patient ID	Time		Screening Imp
	test2^test2	000000660	test12	Sep 28 09:26		
Images	test^test	0000062529	esrtabh	Oct 03 17:56		
Images	Test [^] TEST	0000022598	0000022598	Oct 09 10:22		
	TEST2^TEST2	000001380	TEST2	Oct 09 11:03		cc cc
	Tester^Teresa	0000022609	123456test	Oct 10 15:26		MLO MLO
Patient Informati						
s23075cl.img TEST FILM FOR I R2 TECHNOLOGY, 325 DISTEL CIRC						
- SZS DISTED CIRC						tatus
						Ready
Patient ID: tes					Cancel	
? Studies						Start 📄
					10 Jan - 19 19	

A Note: To move the window, touch and drag the Search Results title bar.

If more than one study appears, touch the study you want to see. The Search Results window closes and the results of the study you selected appear on the Studies tab.

3.13. Using Queue Manager

Queue Manager enables you to monitor and manage the queue of cases being processed by the R2 system.

► To enter Queue Manager

1 Using any web browser on the same network as the R2 system, in the Address box type the IP address for the processing unit followed by /qman, and press ←Enter. For example:

http://12.345.67.89/qman

The browser will display the log-in window.

Pa		help
NY NY		Queue Manager
© 2005-2006 R2 Technol	Password:	login

2 Enter your Queue Manager database password (h0spital spelled with a 'zero'), and click login.

R2 queu						
earch						
atient Name		Birth Date (YYYYMM	1DD)	Study Date From		search
Patient ID		Sł	atus All	(YYYYMMDD) To		clear
to 100 of 217 cases Ne	xt 100 ca	ses >>			_	
Patient Name	Sex	Birth Date	Patient ID	Study Date	Priority	Status
Patient916, Patient916	F		Patient916	20061030	Medium	Processing
Patient915, Patient915	F		Patient915	20061030	Medium	Processing
Patient924, Patient924	F		Patient924	20061030	Medium	Scheduled
Patient916, Patient916	F		Patient916	20061030	Medium	Scheduled
Patient905, Patient905	F		Patient905	20061030	Medium	Scheduled
Patient904, Patient904	F		Patient904	20061030	Medium	Scheduled
Patient918, Patient918	F		Patient918	20061030	Medium	Sending Outputs
run next			delete		stop	
escription			_	_		
Case Index:		Algor	ithm Used:	Auto	Link:	
Patient Name: Sex:		9	Priority: tudy Date:	Completion Acces		
Birth Date:			ing Status:	Input Sou		
Patient ID:		Number	of Images:	Case Li	evel:	
Output Jobs: (Destinatio	on Forma	at Status)	SubCase Li	st: (PatientName Patie	ntID StudyD	ate)
	_					
© 2005-2006 R2 Technolog	, Inc.					

The system displays the Queue Manager screen, showing the list of cases and the status of each case. Click Help at the upper-right corner for detailed instructions.

- You can filter the list of cases by entering search criteria in any of the fields in the Search panel at the top of the screen and then clicking the search button.
- You can sort the list of cases by clicking on any of the column headers. Clicking it again reverses the sort order.
- When you click on a patient name, the information associated with the study is shown in the Description panel at the bottom of the screen.
- **3** Select one or more cases using the checkboxes on the left side of the queue.
- **4** Use the **run next**, **delete**, and **stop** buttons to move the selected case(s) to the top of the queue (run next), delete them, or stop processing them.
- 5 When you are finished, exit the web browser.

Part 4: Alerts

- ► 4.1. Alerts Listed Alphabetically
- ► 4.2. Common Questions

•

Part 4 provides a listing of alert conditions with instructions for resolving the alerts and diagnosing and solving common problems.

The Alerts are designed to help you and Hologic service personnel diagnose problems when they arise. When they occur, the alerts appear in two places:

- On the Studies screen, just below the Status field for the study currently displayed.
- R2 Patient ID

On the Alerts screen, where you can view all recent alerts.

Touch to sort alerts —	Time • PID / R2 ID Description	Auto Screening
by Time, R2 ID, or alert Description.	Oct 10 14:13 0000000541 No patient record information is associate Oct 10 14:09 000000655 No patient record information is associate Oct 10 14:05 000001691 No patient record information is associate	Uni Right Alert Uni Left Alert Screening Imp
Date and time alert —— was generated.	Oct 10 14:03 0000001376 No patient record information is associate Oct 09 23:00 Calibrating the scanner. It may take up to Oct 09 15:29 Rebooting the operating system, please v Oct 09 14:51 Shutting down the R2 software, please v	Alert Any Alert C C C
R2 ID ——	Oct 09 12:10Restarting R2 software, please wait.Oct 09 12:07Restarting R2 software, please wait.Oct 09 10:040000022598Oct 09 10:040000022598Not all films in case #0000022598 are speceeeeeeeeeeeeeeeeeeeeeeeeeeeeeeeeee	Alert
Touch to view or hide alerts you've hidden.	Oct 09 10:04 No output jobs scheduled for case 00000 Oct 09 09:16 Calibration is complete. The scanner is re Oct 09 09:14 Calibrating the scanner. It may take up to Oct 09 09:14 The scanner hasn't been calibrated in 24	Alert
Touch to hide all alerts.	Oct 09 09:11 Rebooting the operating system, please to 00 00-10 The scanner Book's hear calibrated in 24	Status
Touch to hide a selected alert.	Hide Hide All Alerts View Past Alerts ? Studies Alerts ? Controls	Ready Start

Also, whenever the system generates a new alert, an alert icon (Δ) appears on the Alert tab.

4.1. Alerts Listed Alphabetically

The alert messages are presented here in alphabetical order, with instructions for addressing each problem.

Archiving was cancelled for case (3045)

The barcode has no associated patient information in the patient information database.

Enter the patient information into the patient database, including the R2 barcode on the separator sheet scanned with the case, then reprocess the case via the Image Correction screen. See '3.10. Correcting Scanned Images' for more information.

Cannot process image successfully (1063)

The system is unable to perform CAD processing or image enhancement on the image due to poor film quality. A yellow 'X' appears on the image thumbnails on the Studies tab. Even though no processing was possible, the scanned images are archived as usual.

- Verify that the images have the correct view labels, if not correct the view labels on the Image Correction screen. See '3.10. Correcting Scanned Images' for more information.
- Verify that the image views and view modifiers are supported for CAD processing. See '2.6. System Inputs and Outputs' for more information.

Cannot process this case (1062)

The R2 software cannot work with the images as they are.

- Correct the images as necessary via the Image Correction screen, and reprocess the study. See '3.10. Correcting Scanned Images' for more information.
- If films for two or more patients were scanned under a single separator sheet, rescan the case.
- If these steps do not solve the problem, call Hologic Technical Support.

ImageChecker attempted but could not send case (3016, 3018, 3031, 3032)

The system is trying to transfer the images but cannot find the specified output device (display unit, PACS, etc.). The system will try repeatedly to send the images. Usually, it is either a display unit problem or your computer network is down. To fix:

- Make sure the display unit is up and running. If this was the problem, the studies should transfer within about ten minutes.
- If this isn't the problem, check to be sure all cables between the processing unit and the display unit are connected and that the computer network is up.
- If your network is up, all cables are connected, and the display unit is in working condition, call Hologic Technical Support.

ImageChecker attempted but could not print reimbursement report (2008)

The system has canceled the print request for the study. You will only see this alert after several 'Printer not found...' alerts. The system tries repeatedly to print and displays this message when it gives up. To fix:

- Verify that the printer is on, on-line, and has enough paper. After correcting any of these problems, ensure that printing is set to 'on' and reprocess the study in the Image Correction screen.
- If the 'Printer not found...' or similar alert appears again, check that all cables between the printer and the processing unit are connected. Check with your system administrator that your network is up. Reprocess the study in the Image Correction screen and print.
- If these steps do not solve the problem, call Hologic Technical Support.

ImageChecker cannot contact the scanner (1010)

This message may occur during system start-up, or if the scanner was inadvertently turned off. The system does not allow scanning films until the problem is fixed.

- If the system is rebooting, wait about 5 minutes.
- If the system does not restart properly, remove the front panel and verify that the scanner power supply is ON. (If you see green lights, then power is ON.) If not, turn on the scanner power supply and replace the front panel.
- Power-cycle the system. See '6.2. Power-Cycling the System'.
- If these steps do not solve the problem, call Hologic Technical Support.

ImageChecker cannot identify the lead markers (3023)

This message appears when scanning with the Auto protocol. It means that the system did not recognize the lead markers. Either they are not one of the supported types or there is a problem with marker position and/or placement on the film(s). To fix:

• Identify and label the views on the Image Correction screen and reprocess the study. If this doesn't fix the problem, call Hologic Technical Support.

ImageChecker cannot read the barcode (1041)

The scanner is not able to read the barcode on the separator sheet.

- Ensure that the separator sheet is in the proper position (e.g., is not half-fed in the feeder) and is not damaged.
- Reload the stack and restart scanning.
- If these steps do not solve the problem, call Hologic Technical Support.

ImageChecker is busy (3012, 1029, 1030)

The images for the study have not yet been sent.

- Wait about ten minutes the problem should resolve itself.
- If it doesn't, ensure the destination (i.e., display unit or printer) is up and ready to receive images.

- Check that all cables between the processing unit and the destination are connected and that the network is up.
- If these steps do not solve the problem, call Hologic Technical Support.

ImageChecker license not found (1024)

This message means the system could not verify a valid license.

• Ensure that the license dongle is present and properly connected. If this does not solve the problem, call Hologic Technical Support.

ImageChecker could not print a CAD report (2007)

The system has canceled the print request for the study. You will only see this alert after several 'Printer not found...' alerts. The system tries repeatedly to print and displays this message when it gives up. To fix:

- Verify that the printer is on, on-line, and has enough paper. After correcting any of these problems, ensure that printing is set to 'on' and reprocess the study in the Image Correction screen.
- If the 'Printer not found...' or similar alert appears again, check that all cables between the printer and the processing unit are connected. Check with your system administrator that your network is up. Reprocess the study in the Image Correction screen and print.
- If these steps do not solve the problem, call Hologic Technical Support.

ImageChecker could not send results (2005)

The system has tried to send the results to the desired output destination, but the time limit to retry has expired without successfully sending the results.

- Reprocess the images via the Image Correction screen. See '3.10. Correcting Scanned Images' for more information.
- If the problem persists, call Hologic Technical Support.

Images didn't transfer (1031)

Images couldn't be sent to the output device (display unit, PACS, etc.) and the transfer was canceled by the system. You will see this alert only after several 'ImageChecker ... could not send case #...' alerts (#3016) have appeared. The system tries repeatedly to send the studies and displays this message when it gives up trying.

- Be sure the display unit is up and running.
- Check cable connections between the processing unit and the display unit (i.e., the network cable from the system to the wall and the display unit network cable to the wall). Reprocess the study in the Image Correction screen.
- If this doesn't fix the problem, ask your system administrator if the network is up.
- If these steps do not solve the problem, call Hologic Technical Support.

Image transfer was interrupted (3013)

The system is trying to transmit results but cannot find the specified output device (display unit, PACS, etc.). The system will try repeatedly to send the results. To fix:

- Make sure the destination device on. If this was the problem, the results should appear within about ten minutes.
- Check that all cables between the processing unit and the destination device are connected and that the network is up.
- If these steps do not solve the problem, call Hologic Technical Support.

No output jobs scheduled for case (3054)

The R2 system was not configured with any output destinations for CAD results.

If your system is set up to output to the printer only verify that printing is turned on at the Controls screen. See '5.6. Printing Options' for more information

No patient record information is associated with barcode (3501)

This alert appears when you are attempting to transmit the study results to an archive. It means that there is no patient record in the database with that specific barcode. When this alert is issued, the results will not be archived. Although you can print the results for the study, any patient information in the database will not be listed on the printout.

The images in the R2 system are linked to the patient record in the database by the separator sheet barcode number (R2 ID). When you scan the films, the R2 system searches for the patient record in the database so it can acquire the patient information.

Always enter the R2 barcode number into the patient record in the database before you scan the films.

If there is no patient information associated with the R2 ID, you can either:

- Enter the information into the R2 Patient ID system and reprocess the case, or
- Enter the information into PenRad, MRS or MagView and reprocess the case (to requery from their database)

Not all films in case are specified by scanning protocol (3024)

There are more films in the case than defined by the selected scanning protocol. Identify the film views via the Image correction screen and reprocess. See '3.10. Correcting Scanned Images' for more information.

Printer not found (3014)

The processing unit is attempting to send the study results to the printer, but the printer is not responding. The printer may be unable to respond, or there may be a network or configuration problem. The system will continue to try to send the study results.

- Verify that the printer is running.
- Check all cable connections between the processing unit and the printer (i.e., the network cable from the processing unit to the wall and the printer network cable to the wall). Sometimes a cable becomes loose. If so, wait about ten minutes, the system will resend the study results.

- Ask your system administrator if the network is down. If it is, and it is down long enough, the system will finally give up, and you will have to reprocess the study with printing turned on.
- If these steps do not solve the problem, call Hologic Technical Support.

Rebooting the operating system, please wait (5021)

You'll see this message when the system is rebooting (starting up again after being shut down and turned off).

- Wait about 10 minutes. Once the system comes back up, you will see the Ready message above the Start button.
- If the system doesn't come back up after 15 minutes, call Hologic Technical Support.

Reporting system query error (3502)

The R2 system could not communicate with the patient database, and possibly the system is not set up correctly. Contact Hologic Technical Support and they will diagnose and fix the problem.

Restarting R2 software, please wait (5019)

- Wait about 10 minutes. Once the system comes back up, you will see the 'Ready' message above the Start button.
- If the system doesn't come back up after 15 minutes, call Hologic Technical Support.

Scanner not ready! (5002)

For some reason, the scanner did not turn on properly.

- Power-cycle the system. See '6.2. Power-Cycling the System'.
- If this doesn't fix the problem, call Hologic Technical Support.

Scanner test failed (3025)

This message appears when you run the weekly test films and the scanner test fails. Usually you can clear the alert by rerunning ALL tests again.

- First, make sure the test films and separator sheets are not bent or damaged in any way. If they are, call Hologic Technical Support for replacements.
- Rerun all test films. See '6.4. Running the Weekly Tests'. Usually, this fixes the problem.
- If any tests fail again, power-cycle the system (see '6.2. Power-Cycling the System') and rerun the tests again.
- If any tests fail at this point, call Hologic Technical Support. Hologic does not recommend that you use the system if any of the tests failed.

Scanning is disabled, remote maintenance in progress (5010, 5059)

Hologic Technical Support is working on your system, and you'll need to wait until they are done.

- Wait two minutes. The Ready message will appear above the Start button.
- If your Hologic Technical support is done and the message does not go away, power-cycle the system. See '6.2. Power-Cycling the System'.
- If this doesn't fix the problem, call Hologic Technical Support.

Scheduled reimbursement report (5050)

Appears when the system prints a reimbursement report automatically (weekly or monthly, depending upon the system configuration). This is normal. The message does not appear when you print a reimbursement report manually. See '5.6. Printing **Options**' for more information.

Shutting down the operating system, please wait (5020)

This means the system is shutting down.

- Let the system shut down. Usually, it shuts down only when instructed to do so.
- If this happened on its own, power-cycle the system to bring the system back up. See '6.2. Power-Cycling the System'.
- If the alert continues to appear, contact Hologic Technical Support.

Shutting down the R2 software, please wait (5018)

This means the system is shutting down. It will shut down the R2 software, then the operating system.

- Let the system shut down. Usually, it shuts down only when instructed to do so.
- If this happened on its own, power-cycle the system to bring the system back up. See '6.2. Power-Cycling the System'.
- If it happens again, contact Hologic Technical Support.

The same barcode has been assigned to multiple patients (3500)

This alert appears when you are sending results to the patient database. It means you assigned the same R2 barcode number to two different patients in the database.

The images in the R2 system are associated to the patient record in the database by the barcode number on the separator sheet. When you scan the studies, the R2 system searches for the patient record in the database that has that barcode and associates the images to that patient record. If there is more than one patient record with the same barcode, the system does not know which patient the images belong to.

- Access the database and check the patient records.
- Replace the duplicate barcode number with a new barcode number. Use this new sheet to scan the case.

The scanner is not responding (1007)

The internal scanner software is not responding to signals from the processing unit. By power-cycling the system, you can reset the internal scanner software.

- Power-cycle the system. See '6.2. Power-Cycling the System'.
- If this doesn't fix the problem, call Hologic Technical Support.

The scanner might be empty (3033)

The scanner is not detecting any films in the scanner.

- Ensure that the films and separator sheets are properly loaded (e.g., are not half fed in the feeder).
- Reload the case and restart scanning.
- If these steps do not solve the problem, call Hologic Technical Support.

X films were found when Y were expected. (1017, 1018)

This usually happens when (1) a separator sheet is missing from the stack or is upside down, (2) the system failed to read the barcode for the next case in the stack (the barcode is obscured, smudged, or damaged), or (3) a case did not have the number of films specified by protocol.

- Check to see if the next case is missing its separator sheet or if the barcode is obscured somehow. Add a separator sheet, or clean the separator sheet. If films for two or more patients were scanned under a single separator sheet, rescan the case.
- In many situations, you can simply correct the image labels and reprocess the images via the Image Correction screen. See '3.10. Correcting Scanned Images' for more information.
- If these steps do not fix the problem, call Hologic Technical Support.

4.2. Common Questions

This Section provides common issues and solutions for R2 systems.

CAD does not run on views with implants

CAD does not run on images with the Implant Present modifier enabled, even if Implant Displaced is also enabled. If it is necessary to run CAD on Implant Displaced images, do not enable the Implant Present modifier either on the Image Correction screen or in scanning protocols.

CAD marks do not appear on thumbnails

Marks do not display on the thumbnails. They only appear on the display unit, print out, and/or archive system (if configured).

Case icons become yellow alerts; there are no images

Typically, an alert will be generated when an image fails to process. If it did, follow the instructions in the alert, or turn to the Alerts tab and look for that alert.

If an alert did not pop up, and there are no alerts in the Alerts screen, rescan the case using the appropriate Manual scanning protocol.

Film is 'missing' inside the scanner

If a film jams in the scanner, see 'Ejecting Films' on page 68.

Image is missing from a case

Occasionally, the film scanner reads the separator sheet barcode *through* the last film of a case. Although extremely rare, this can result in a case where the last image is missing. Sometimes an alert will appear ('X films were found when Y were expected'), but not always. This situation is most likely to occur with an LMLO image from a film with a light pectoral muscle, or with any film that has an extremely light area in the upper-left corner.

If you suspect that the film scanner has read a separator sheet barcode through a film, then reorient the film and rescan the case. As an alternative, run the case at the bottom of the next stack of cases.

Images are reversed in the thumbnails

Typically, this happens either when you are scanning with a Manual protocol and did not stack the films in the order expected by the R2 system or when you are scanning with the Auto protocol and the lead markers are bad or obscured.

- Use the Image Correction screen to flip the images to the correct orientation and reprocess. See '3.10. Correcting Scanned Images'.
- If the problem persists, contact Hologic Technical Support.

Images at PACS/workstation are of poor quality

Contact Hologic Technical Support. In some cases, a service representative may need to adjust the contrast and brightness settings for your system.

Images didn't print out

- Make sure the study processed successfully, or that there were no printer alerts. If a case did not scan and/or process successfully, it will not print out.
- If the system couldn't find the printer, it will generate an alert. To find out if there was an alert, check the Alerts screen.
- If the study processed correctly, check to make sure that printing is enabled. See 'Configuring CAD Results Printing' on page 70 for information on turning printing on/off. If printing is turned off, turn it on and reprocess the study in the Image Correction screen.
- If none of these steps fixes the problem, contact Hologic Technical Support.

Note: Not all scanned images are printed. Only four images appear on the printout. Images which have not been processed by CAD, and reversed views do not print.

Images do not appear on my display unit

Typically, this happens when there is a delay on the processing unit.

- Check the processing unit for alerts.
- If there were no alerts, wait ten minutes, check the display unit again.
- Ask your system administrator if there have been network problems.
- If after ten minutes the images don't display and your network is up, contact your Hologic Service Representative.

Note: R2 display units do not display all scanned images. For more information, see **'2.6. System Inputs and Outputs**'.

Monitor isn't working

The monitor power switch may have been accidentally turned off. The processor unit is probably still working. Switch the monitor back on. If this doesn't work, verify that the monitor power cable is securely fastened.

Main screen is frozen and I can't do anything

Contact Hologic Technical Support. They will log in remotely using HologicConnect and troubleshoot the system. DO NOT just turn off the system; if you do, the software may not shut down properly, which can result in other problems.

Patient ID flash on the printout is backwards

- Typically, this happens when you scan the case with a Manual protocol and loaded the stack or case upside down. See '3.10. Correcting Scanned Images' for instructions on adjusting the image orientation.
- This problem can also occur if the service configuration is incorrect. If the problem persists, contact Hologic Technical Service.

Radiologist needs CAD quickly

For sites that archive images to PACS, a radiologist may request a quick turnaround for CAD results from certain films without waiting for the system to scan all films. Splitting the films into two cases may be undesirable because that produces separate studies at the PACS for the case. In such a situation, you can turn off Archiving (see **5.2.** Connectivity Options), scan and process the requested views, and send the results to the CAD display unit. Then, turn Archiving back on, scan and process all films for the case, and send the results to the PACS.

Scanned case before entering patient information

You can enter the patient information after scanning and reprocess the study to link the images to the proper patient. You must include the barcode from the separator sheet.

- See 'Part 7: Using DigitalNow and the R2 Patient ID System' for more information on entering patient information.
- See '3.10. Correcting Scanned Images' for information on reprocessing a study.

Yellow 'X' appears on my image thumbnails

The yellow 'X' icon indicates that CAD processing could not be performed on an image.

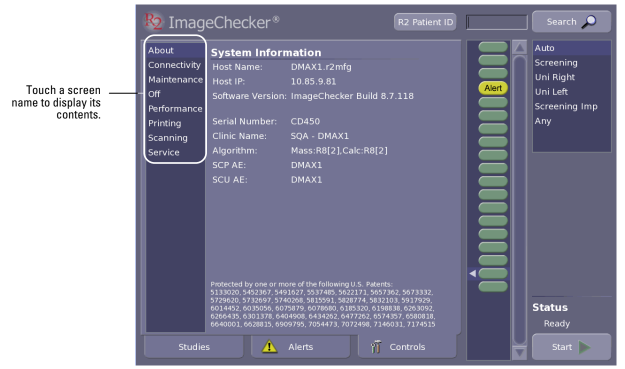
- Verify that the images have the correct view labels. If they do not, correct the view labels on the Image Correction screen. See '3.10. Correcting Scanned Images' for more information.
- Verify that the image views and view modifiers are supported for CAD processing. See '2.6. System Inputs and Outputs' for more information.

Even though no processing was possible, the scanned images are archived as usual.

Part 5: Controls

- ▶ 5.1. About Option
- 5.2. Connectivity Options
- 5.3. Maintenance Options
- 5.4. Shutdown and Reboot Options
- ► 5.5. Performance Options
- **5.6.** Printing Options
- 5.7. Scanning Protocols
- ► 5.8. Service Utilities

Part 5 provides instructions for using the system configuration and maintenance options that appear on the **Controls** screen. The **Controls** screen is used to configure system settings to meet your particular needs, run maintenance utilities, and display system information.



Note: The **Connectivity** and **Printing** options appear only if they were configured by the Hologic service engineer when the system was installed.

5.1. About Option

The About option on the Controls screen displays information about the configuration of your R2 film-scanning system on the network. This information is configured by service personnel when the system is installed.

► To display the About screen

Touch the **Controls** tab and then select **About** (if the About screen does not appear).

Ro Imag	eChecker®		R2 Patient ID	Search 🔎
About Connectivity Maintenance Off Performance Printing Scanning Service	System Inform Host Name: Host IP: Software Version: Serial Number: Clinic Name: Algorithm: SCP AE: SCU AE: SCU AE:	mation DMAX1.r2mfg 10.85.9.81 ImageChecker Build 8. ⁻ CD450 SQA - DMAX1 Mass:R8[2],Calc:R8[2] DMAX1 DMAX1 DMAX1 Solution Science Sc	: 62, 5673332, 03, 5917929, 38, 6263092, 357, 6580018,	Auto Screening Uni Right Uni Left Screening Imp Any Status Ready
Studie			Controls	Start

This screen provides some important data that you may need to provide to a Hologic service support specialist:

- Host IP address
- Software version number
- Instrument serial number
- CAD algorithm operating points (The example above shows that operating point [2] has been selected for both calcifications and masses.)

5.2. Connectivity Options

The Connectivity option on the Controls screen allows you to specify whether to:

- Retrieve patient information when you scan a new case
- Enable or disable the DigitalNow archiving feature
- Verify DigitalNow images before sending them to an archive

In order to use these features, a Hologic service engineer must (1) configure the system with the DigitalNow license and enable the license, and (2) configure your system to receive information from a patient database and transmit results to an external storage archive.

Note: The Connectivity options appear only if the system has been configured with the DigitalNow license.

To display the Connectivity screen

Touch the **Controls** tab and then select **Connectivity**.

R2 Imag	eChecker [®] R2 Patient ID	Search 🔎
About Connectivity Maintenance Off Performance Printing Scanning Service	Retrieve Patient Information for New Cases On Off DigitalNow On On Off On Off Verification before sending Output On Off	Auto Screening Uni Right Uni Left Screening Imp Any
Studie	s 🛕 Alerts 🛉 Controls	Status Ready Start

For more on these settings, refer to the next page.

Retrieving Patient Information

This option on the Connectivity screen allows you to specify whether the R2 system should retrieve information from your patient database. If set up properly, when you scan the R2 ID, the R2 system will request that the database system transmit the patient information associated with the case.

If a Hologic service engineer has configured the R2 system to communicate with the patient database, then you can enable or disable the data transfer using the Retrieve Patient Information setting on the Connectivity screen.

Retrieve Patient I	nformation for New Cases
On Off	

To enable or disable this feature, simply touch On or Off, as desired.

Enabling DigitalNow

The DigitalNow option on the Connectivity screen allows you to enable or disable DigitalNow.



To enable or disable this feature, touch **On** or **Off**, as desired.

Enabling Verification

The Verification option on the Connectivity screen allows you to specify whether a verification step will be required after you scan the films. For more information, see '3.9. Verifying Scanned Cases'.



To enable or disable this feature, touch **On** or **Off**, as desired.

▲ **Note:** Changing the Verification status ('On' or 'Off') has no effect on studies currently waiting to be verified or those currently processing. You must still manually verify each case.

Note: Verification can be turned 'On' only if the DigitalNow option is also 'On'.

5.3. Maintenance Options

The Maintenance option on the Controls screen allows you to review whether scheduled maintenance is needed.

► To display the Maintenance screen

Touch the **Controls** tab and then select **Maintenance**.

R2 Imag	eChecker®		R2 Patient ID	Search 🔎
About Connectivity Maintenance Off Performance Printing Scanning Service	Test Films: Reboot: M Service	Information OK, 6 days left OK, 7 days left Needed Tests: OK, all tests passed		Auto Screening Uni Right Uni Left Screening Imp Any Status Ready
Studie	s 🛕	Alerts	Controls	Start

The screen identifies any required maintenance procedures with a yellow alert icon. For further information, see 'Part 6: Maintenance'.

Maintenance Information					
🛕 Test Films:	Needed				
Reboot:	Ok, 4 days left				
Scanner Cleaning:	Ok, 7 days left				
PM Service:	Ok, 81 days left				
Diagnostic Tests:	Ok, all tests passed				

5.4. Shutdown and Reboot Options

The Off option on the Controls screen allows you to:

- Reboot (restart) the system
- Shut down the system

► To display the Off screen

Touch the Controls tab and then select Off.

R2 Imag	eChecker®	R2 Patient ID	Search 🔎
About Connectivity Maintenance Off Performance Printing Scanning Service	Turning the System Off Reboot Stop and restart the Shutdown Safely power down		Auto Screening Uni Right Uni Left Screening Imp Any Status Ready
Studie	s 🔥 🔥 Alerts	🗍 Controls	Start 🕨

Use the Shutdown option whenever you need to power-cycle the system. Normally, you will shut the system down on a weekly basis during routine maintenance and sometimes during troubleshooting. For further information, see '6.2. Power-Cycling the System'.

When you touch Reboot or Shutdown, a confirmation window appears which requires you to either confirm or cancel the power down command.

5.5. Performance Options

The Performance option on the Controls screen provides access to the:

- Film ejection procedure
- Scanner cleaning procedure
- Results of system verification testing

► To display the Performance screen

Touch the **Controls** tab and then select **Performance**.

R2 Imag	eChecker®		R2 Patient ID		Search 🔎
About Connectivity Maintenance Off Performance Printing Scanning Service		formance ent Film In case	a film gets stuck		Auto Screening Uni Right Uni Left Screening Imp Any
	Processor Test Scanner Film #:	Jul 24, 2008 PAS Jul 24, 2008 CO Jul 24, 2008 CO Jul 23, 2008 PAS Jul 23, 2008 PAS	MPLETE MPLETE SSED	1 100000	Status Ready
Studie	es 🛕 🛕	Alerts	າ) Controls		Start

Ejecting Films

Use the following procedure if a film seems to be jammed inside the scanner.

► To remove a jammed film

- 1 Display the Performance screen by touching the **Controls** tab and selecting **Performance**.
- 2 Touch the Eject Current Film button and wait to see if the scanner ejects the film.
- 3 If the film is not ejected, then gently try to remove the film by hand.
- 4 If the preceding step doesn't eject the film, power-cycle the system. See '6.2. Power-Cycling the System'.
- 5 If the preceding steps do not fix the problem, call Hologic Technical Support.



Reviewing System Verification Results

The Test Results portion of the Performance screen provides the results of the most recent system verification testing, including the scanner test.

Test Results		
CPU	Jul 24, 2008	PASSED
Algorithm Test	Jul 24, 2008	COMPLETE
Processor Test	Jul 24, 2008	COMPLETE
Scanner Film #1	Jul 23, 2008	PASSED
Scanner Film #2	Jul 23, 2008	PASSED

For more information on the tests, see '6.4. Running the Weekly Tests'.

5.6. Printing Options

The Printing option on the Controls screen allows you to:

- Configure the system to send images and CAD results to a printer on your network. See 'Configuring CAD Results Printing' on page 70.
- Configure the system to print the barcode reimbursement report on a weekly or monthly basis (or not at all).
- Print the barcode reimbursement report. See 'Setting When the Reimbursement Barcode Report will be Printed' on page 70.

Note: Printing options appear only if configured by the Hologic service engineer when the system was installed.

Note: Not all scanned images are sent to the printer. For more information, see '**2.6**. **System Inputs and Outputs**'.

Once configured, you can use this screen to decide how you want to use the printer.

► To display the Printing screen

Touch the **Controls** tab and then select **Printing**.

R2 Imag	eChecker®	R2 Patient ID	Search 🔎
About Connectivity Maintenance Off Performance Printing Scanning Service	Print CAD results for every case On Off On Off Reimbursement Barcode Report Weekly Off Monthly Print Now Print Now Print a reimbursement reimbursemen	eport right now	Auto Screening Uni Right Uni Left Screening Imp Any Status Ready
Studie	s 🔥 Alerts	Controls	Start

Configuring CAD Results Printing

You can configure the system so that it

- Prints images with CAD results for every case, or
- Does not print at all.

To enable or disable this feature, touch **On** or **Off**, as desired.



Setting When the Reimbursement Barcode Report will be Printed

You can choose to print a Reimbursement Barcode Report either automatically on a weekly or monthly basis, or not at all.

- To specify automatic printing of the reimbursement report, touch **Weekly** or **Monthly**.
- To disable automatic printing of the reimbursement report, touch Off.

Reimbursement	Barcode Report
🔵 Weekly	Off
Monthly	
Print Now	Print a reimbursement report right now

Printing a Reimbursement Barcode Report

- ► To print a reimbursement report
 - 1 Touch Print Now. A calendar window appears.
 - 2 Specify the date range and touch **Print**.

An example of a Reimbursement Report is shown below. The clinic information in the header can be configured by a Hologic service engineer. For each case, the report prints the R2 ID from the separator sheet, or the patient information, if the system was able to retrieve the patient information from the patient database.

K5 ∎	nageChecker [®] Computer Aided Detection	1234 Eastside Drive Northside, CA 95014 United States
CASE H	ISTORY REPORT: Mar 07	7 2007 - Mar 13 2007
Mar 07 2005 Wed		
Barcode: 0000095458		
Case Time: 17:38:53	ID:	Birthdate:
Barcode: 0000095460	1	
Case Time: 17:41:09	ID:	Birthdate:
Barcode: 0000095461		
Case Time: 17:43:26	ID:	Birthdate:
Name: Sellers, Patricia		
Case Time: 17:45:32	ID: 406893192	Birthdate: 19510501
Name: Eldridge, Jessica		
Case Time: 17:47:53	ID: 289450104	Birthdate: 19180702
Name: Binder, Taylor		
Case Time: 17:49:59	ID: 338495028	Birthdate: 19530401
Name: Keefer, Ruby		

5.7. Scanning Protocols

The Scanning option allows you to perform the following procedures:

- 'Changing Scanning Protocol Order' on page 73
- 'Changing the Default Scanning Protocol' on page 73
- 'Deleting a Scanning Protocol' on page 73
- 'Creating Scanning Protocols' on page 74
- 'Editing Scanning Protocols' on page 75

► To display the Scanning screen

Touch the **Controls** tab and then select **Scanning**.

R2 Imag	eChecker®	R2 P	atient ID	Search 🔎
About Connectivity Maintenance Off Performance Printing Scanning Service	eChecker® Scanning Proto Auto Screening Uni Right Uni Left Screening Imp Any			Search O
Studie	s A	Move Up Move Down lerts Ŷ∏ Contro		Status Ready Start

Changing Scanning Protocol Order

You can change the order in which the scanning protocols appear in the scanning protocol list on the Studies screen.

- ► To change the scanning protocol order
 - 1 Touch a scanning protocol in the list at the left of the screen.
 - 2 Touch the **Move Up** and **Move Down** buttons to move the selected scanning protocol up or down in the list.

Changing the Default Scanning Protocol

The default scanning protocol appears at the top of the scanning protocol list.

► To change the default scanning protocol

- 1 Touch the desired default scanning protocol in the list at the left of the screen.
- 2 Touch the **Move Up** button to bring the desired protocol to the top of the list.

Deleting a Scanning Protocol

You can delete unwanted scanning protocols.

- To delete the default scanning protocol
 - 1 Touch the desired scanning protocol in the list at the left of the screen.
 - 2 Touch Remove.
 - **3** Confirm that you want to delete the protocol.

Note: You can delete any scanning protocol except the Auto protocol.

Creating Scanning Protocols

The Scanning option on the Controls screen allows you to create custom scanning protocols, and edit or delete existing protocols.

► To create a new scanning protocol

- 1 Touch New on the Controls Scanning screen to display the Edit Protocol Screen.
- 2 Touch (or click in) the Protocol Name field, and enter a name for your new protocol.

Protocol Name:	New Protocol				
RCC	Remove	Film: RCC			
		Side	View		Modifiers
		🔵 Left	O cc	🔵 xcc	M DID
		Right	MLO	XCCL	S TAN
		Both	ML	🔵 хссм	RL AT
				FB	RM CV
			LM	SIO	
					RS
			Partia	l View	
	û Up		📕 Implar	nt Present	
	🕀 Down				

- **3** Touch **New** to add a film view to the protocol. Build the film view from left to right: select a side, a view, and any modifiers.
 - a Touch the proper Side selection: Left, Right, or Both (if cleavage view).
 - **b** Touch to select the View. Note the two additional selections below the view area for Partial View and Implant Present.
 - c Touch to select any modifiers needed.

▲ **Note:** As you build the film view, invalid selections are eliminated. For example if you choose Both for the side, the view is set to CC and the modifier is set to CV. Valid modifiers can be chained together as needed. Some modifiers (for example, M and S) are added before the view designation, other modifiers (such as ID and TAN) are added after. For a list of supported views see 'Supported Views' on page 20.

- 4 Repeat step 3 for each film view you wish to add to the scanning protocol.
 - To move a view, touch a view name in the list at left and then touch **Up** or **Down**.
 - To remove the view from the scanning protocol, touch a view name in the list at left and then touch **Remove**.
 - To enter a study description, touch the **DICOM Study Description** field.
- **5** Touch **OK** when finished.

Editing Scanning Protocols

You can edit scanning protocols from the Controls Scanning screen.

► To edit a scanning protocol

- 1 Touch a scanning protocol in the list at the left of the screen.
- 2 Touch Edit to open the Edit Protocol screen.
- **3** Edit the Protocol. You have the following options:
 - To change the view order, touch a view name in the list at the left, then touch **Up** or **Down** as needed.
 - To remove the view from the scanning protocol, touch the view name, then touch **Remove**.
 - To enter a new name for the protocol, touch the Protocol name, then enter a new name.
 - To add a new view to the protocol, touch **New**. Build the view from left to right starting with the side, then a view, then modifiers.
 - To add or change the DICOM study description, touch the field and edit the study description as desired.
- 4 Touch **OK** when finished, or touch **Cancel** to cancel changes to the scanning protocol.

Note: You can edit any scanning protocol except the Auto protocol. If you delete all views from a Manual protocol, the protocol will be equivalent to the Any protocol.

5.8. Service Utilities

The Service option on the Controls screen allows:

- Calibrating the touch screen
- Access to the service tools (only for Hologic-trained service personnel)

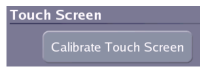
► To display the Service screen

Touch the **Controls** tab and then select **Service**.

R2 Imag	eChecker®	R2 Patient ID	Search 🔎
About Connectivity Maintenance Off Performance Printing Scanning Service	Touch Screen Calibrate Touch Screen Service Tool Connect to the Service Tool		Auto Screening Uni Right Uni Left Screening Imp Any Status Ready
Studie	s 🛕 🛕 Alerts 👘	Controls	Start

Calibrating the Touch Screen

Calibrating the touch screen can make it easier to precisely touch the buttons and other controls.



To calibrate, first touch **Calibrate Touch Screen**, and then follow the simple onscreen instructions. When you finish, you will have to restart the system to save the new settings.

Connecting to the Service Tool

Access to the service tool is password protected and intended solely for use by authorized Hologic service personnel.

Part 6: Maintenance

- ► 6.1. Maintenance Reminders
- ▶ 6.2. Power-Cycling the System
- ▶ 6.3. Cleaning the Scanner
- ▶ 6.4. Running the Weekly Tests
- ▶ 6.5. Resetting the Handheld Barcode Scanner

Part 6 provides maintenance procedures that must be performed **weekly** to ensure optimal system performance. Hologic strongly recommends you follow all maintenance instructions given in this manual.

6.1. Maintenance Reminders

The system automatically issues a reminder message whenever a required maintenance procedure must be performed. The reminder messages are listed below.

Diagnostic tests failed

The scanner diagnostic tests failed. Each time the system starts up, the scanner performs tests to make sure all internal systems are working properly. You cannot scan until the scanner passes its tests. To fix the problem, power-cycle the system. See '6.2. Power-Cycling the System'. If this doesn't resolve the problem, call Hologic Technical Support.

Reminder: Please clean the scanner

Clean the scanner regularly in order to keep the system performing optimally. For instructions on cleaning the scanner, see '6.2. Power-Cycling the System'.

Test Films: Needed

The system test films have not been run, are overdue, or have failed. To fix the problem, see '6.4. Running the Weekly Tests'. (More information about individual film tests is available on the Controls Performance page; see '5.5. Performance Options'.)

Preventive maintenance is required

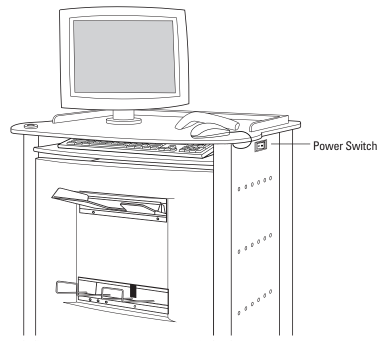
Your system is due for its quarterly preventive maintenance servicing. Call Hologic Technical Support to schedule an appointment. In the meantime, you can continue scanning. If the system is not serviced within 180 days, the alert will appear continuously.

6.2. Power-Cycling the System

'Power-cycling' means to shut down the computer and then switch it back on. By power-cycling once a week, you will help keep the processing unit computer operating properly.

► To power-cycle the system

- 1 Touch the **Controls** tab and then select **Off**.
- **2** Touch the **Shutdown** button. After the software application shuts down, the monitor displays 'Power down...'
- 3 Switch off the unit using the power switch on the side of the cart.



Shutting Off the System

Wait ten (10) seconds. Switch on the system using the power switch. When the Studies screen appears, the system will be ready for use.

6.3. Cleaning the Scanner

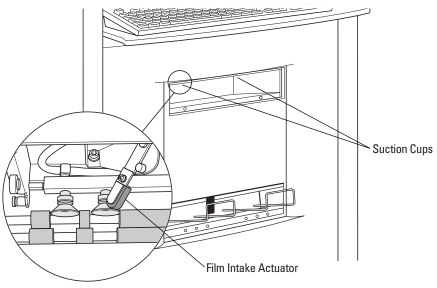
To ensure optimum scanner performance, once a week clean the scanner's suction cups using alcohol and the cleaning swabs provided with the system.

► To clean the suction cups

- 1 Switch off power to the system as described in '6.2. Power-Cycling the System'.
- 2 Remove any films in the scanner's input tray, then remove the input tray itself by carefully sliding it straight out from the scanner. Set the input tray aside.

Caution: Never remove the input tray during a film-feed cycle.

3 Dip a cleaning swab in isopropyl alcohol, and then gently swab the surfaces of the four suction cups inside the scanner.



Suction Cup Location

Important: Inspect the foam tip of the film intake actuator. If the foam appears worn or damaged, call Hologic Service.

4 *Carefully* replace the input tray. Tilt the tray up about 20 degrees and gently slide it in. Be sure that the plastic rails on the sides of the input tray slide under the metal flange on the mounting tracks. Push in tray until it snaps into place.

Caution: After cleaning the suction cups, wait at least five minutes before scanning any films. Residual alcohol can damage the films!

5 Switch on the power as described in '6.2. Power-Cycling the System'.

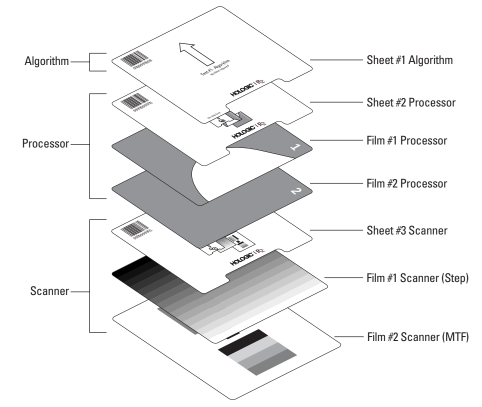
6.4. Running the Weekly Tests

Three tests must be run weekly to ensure optimal system performance. Always run the tests together and in this order: Algorithm on top, then Processor, then Scanner – think 'APS'!

To run the weekly tests

1 Stack the test sheets and films as shown. Note that the test films are ordered as if they were ordinary films.

Important! If your system does not have the CAD license, do not run the Algorithm and Processor tests. Run only the Scanner test films.



- **2** Place the stack in the input tray and touch **Start**. The computer recognizes the test film barcode numbers and automatically changes to Test mode. The test requires about five minutes to complete.
- **3** When done, review the results by touching the **Controls** tab and then selecting the **Performance** option.
- 4 Verify that all test results read 'PASSED'.
 - If all tests pass, you're done for the week.
 - If any test reads FAILED, rerun ALL the tests. Be sure that the test sheets and films are loaded in the correct order. If any test fails again, power-cycle the system (see '6.2. Power-Cycling the System') and rerun all the tests.
 - If any test fails again, call your Hologic service representative.

6.5. Resetting the Handheld Barcode Scanner

Occasionally it is necessary to reset the handheld barcode scanner software to its default settings. If the barcode scanner stops reading barcodes properly, use the barcode labels provided in this section to reset it.

► To reset the handheld barcode scanner

1 Scan the 'Enter/Exit Configuration Mode' barcode shown below. When you scan the barcode, the scanner should beep three times.



2 Scan the 'Recall Defaults' barcode. When you scan the barcode, the scanner should beep once. This erases all previous settings and returns the barcode scanner to its factory default communications settings.



Part 7: Using DigitalNow and the R2 Patient ID System

- ► 7.1. Overview
- ► 7.2. DigitalNow Archiving
- 7.3. Using the R2 Patient ID System
- 7.4. Querying an External Database

Part 7 provides information about DigitalNow, which provides archiving features and the R2 Patient ID system.

7.1. Overview

R2 DMax supports the archiving of digitized film images and CAD results for supported DICOM views. You can enable and disable the archiving features by means of the Connectivity option on the Controls screen. At the time of installation, the features are set either to On or Off, based on your preference.

The Archiving feature requires a patient information database from which to draw the necessary patient information. This ensures that the images and CAD results are stored properly in the archive. Therefore, the Patient ID feature must be enabled and the system configured to communicate with a patient information database in order for the Archiving feature to function. In addition, the R2 system must be configured to communicate with the archive system.

If both Patient ID and Archiving are enabled, the images and any CAD results are sent to the archive system after the images are processed.

The archived images and CAD results, like any other record in the archive system, can then be retrieved and viewed on softcopy review workstations and/or printed.

7.2. DigitalNow Archiving

The workflow for DigitalNow archiving is as follows:

- 1 At any point prior to scanning film mammograms for a case, you must enter the R2 barcode on the separator sheet used for the case into the record for the appropriate patient in the patient information database.
- **2** Then, you can scan the films. The system retrieves the patient information from the patient database, and transfers the images to the archive system.

If Verification is set to 'On', you must verify the images are correct before they can be sent to the archive. For more information, see '3.9. Verifying Scanned Cases'.

Once the images are stored in the archive, they can be manipulated like any other record in the archive system – for example, you can send them to the softcopy digital review workstation. Keep in mind that once film mammograms are digitized and archived, they are treated like digital images and will not be sent to display units other than a softcopy digital review workstation.

If archiving is enabled, and no record in the patient database matches the R2 barcode, the system does not process the case. You can correct this by entering patient information in the R2 database after the case is scanned and reprocessing the case.

7.3. Using the R2 Patient ID System

For sites that do not have a patient database but want to use DigitalNow, Hologic provides the R2 Patient ID system.

The purpose of this patient identification database is to enter the patient information necessary to archive images and/or CAD results. You cannot query the database, view or manipulate records. The information remains in the R2 database for only 96 hours (four days) from time of entry.

The R2 Patient ID system can be used in conjunction with the DICOM Modality Worklist, Study Root, or Patient Root Queries (as described in '7.4. Querying an External Database' on page 88) to retrieve information from your Modality Worklist or PACS.

To enter patient information, your service representative must have configured your system to use the R2 Patient ID system. In addition, connectivity with the patient database must be enabled by a service engineer.

You need the following information:

- Database password: h0spital (the second character is a zero, not the letter 'O')
- IP address for the R2 DMax system (see the Controls, About screen)

Once you have this information you can enter patient information.

▲ **Note:** For sites with multiple film-scanning units: When you access the Patient ID database, make sure you access the R2 system that you will be using to scan the studies. Each system maintains its own database – you must enter the patient information into the database of the R2 system on which you plan to scan the images. Each film-scanning unit can be given a unique name during installation, and this name is displayed in the R2 Patient ID web page.

To enter patient information

1 Touch the **R2 Patient ID** button at the top of the screen to access the Patient ID system directly from the system.

OR

To access the Patient ID system from any computer on the same network as the R2 system, using any web browser, type the IP address for the processing unit followed by /id in the Address box, and press (-Enter). For example:

http://12.345.67.89/id

The browser will display the log-in window.

R2	Patient ID
	DMAX2
Password:	
© 2004-2008 R2 Technology, Inc.	login

2 Enter your Patient ID database password (h0spital), and click login.

\sim	
- K2	
	Patient ID
	SQA - DMAX1
Patient Information	
Last Name First Name	_
query	-)
Date of Birth (MM/DD/YYYY)	
Female -	
Site Patient ID / Medical Record # *	
query	
	_
Study Information	
Study Date (MM/DD/YYYY) *	
07 24 2008	
Accession Number	
query	
R2 Barcode *	
Referring Physician	—
Last Name First Name	
* - Required fields.	
	ear submit
© 2004-2008 Hologic, Inc.	

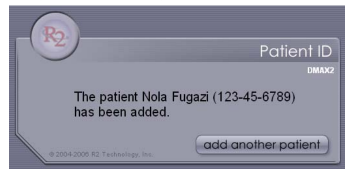
The system displays the patient information entry page:

▲ **Note:** The **query** buttons appear on this screen only if they were configured to appear by the Hologic service engineer when the system was installed. The service engineer can set any of the three **query** fields to be the default field when the screen appears.

- **3** Enter the patient information, Study Date, and R2 barcode number. Required fields are marked with an asterisk (*).
 - Enter the patient information either by typing it in, or by querying a PACS/ workstation (as described under '7.4. Querying an External Database' on page 88).
 - Enter the correct Study Date. (The Study Date defaults to the current date.)
 - Enter the R2 barcode number into the 'Accession Number' field by either scanning the barcode on the separator sheet, or typing it in.

▲ **Note:** Some archives generate a Study Instance UID (unique ID) that is used to identify a given study. If you query to retrieve patient information and then enter new information (or change the information), then the R2 system generates a new Study Instance UID that it will send to the PACS with the study results.

4 Click **submit**. The system confirms that you have added the patient.



You will not be able to verify that the information you enter is correct. If you believe you made a mistake, enter the information again. If an identical record is entered, it will simply overwrite the previous one.

Once you have submitted the patient information and R2 barcode, you can scan the case.

(A) Important! Be careful when entering the information. This is the mechanism that links the images to the correct patient. When viewing digitized images from an archive, always check the patient flash to ensure the case corresponds to the desired patient.

To enter multiple studies from different dates for the same patient, you will need to enter the studies separately using a different separator sheet (with a different R2 barcode) for each study. To avoid entering the patient data twice, use the back arrow on your web browser to return to the patient data entry page. (The keyboard shortcut is [Alt] + [left arrow].) At that point, you would need to change only the study date and accession number.

7.4. Querying an External Database

You can enter patient information onto the patient information entry page by querying a PACS/workstation (configured by the Hologic service engineer when the system was installed).

There are three types of queries; the type you can run was configured by the Hologic service engineer. The following types are available:

- **DICOM Patient Root Query**. If this was configured, only the **query** buttons in the Patient Information section of the patient information entry page are active. When you query by touching the **query** button to the right of the patient name fields, the system queries using only those fields. When you query by touching the **query** button to the right of 'Site Patient ID / Medical Record #' field, the system queries using only that field. With either button, the query result returns data for just the fields in the Patient Information section.
- **DICOM Study Root Query.** If this was configured, only the **query** button in the Study Information section of the patient information entry page is active. When you query by touching the **query** button to the right of the 'Accession Number' field, the system queries using only that field. The query returns data for all fields on the page.
- **DICOM Modality Worklist Query.** If this was configured, all three **query** buttons are active. When you query by touching any **query** button, the system queries using only the field(s) to its left. The query result returns data for all fields on the page; however, the data entered into 'Study Date (MM/DD/YYYY)' comes from the Scheduled Procedure Step Start Date field in the database.

▲ **Note:** These queries do not enter the R2 barcode number. You'll need to enter it as described in step 3 in the preceding procedure, '**7.3. Using the R2 Patient ID System**'.

If you leave the field used by the query blank, the query will return a list of all studies in the database. If you enter into the field one or more characters that appear in the data you want to match, the system will return a list of just the studies containing the matching data.

2			Patient	
The query returned multiple matches. Please select one of the following.				
Patient Name	MRN	Study Date	Accession #	
<u>1, LCC</u>	ID1LCC	05/21/2003	<u>^</u>	
<u>1, LMLO</u>	ID1LMLO	04/24/2003	=	
1, RCC	ID1RCC	04/19/2003	<u> </u>	
1, RML0	ID1RMLO	04/17/2003		
2, LCC LMLO.	ID2LCCLMLO	09/03/2003		
2, LCC RMLO.	ID2LCCRMLO	09/03/2003		
2, RCC LCC.	ID2RCCLCC	05/09/2003		
2, RCC LMLO.	ID2RCCLMLO	04/10/2003		
2, RCC RMLO.	ID2RCCRMLO	04/10/2003		
	ID2RML OL ML O	00/02/2002	× 1	

To select a study and copy its data onto the patient information entry page, touch its entry in the Patient Name column.

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