Horizon®



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User Guide For Windows 10

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Table of Contents

List of Tables xv 1: Overview 1 1.1 Indications for Use. 1 1.1.1 APEX Indications 1 1.1.2 IVA Indications 1 1.1.3 Body Composition Indications 1 1.1.4 Visceral Fat Software 2 1.1.5 IO-year Fracture Risk Indications 2 1.1.6 Hip Structure Analysis Indications 2 1.1.7 Single Energy (SE) Femur Exam Indications 3 1.2 Contraindications 3 1.4 Warnings and Cautions 3 1.4.1 EMI 3 1.4.2 Accessories 4 1.4.3 Coution 4 1.5 Labels 4 1.6 Symbols 7 1.7 Support Materials 8 1.7.1 QDR Reference Manual 8 1.7.2 Online Help 8 1.7.3 QDR Series Technical Specifications Manual 8 1.7.4 QDR Cyber-Security Information 8 1.7.3 QDR Series Technical Specifications Manual 8 1.7.4 QDR Cyber-Security Information 10 3.1 System Startup and Shutdown 9	Lis	ist of Figures	
1.1 Indications for Use	Lis	st of Tables	xv
1.1 Indications for Use. 1 1.1.1 APEX Indications. 1 1.1.2 IVA Indications. 1 1.1.3 Body Composition Indications 1 1.1.4 Visceral Fat Software. 2 1.1.5 10-year Fracture Risk Indications 2 1.1.6 Hip Structure Analysis Indications 2 1.1.7 Single Energy (SE) Femur Exam Indications 3 1.2 Contraindications. 3 1.4 Warnings and Cautions 3 1.4.1 EMI 3 1.4.2 Accessories 4 1.4.3 Caution 4 1.4.4 Accessories 4 1.4.5 Labels 4 1.6 Symbols 7 7.7 Support Materials 8 1.7.1 QDR Reference Manual 8 1.7.3 QDR Series Technical Specifications Manual 8 1.7.4 QDR Cyber-Security Information 8 1.7.4 QDR Cyber-Security Information 11 2. System Startup 11	1: (Overview	1
1.1.2 IVA Indications 1 1.1.3 Body Composition Indications 1 1.1.4 Visceral Fat Software. 2 1.1.5 10-year Fracture Risk Indications. 2 1.1.6 Hip Structure Analysis Indications. 2 1.1.7 Single Energy (SE) Femur Exam Indications. 3 1.2 Contraindications. 3 1.4 Warnings and Cautions 3 1.4.1 ECR egulations. 3 1.4.1 EMI 3 1.4.2 Accessories 4 1.4.3 Caution. 4 1.5 Labels. 4 1.6 Symbols. 7 1.7 ODR Reference Manual 8 1.7.1 QDR Reference Manual 8 1.7.2 Online Help 8 1.7.3 QDR Series Technical Specifications Manual 8 1.7.4 QDR Cyber-Security Information 8 1.7.4 QDR Cyber-Security Information 11 2.5 System Startup and Shutdown 11 2.1 System Startup	1.1		
1.1.3 Body Composition Indications 1 1.1.4 Visceral Fat Software 2 1.1.5 10-year Fracture Risk Indications 2 1.1.6 Hip Structure Analysis Indications 2 1.1.7 Single Energy (SE) Femur Exam Indications 3 1.2 Contraindications 3 1.3 IEC Regulations 3 1.4 Warnings and Cautions 3 1.4.1 EMI 3 1.4.2 Accessories 4 1.4.3 Caution 4 1.5 Labels 4 1.6 Symbols 7 1.7 Support Materials 8 1.7.1 QDR Reference Manual 8 1.7.2 Online Help 8 1.7.3 QDR Cyber-Security Information 8 1.7.4 QDR Cyber-Security Information 8 1.7.4 QDR Cyber-Security Information 11 2.1 System Startup and Shutdown 11 2.1 System Startup 11 3.2 Auto QC 13		1.1.1 APEX Indications	
1.1.4 Visceral Fat Software		1.1.2 IVA Indications	
1.1.5 10-year Fracture Risk Indications 2 1.1.6 Hip Structure Analysis Indications 2 1.1.7 Single Energy (SE) Femur Exam Indications 3 1.2 Contraindications 3 1.3 IEC Regulations 3 1.4 Warnings and Cautions 3 1.4.1 EMI 3 1.4.2 Accessories 4 1.4.3 Caution 4 1.5 Labels 4 1.6 Symbols 7 1.7 Support Materials 8 1.7.1 QDR Reference Manual 8 1.7.2 Online Help 8 1.7.3 QDR Series Technical Specifications Manual 8 1.7.4 QDR Cyber-Security Information 8 1.8 Main Window 9 2: System Startup and Shutdown 11 11 2: System Startup Control Procedure 13 3.1 System Test 13 3.2 Auto QC 13 3.3 Automatic Body Composition Calibration 14		1.1.3 Body Composition Indications	
1.1.6 Hip Structure Analysis Indications 2 1.1.7 Single Energy (SE) Femur Exam Indications 3 1.2 Contraindications 3 1.3 IEC Regulations 3 1.4 Warnings and Cautions 3 1.4 EMI 3 1.4.1 EMI 3 1.4.2 Accessories 4 1.4.3 Caution 4 1.4.4 Counton 4 1.5 Labels 4 1.6 Symbols 7 7 Support Materials 8 1.7.1 QDR Reference Manual 8 1.7.2 Online Help 8 1.7.3 QDR Cyber-Security Information 8 1.7.4 QDR Cyber-Security Information 8 1.8 Main Window 9 2: System Startup and Shutdown 11 2.1 System Startup 11 2.2 System Startup 13 3.1 System Test 13 3.2 Auto QC 13		1.1.4 Visceral Fat Software	2
1.1.7 Single Energy (SE) Femur Exam Indications 3 1.2 Contraindications 3 1.3 IEC Regulations 3 1.4 Warnings and Cautions 3 1.4.1 EMI 3 1.4.2 Accessories 4 1.4.3 Caution 4 1.4.3 Caution 4 1.4.5 Labels 4 1.6 Symbols 7 1.7 Support Materials 8 1.7.1 QDR Reference Manual 8 1.7.2 Online Help 8 1.7.3 QDR Series Technical Specifications Manual 8 1.7.4 QDR Cyber-Security Information 8 1.7.4 QDR Cyber-Security Information 8 1.8 Main Window 9 2: System Startup and Shutdown 11 2.1 System Startup 11 2.1 System Startup 13 3.1 System Startup 13 3.2 Auto QC 13 3.3 Automatic Body Composition Calibra		1.1.5 10-year Fracture Risk Indications	2
1.2 Contraindications		1.1.6 Hip Structure Analysis Indications	2
1.3 IEC Regulations 3 1.4 Warnings and Cautions 3 1.4.1 EMI 3 1.4.2 Accessories 4 1.4.3 Caution 4 1.4.1 EMI 3 1.4.2 Accessories 4 1.4.3 Caution 4 1.4.3 Caution 4 1.5 Labels 4 1.6 Symbols 7 1.7 Support Materials 8 1.7.1 QDR Reference Manual 8 1.7.2 Online Help 8 1.7.3 QDR Speires Technical Specifications Manual 8 1.7.4 QDR Cyber-Security Information 8 1.8 Main Window 9 2: System Startup and Shutdown 11 2.1 System Startup and Shutdown 11 2.2 System Startup 11 3.1 System Test 13 3.2 Auto QC 13 3.3 Automatic Body Composition Calibration 14 <td< td=""><td></td><td>1.1.7 Single Energy (SE) Femur Exam Indications</td><td></td></td<>		1.1.7 Single Energy (SE) Femur Exam Indications	
1.4 Warnings and Cautions 3 1.4.1 EMI 3 1.4.2 Accessories 4 1.4.3 Caution 4 1.4.4 Caution 4 1.4.5 Labels 4 1.6 Symbols 7 7 Support Materials 8 1.7.1 QDR Reference Manual 8 1.7.2 Online Help 8 1.7.3 QDR Series Technical Specifications Manual 8 1.7.4 QDR Cyber-Security Information 8 1.7.4 QDR Cyber-Security Information 8 1.8 Main Window 9 2: System Startup and Shutdown 11 2.1 System Startup and Shutdown 11 2.2 System Shutdown 11 3.1 System Test 13 3.1 System Test 13 3.2 Auto QC 13 3.3 Automatic Body Composition Calibration 14 4: Patient Records 15 4.1 Retrieving a Patient Record 15 <	1.2	Contraindications	
1.4.1 EMI	1.3	IEC Regulations	
14.2 Accessories 4 14.3 Caution 4 14.3 Caution 4 1.5 Labels 4 1.6 Symbols 7 1.7 Support Materials 8 1.7.1 QDR Reference Manual 8 1.7.2 Online Help 8 1.7.3 QDR Series Technical Specifications Manual 8 1.7.4 QDR Cyber-Security Information 8 1.8 Main Window 9 2: System Startup and Shutdown 11 2.1 System Startup 11 2.2 System Startup 11 2.1 System Startup 11 2.2 System Test 13 3.1 System Test 13 3.2 Auto QC 13 3.3 Automatic Body Composition Calibration 14 4: Patient Records 15 4: Retrieving a Patient Record 15 4: Creating a Patient Record 15 4: Batieting a Patient Record 15 </td <td>1.4</td> <td>0</td> <td></td>	1.4	0	
14.3 Caution 4 1.5 Labels 4 1.6 Symbols 7 1.7 Support Materials 8 1.7.1 QDR Reference Manual 8 1.7.2 Online Help 8 1.7.3 QDR Series Technical Specifications Manual 8 1.7.4 QDR Cyber-Security Information 8 1.8 Main Window 9 2: System Startup and Shutdown 11 2.1 System Startup 11 2.2 System Startup 11 2.1 System Startup 11 2.2 System Startup 11 2.3 System Test 13 3.1 System Test 13 3.2 Auto QC 13 3.3 Automatic Body Composition Calibration 14 4: Patient Records 15 4: Retrieving a Patient Record 15 4: Creating a Patient Record 15 4: Bratient Record 15 4: Bratient Record 15 <		1.4.1 EMI	
1.5 Labels		1.4.2 Accessories	4
1.6 Symbols 7 1.7 Support Materials 8 1.7.1 QDR Reference Manual 8 1.7.2 Online Help 8 1.7.3 QDR Series Technical Specifications Manual 8 1.7.4 QDR Cyber-Security Information 8 1.8 Main Window 9 2: System Startup and Shutdown 11 2.1 System Startup 11 2.2 System Startup 11 2.1 System Test 13 3.1 System Test 13 3.2 Auto QC 13 3.3 Automatic Body Composition Calibration 14 4: Patient Records 15 4.1 Retrieving a Patient Record 15 4.2 Creating a Patient Record <		1.4.3 Caution	4
1.7 Support Materials 8 1.7.1 QDR Reference Manual 8 1.7.2 Online Help 8 1.7.3 QDR Series Technical Specifications Manual 8 1.7.4 QDR Cyber-Security Information 8 1.8 Main Window 9 2: System Startup and Shutdown 11 2.1 System Startup and Shutdown 11 2.2 System Startup 11 2.1 System Startup 11 2.2 System Startup 11 2.1 System Startup 11 2.2 System Startup 11 2.3 System Startup 11 2.4 System Startup 11 2.5 System Test 13 3.1 System Test 13 3.2 Auto QC 13 3.3 Automatic Body Composition Calibration 14 4: Patient Records 15 4.1 Retrieving a Patient Record 15 4.2 Creating a Patient Record 15 4.3 Editing a Patient Recor	1.5		
1.7.1 QDR Reference Manual 8 1.7.2 Online Help 8 1.7.3 QDR Series Technical Specifications Manual 8 1.7.4 QDR Cyber-Security Information 8 1.8 Main Window 9 2: System Startup and Shutdown 11 2.1 System Startup 11 2.2 System Startup and Shutdown 11 2.1 System Startup 11 2.2 System Startup 11 2.1 System Startup 11 2.2 System Startup 11 2.3 System Startup 11 3.4 System Test 13 3.1 System Test 13 3.2 Auto QC 13 3.3 Automatic Body Composition Calibration 14 4: Patient Records 15 4.1 Retrieving a Patient Record 15 4.2 Creating a Patient Record 15 4.3 Editing a Patient Record 15	1.6	Symbols	7
1.7.2 Online Help 8 1.7.3 QDR Series Technical Specifications Manual 8 1.7.4 QDR Cyber-Security Information 8 1.8 Main Window 9 2: System Startup and Shutdown 11 2.1 System Startup 11 2.2 System Startup 11 2.1 System Startup 11 2.2 System Startup 11 2.3 System Startup 11 3.4 System Test 13 3.1 System Test 13 3.2 Auto QC 13 3.3 Automatic Body Composition Calibration 14 4: Patient Records 15 4.1 Retrieving a Patient Record 15 4.2 Creating a Patient Record 15 4.3 Editing a Patient Record 15	1.7	11	
1.7.3 QDR Serie's Technical Specifications Manual 8 1.7.4 QDR Cyber-Security Information 8 1.8 Main Window 9 2: System Startup and Shutdown 11 2.1 System Startup 11 2.2 System Startup and Shutdown 11 2.1 System Startup 11 2.2 System Startup 11 3.1 System Shutdown 11 3.2 Quality Control Procedure 13 3.1 System Test 13 3.2 Auto QC 13 3.3 Automatic Body Composition Calibration 14 4: Patient Records 15 4.1 Retrieving a Patient Record 15 4.2 Creating a Patient Record 15 4.3 Editing a Patient Record 15		1.7.1 QDR Reference Manual	
1.7.4 QDR Cyber-Security Information 8 1.8 Main Window 9 2: System Startup and Shutdown 11 2.1 System Startup 11 2.2 System Shutdown 11 3.1 System Startup 13 3.2 Auto QC 13 3.3 Automatic Body Composition Calibration 14 4: Patient Records 15 4.1 Retrieving a Patient Record 15 4.2 Creating a Patient Record 15 4.3 Editing a Patient Record 15		1	
1.8 Main Window		1	
2: System Startup and Shutdown 11 2.1 System Startup 11 2.2 System Shutdown 11 3: Quality Control Procedure 13 3.1 System Test 13 3.2 Auto QC 13 3.3 Automatic Body Composition Calibration 14 4: Patient Records 15 4.1 Retrieving a Patient Record 15 4.2 Creating a Patient Record 15 4.3 Editing a Patient Record 15		· · ·	
2.1 System Startup 11 2.2 System Shutdown 11 3: Quality Control Procedure 13 3.1 System Test 13 3.2 Auto QC 13 3.3 Automatic Body Composition Calibration 14 4: Patient Records 15 4.1 Retrieving a Patient Record 15 4.2 Creating a Patient Record 15 4.3 Editing a Patient Record 15	1.8	Main Window	
2.1 System Startup 11 2.2 System Shutdown 11 3: Quality Control Procedure 13 3.1 System Test 13 3.2 Auto QC 13 3.3 Automatic Body Composition Calibration 14 4: Patient Records 15 4.1 Retrieving a Patient Record 15 4.2 Creating a Patient Record 15 4.3 Editing a Patient Record 15	2: S	System Startup and Shutdown	11
3: Quality Control Procedure 13 3.1 System Test 13 3.2 Auto QC 13 3.3 Automatic Body Composition Calibration 14 4: Patient Records 15 4.1 Retrieving a Patient Record 15 4.2 Creating a Patient Record 15 4.3 Editing a Patient Record 15			
3.1 System Test 13 3.2 Auto QC 13 3.3 Automatic Body Composition Calibration 14 4: Patient Records 15 4.1 Retrieving a Patient Record 15 4.2 Creating a Patient Record 15 4.3 Editing a Patient Record 15	2.2	System Shutdown	
3.1 System Test 13 3.2 Auto QC 13 3.3 Automatic Body Composition Calibration 14 4: Patient Records 15 4.1 Retrieving a Patient Record 15 4.2 Creating a Patient Record 15 4.3 Editing a Patient Record 15	3: Ç	Quality Control Procedure	13
3.3 Automatic Body Composition Calibration		-	
3.3 Automatic Body Composition Calibration	3.2	Auto QC	
 4.1 Retrieving a Patient Record	3.3	Automatic Body Composition Calibration	
 4.1 Retrieving a Patient Record	4: P	Patient Records	15
 4.2 Creating a Patient Record			
4.3 Editing a Patient Record		8	
0		8	
		8	

4.5	Confirming Patient Information	
	4.5.1 Biographical Information	
	4.5.2 Exam Information	
5: P	Performing an Exam	17
5.1	Patient Interview	
5.2	Patient Preparation	
5.3	Patient Selection	
5.4	Choosing the Scan Type	
5.5	Performing a Scan	
6: N	Manual Scan Analysis	19
6.1	Analysis Step Buttons	
6.2	Toolboxes	
	6.2.1 Bone Map Toolbox	
	6.2.2 Lines Toolbox (Spine)	
	6.2.3 Neck Toolbox (Hip)	
	6.2.4 Results Toolbox (Whole Body)	
	6.2.5 Sub Region Toolbox	
	6.2.6 Toolbox Controls	
6.3	Brightness/Contrast Control	24
7: A	AP Lumbar Spine Examination	25
	Positioning the Patient	
7.1	rositioning the ratient	
7.1 7.2	Positioning the C-arm	
	8	
7.2	Positioning the C-arm Starting the AP Lumbar Spine Scan 7.3.1 Repositioning the Scan (if necessary)	
7.2	Positioning the C-arm Starting the AP Lumbar Spine Scan 7.3.1 Repositioning the Scan (if necessary) Analyzing the Scan	
7.2 7.3	Positioning the C-arm.Starting the AP Lumbar Spine Scan7.3.1Repositioning the Scan (if necessary).Analyzing the Scan7.4.1One-Time™ Auto Analysis	
7.2 7.3	Positioning the C-arm.Starting the AP Lumbar Spine Scan	
7.2 7.3 7.4	Positioning the C-arm.Starting the AP Lumbar Spine Scan7.3.1Repositioning the Scan (if necessary).Analyzing the Scan7.4.1One-Time™ Auto Analysis	26 27 28 29 29 30
7.27.37.47.57.6	Positioning the C-arm.Starting the AP Lumbar Spine Scan	
7.27.37.47.57.6	Positioning the C-arm	26 27 28 29 29 30 30 31
 7.2 7.3 7.4 7.5 7.6 8: H 	Positioning the C-arm	
7.2 7.3 7.4 7.5 7.6 8: H 8.1	Positioning the C-arm	26 27 28 29 30 30 30 31 31 32 33
7.2 7.3 7.4 7.5 7.6 8: H 8.1	Positioning the C-arm	26 27 28 29 30 30 30 31 31 31 32 33 33
7.2 7.3 7.4 7.5 7.6 8: H 8.1	Positioning the C-arm	26 27 28 29 30 30 30 31 31 31 32 33 33
7.2 7.3 7.4 7.5 7.6 8: H 8.1 8.2	Positioning the C-arm	26 27 28 29 30 30 30 30 30 33 31 31 32 33 33 33 33
7.2 7.3 7.4 7.5 7.6 8: H 8.1	Positioning the C-arm	26 27 28 29 30 30 30 30 30 30 30 33 33 33 33
7.2 7.3 7.4 7.5 7.6 8: H 8.1 8.2	Positioning the C-arm	26 27 28 29 29 30 30 30 31 31 32 33 33 33 33 33 33 33 33 33 33 33 33
7.2 7.3 7.4 7.5 7.6 8: H 8.1 8.2	Positioning the C-arm	26 27 28 29 30 30 30 30 30 30 30 30 33 33 33 33

8.4	Analyzing the Scan	
	8.4.1 One-Time Auto Analysis	
8.5	SE (Single Energy) Femur Examination	
	8.5.1 Positioning for the SE Femur scan	
	8.5.2 Positioning for the SE Femur Scan after a Hip Scan	
	8.5.3 Starting the SE Femur Scan	
	8.5.4 Analysis of SE Femur Scan	
	8.5.5 Visual Assessment	
	8.5.6 Image Toolbox	
	8.5.7 Analysis Tools Control Tab	
	8.5.8 Assessment	
8.6	References	
	8.6.1 Dual-Hip Scans	
8.7	Generating and Printing Reports	
9: Fo	orearm Examination	47
9.1	Measuring the Patient's Forearm	
9.2	Positioning the Patient	
	9.2.1 Positioning the Patient for Supine Forearm Examinations	
9.3	Positioning the C-arm	
9.4	Starting the Forearm Scan	
	9.4.1 Repositioning the Scan (if necessary)	
	9.4.2 Repositioning the Patient (if necessary)	
9.5	Analyzing the Scan	
	9.5.1 Entering the Forearm Length	
	9.5.2 Defining the Global ROI	
	9.5.3 Viewing the Bone Map	
	9.5.4 Checking the MID/UD Region	
9.6	Viewing Results	
9.7	Exiting Analysis	
9.8	Generating and Printing Reports	
10: V	Whole Body Examination	57
10.1	Body Composition Analysis	
10.2		
10.3	0	
10.4	· ·	
10.5		
	10.5.1 Body Composition Analysis	
	10.5.2 Default Placement of Whole Body Regions	
	10.5.3 Adjusting A/G Regions (if necessary)	
	10.5.4 Visceral Adipose Tissue	
	10.5.5 Adjusting VAT Regions (if necessary)	
	10.5.6 VAT References	

10.6	Viewing Results	67
	10.6.1 Rulers	67
10.7	Exiting Analysis	68
10.8	User-Defined SubRegions	68
10.9	Resolving Asymmetric Results with Reflection	69
10.10	Enable NHANES BCA	70
10.11	Generating and Printing Reports	70
11: 5	Supine AP/Lateral Spine BMD Examination (Horizon A)	71
11.1	Table Safety Feature	71
11.2	Positioning for the AP/Lateral Scan	
11.3	Starting the AP Scan	
11.4	Analyzing the AP Scan	
11.5	Starting the Lateral Scan	
11.6	Analyzing the Lateral Scan	
	11.6.1 Defining the Global ROI	
	11.6.2 Adjusting the Vertebral Boundaries	
	11.6.3 Adjusting the Vertebral Bodies	
	11.6.4 Viewing the Bone Map	
	11.6.5 Adjusting the Mid Regions	
11.7	Viewing Results	
11.8	Exiting Analysis	
11.9	Generating and Printing Reports	
	Decubitus Lateral Spine BMD Examination	
12.1	Performing and Analyzing the AP Scan	
12.2	Positioning the Patient for the Decubitus Lateral Scan	
12.3	Positioning the C-arm for the Decubitus Lateral Scan	
12.4	Starting the Decubitus Lateral Scan	
	12.4.1 Repositioning the Scan (if necessary)	
12.5	Analyzing the Scan	
	12.5.1 Defining the Global ROI	
	12.5.2 Adjusting the Vertebral Boundaries	
	12.5.3 Adjusting the Vertebral Bodies	
10 (12.5.4 Viewing the Bone Map Viewing Results	
12.6 12.7	Exiting Analysis	
	Generating and Printing Reports	
12.8	Generating and Finning Reports	
	VA, IVA HD Imaging on Horizon C, W; IVA Imaging on Horizon Ci, Wi	
13.1	Choosing the Scan Type	
13.2	Positioning the Patient for the AP IVA Scan	
13.3	Starting the AP IVA Scan	
13.4	Positioning the Patient and C-arm for the Lateral IVA Scan	
13.5 13.6	Starting the Lateral IVA Scan IVA Analysis for Lateral IVA Scan	
136	IVA Analysis for Lateral IVA Scan	90

14:]	IVA and IVA HD Imaging on a Horizon A System	91
14.1	Positioning the Patient	
14.2	Choosing the Scan Type	
14.3	Starting the AP IVA Scan	92
14.4	Starting the Lateral IVA Scan	
14.5	IVA Analysis for Lateral IVA Scan	
15: I	IVA with BMD Examination	95
15.1	Positioning the Patient	
15.2	Choosing the Scan Type	
15.3	Perform the AP IVA Scan	
15.4	Perform and Analyze the AP BMD Scan	
15.5	Perform the Lateral BMD Scan	96
15.6	Perform the Lateral IVA Scan	96
15.7	IVA Analysis	96
16: I	IVA Analysis	97
16.1	Viewer Window, Left Panel	
16.2	Viewer Window, Middle Panel	
16.3	Viewer Window, Right Panel	
16.4	Viewer Window, Left Panel, Visual Tools Tab	
16.5	Viewer Window, Left Panel Analysis Tools Tab	
16.6	Image Controls	
16.7	DE Scan	
17: I	Interpreting the IVA Image	103
18: I	Markers	105
18.1	Using Markers	
	18.1.1 Flipping the Image	
	18.1.2 Adding Markers	
	18.1.3 Selecting Markers	
18.2	Result Block	
	18.2.1 Result Block for Lateral IVA Scans	
	18.2.2 Result Block for AP IVA Scans	
18.3	Printing	
	18.3.1 Print Report	
	18.3.2 Print Image	
19: (Compare and Follow-up	111
19.1	Restore a Baseline or Prior Scan	
19.2	Evaluate the Baseline or Prior Scan	
19.3	Perform the Follow-Up Scan	
19.4	Analyze the Follow-up Scan Using Compare Analysis	
19.5	Generate the Rate of Change Report	

20: I	Report	s	113
20.1	_	t Information Blocks	
20.2	Edit C	omments	
20.3	Rate o	f Change Report	
	20.3.1	Remove Asterisks (*) and Pound Signs (#) from Reports	
	20.3.2	Create Hip Pairs for Dual Hip Rate of Change Reports	
20.4		Composition Reports	
	20.4.1		
	20.4.2	BCA Rate of Change Results	
	20.4.3	Body Composition Reports and Reference Database Comparisons	
20.5	Pediat	ric Reports	
20.6	DICO	M Report	
	20.6.1	Select a DICOM BMD Report Type	
	20.6.2	View Scan Details and Enter Patient Biography Fields	
	20.6.3	Enter the Accession Number and User Defined Entries	
	20.6.4	Preview a DICOM Report	
	20.6.5	Print a DICOM Report	
	20.6.6	Save a DICOM Report	
	20.6.7	Send a DICOM Report	
	20.6.8	Sort the Scan List	
	20.6.9	View the Queue	
	20.6.10) Close a DICOM Report	
20.7	DxRep	port	
	20.7.1	Create a DxReport	
21: I	nterpr	reting Results	123
21.1	About	FRAX	
21.2	FRAX	Limiting Criteria	
	21.2.1	Previous hip or vertebral fracture	
	21.2.2	Treatment for osteoporosis	
	21.2.3	Premenopausal woman	
	21.2.4	Configuring FRAX	
21.3	About	10-year Fracture Risk - All Countries	
21.4	FRAX	Risk Factors	
21.5	Refere	nces	
22: 5	Scans		129
00.1	 	0	100

	129
Archive Scans	
Locate Scans	
Restore Scans	
Copy Scans	
Query/Retrieve Scans	
23: Perform System Backup	
	Archive Scans Locate Scans Restore Scans Copy Scans Query/Retrieve Scans

24: 0	Clean the System	133
24.1	Clean the QDR and Computer Components	
24.2	Clean the Table Pad	
24.3	Disinfect the Table Pad	
24.4	Clean Accidental Spills	
25: I	Emergency Procedures	135
25.1	Power Failure	
	25.1.1 Shutting Down	
25.2	Failure During Operation	
25.3	Loss of Power	
26: I	Dose Area Product Meter	137
26.1	Turning the DAP Meter On and Off	
27: U	Utilities	139
27.1	System Configuration	
27.2	Usage	
27.3	Database Tools	
	27.3.1 Patient Management	
	27.3.2 Export	
	27.3.3 Import	
	27.3.4 Reconcile	
	27.3.5 Patient Callback	
	27.3.6 Auto Baseline Utility	
27.4	Scan File Look	
27.5	Scan File Plot	
27.6	Emergency Motion	
27.7	AP Reposition	
27.8	Factory Utilities	
27.9	Service Utilities	
27.10) Reference Curve	
	27.10.1 Editor	
	27.10.2 Add Ethnicity	
07.14	27.10.3 Restore	
	I Rebuild Archive Index	
27.12	2 Install Options	141
	Reference Curve	
28.1	Starting Reference Curve Editor	
28.2	Viewing Reference Curve Data	
28.3	Creating New Reference Curve Records	
28.4	Copying a Reference Curve Record	

28.5

28.7	Adding	; New Ethnic Groups	
28.8	Restori	ng Reference Curve Database	
29: I	DICOM	I Option	149
29.1		ure DICOM Option	
	29.1.1	DICOM Configuration Tabs	149
	29.1.2	Modality Worklist	149
	29.1.3	Query Parameters	
	29.1.4	Auto Query Interval	
	29.1.5	Query Retry Parameters	
	29.1.6	Purge Interval	
	29.1.7	Input From File	
	29.1.8	Worklist Provider	
	29.1.9	Map Worklist Fields	
	29.1.10	DICOM Send Destinations	
	29.1.11	DICOM Storage Commitment Destinations	
	29.1.12	DICOM Query/Retrieve Destinations	
	29.1.13	Host	
30: F	RAX F	AQs	171

List of Figures

Figure 1: Horizon Main Window	9
Figure 2: Spine Phantom Position	13
Figure 3: AP Lumbar Spine Positioning	
Figure 4: Leg Positions	
Figure 5: AP Lumbar Spine	
Figure 6: Spine Repositioning	
Figure 7: AP Lumbar Spine Analysis	
Figure 8: Left Hip Positioning without Hip Autopositioning	
Figure 9: Right Hip Positioning without Hip Autopositioning	
Figure 10: Foot Positioning (Dual Hip)	
Figure 11: Left Hip Positioning with Hip Autopositioning	
Figure 12: Hip Repositioning	
Figure 13: Properly Analyzed Hip Scan	
Figure 14: Left SE Femur Positioning	
Figure 15: Measuring the Left Forearm	
Figure 16: Left Forearm Positioning	
Figure 17: Laser Position on Left Arm	
Figure 18: Right Forearm Positioning	
Figure 19: Laser Position on Right Arm	
Figure 20: Patient On/Off Position for Left Supine Forearm	
Figure 21: Patient Position for Left Supine Forearm	51
Figure 22: Patient Position for Right Supine Forearm	51
Figure 23: Forearm Repositioning	
Figure 24: Forearm ROI	
Figure 25: Forearm Bone Map	
Figure 26: Forearm MID/UD	
Figure 27: Whole Body Positioning	
Figure 28: Whole Body Analysis (Horizontal and Lower Pelvic Divider Lines)	60
Figure 29: Whole Body Analysis (Vertical Lines)	
Figure 30: A/G Regions	63
Figure 31: VAT Regions Pictured as CT Slice	
Figure 32: Abdominal Tissue Features	
Figure 33: VAT Regions	
Figure 34: Asymmetric Results Warning	
Figure 35: AP/Lateral Spine Positioning	
Figure 36: Lateral Spine	
Figure 37: Lateral ROI	74
Figure 38: Vertebral Boundaries	75
Figure 39: Vertebral Bodies	76
Figure 40: Lateral Bone Map	
Figure 41: Decubitus Lateral Positioning	
Figure 42: Spine Position	

Figure 43: Decubitus Spine	81
Figure 44: Decubitus Repositioning	
Figure 45: Lateral ROI	
Figure 46: Adjusting Vertebral Boundaries	
Figure 47: Vertebral Bodies	
Figure 48: Lateral Bone Map	
Figure 49: AP IVA Positioning	
Figure 50: Lateral IVA Positioning	
Figure 51: Spine Position	90
Figure 52: AP IVA Positioning	91
Figure 53: Lateral IVA Positioning	93
Figure 54: IVA Viewer Window	97
Figure 55: DE Scan 1 (Top of Region)	101
Figure 56: DE Scan 2 (Top to Bottom of Region)	
Figure 57: The Human Spine	
Figure 58: Vertebrae Deformities	104
Figure 59: Marker Placement	
Figure 60: Adding a Marker	
Figure 61: Selecting a Marker	
Figure 62: Marker Data in the Result Block	106
Figure 63: Result Block	
Figure 64: Report Blocks	113
Figure 65: Advanced Body Composition Report	116
Figure 66: Advanced Body Composition Rate of Change Report	117
Figure 67: Pediatric Spine Report	
Figure 68: Circuit Breaker and Indicator	135

List of Tables

Table 1: Symbols	7
Table 2: Analysis Step Buttons	19
Table 3: Global ROI Toolbox	21
Table 4: Bone Map Toolbox	21
Table 5: Neck Toolbox (Hip)	
Table 6: Neck Toolbox (Hip)	
Table 7: Sub Region Toolbox	
Table 8: Toolbox Controls	23
Table 9: Image Toolbox	41
Table 10: Assessment	
Table 11: Adding and Modifying Rulers	
Table 12: Display Options	
Table 13: Change Annotations	44
Table 14: Atypical Femoral Fracture Assessment	44
Table 15: Ulnar Styloid Process	54
Table 16: Viewer Window, Left Panel	97
Table 17: Viewer Window, Middle Panel	
Table 18: Viewer Window, Right Panel	
Table 19: Viewer Window, Left Panel, Visual Tools Tab	
Table 20: Assessment Area	
Table 21: Display Options Area	
Table 22: Change Annotations Area	100
Table 23: Image Controls	
Table 24: Result Block Labels	
Table 25: Result Block for the AP IVA Scan	
Table 26: Follow-up Scan Procedures	
Table 27: Advanced Body Composition Report Fields	116
Table 28: Advanced Body Composition Graph Fields	116
Table 29: Advanced Body Composition Rate of Change Report Fields	117
Table 30: Advanced Body Composition Rate of Change Graph Fields	
Table 31: FRAX Risk Factors	
Table 32: Reference Curve Description Fields	145
Table 33: Query Parameters	
Table 34: Auto Query Interval	
Table 35: Query Retry Parameters	152
Table 36: Purge Interval	
Table 37: Input From File	
Table 38: Worklist Provider Features	154
Table 39: Worklist Provider	
Table 40: Performed Procedure Step	
Table 41: Provider	
Table 42: DICOM Send Destinations	157

Table 43: DICOM Send Destinations	
Table 44: DICOM Send Configuration	
Table 45: DICOM Storage Commitment Destinations	
Table 46: Storage Commitment Destination Editing	
Table 47: Storage Commitment Configuration	
Table 48: DICOM Query/Retrieve Destinations	
Table 49: Add / Edit DICOM Query/Retrieve Destination	
Table 50: Query/Retrieve Configuration	
Table 51: Host Configuration	

Chapter 1 Overview

1.1 Indications for Use

1.1.1 APEX Indications

The APEXTM for QDRTM X-ray Bone Densitometers is indicated for the estimation of bone mineral density (BMD), comparison of measured variables obtained from a given QDR scan to a database of reference values, the estimation of fracture risk, vertebral deformity assessment, body composition analysis, and discrimination of bone from prosthetics using the Hologic QDR X-ray Bone Densitometers.

1.1.2 IVA Indications

IVA scans are intended for the visualization or quantitative assessment of vertebral bone deformities. IVA also allows the visualization of abdominal aortic calcification, and, if present, clinical correlation may be advised since abdominal aortic calcification may be associated with cardiovascular disease.

1.1.3 Body Composition Indications

The Hologic Whole Body DXA Reference Database software used on Hologic QDR bone densitometers measures the:

- regional and whole body bone mineral density,
- lean and fat tissue mass, and
- calculates derivative values of:
 - bone mineral content
 - area
 - soft tissue mass
 - regional soft tissue mass
 - total soft tissue mass
 - fat free mass
 - regional and total soft tissue mass ratios

- % fat, regional
- % fat, total body
- % fat, android
- % fat, gynoid
- % fat, android/gynoid ratio
- body mass index
- The values can be displayed in user-defined statistical formats and trends with color image mapping, and compared to reference populations at the sole discretion of the health care professional.

These body composition values are useful to health care professionals in their management of diseases and conditions where the disease and conditions itself, or its treatment, can affect the relative amounts of fat and lean tissue. The Hologic Whole Body DXA Reference Database software does not diagnose disease, recommend treatment regimens, or quantify treatment effectiveness. Only the health care professional can make these judgments. Some of the diseases (and conditions) for which body composition values are useful include chronic renal failure, anorexia nervosa, obesity, AIDS/HIV, and cystic fibrosis. DXA body composition is a useful alternative to hydrostatic weighting and skin fold measurements.

1.1.4 Visceral Fat Software

The Hologic Visceral Fat Software used on Hologic Horizon[®] bone densitometer total body scans estimates the visceral adipose tissue (visceral fat) content within the android region in an adult male or female population, excluding pregnant women. The content that is estimated is the Visceral Fat Area, Visceral Fat Mass, and Visceral Fat Volume. These values can be displayed in user-defined statistical formats and trends.

The estimated visceral fat content is useful to health care professionals in their management of diseases/conditions where the disease/conditions itself, or its treatment, can affect the relative amounts of visceral fat content in the android region.



Note

The Hologic Visceral Fat Software does not diagnose disease, recommend treatment regimens, or quantify treatment effectiveness. Only the health care professional can make these judgments.

Some of the diseases/conditions for which visceral fat estimation is useful include hypertension, impaired fasting glucose, impaired glucose tolerance, diabetes mellitus, dyslipidemia, and metabolic syndrome.

1.1.5 10-year Fracture Risk Indications

Femoral neck BMD and clinical risk factors are used to estimate 10-year risk of hip fracture and 10-year risk of major osteoporotic fracture using the World Health Organization (WHO) algorithm (FRAX®) in adults. The physician may use the 10-year fracture risk, along with the physician's knowledge of patient history, and apply medical expertise and best practice clinical judgment to determine if therapeutic intervention is indicated.

1.1.6 Hip Structure Analysis Indications

The Hip Structure Analysis (HSA®) for QDR X-ray Bone Densitometers uses data from conventional Dual Energy X-ray Absorptiometry (DXA) scans to measure the distribution of bone mineral mass at specific cross sections of the hip and allows the physician to estimate structural properties of the hip, such as CSA, CSMI, Z and Buckling Ratio.

1.1.7 Single Energy (SE) Femur Exam Indications

Single Energy (SE) femur exams are used to visualize focal reaction or thickening along the lateral cortex of the femoral shaft, which may be accompanied by a transverse radiolucent line. Clinical correlation is advised as these features may be consistent with atypical femur fractures, a complication associated with long term use of antiresorptive therapy.

1.2 Contraindications

Pregnancy and the use of contrast agents within the past 7 days are contraindicated. See *Patient Interview* on page 17 for more information.

1.3 IEC Regulations

QDR Series X-ray Bone Densitometers comply with the requirements of IEC 60601-1. The classification of the QDR Series X-ray Bone Densitometers under this specification is Class 1, Type B.

The QDR Series comply with IEC 60601-1-3 except for Section 29.205.2 which specifies "...focal spot to skin distances (FSSDs) of 45 cm or more in normal use." The QDR Series FSSD, which is approximately 42.5 cm, has been selected to provide optimum spatial resolution and precision with minimum patient exposure.

The QDR Series comply with the following IEC standards:

IEC 60601-1 2014	IEC 60601-2-28 2010
IEC 60601-1-2 2014	IEC 60825-1 2014
IEC 60601-1-3 2013	IEC 60601-1-6 2013

Software used in the QDR Series was developed using IEC 62304 as a guide.

1.4 Warnings and Cautions



Warning:

To avoid risk of electric shock, this equipment must only be connected to a supply mains with protective earth. No modification of this equipment is allowed.

1.4.1 EMI

This instrument is designed to be compatible with the electromagnetic environments specified in IEC60601-1-2 and will operate satisfactorily when placed in an environment that includes other equipment complying with that standard.

1.4.2 Accessories

Do not use any accessories in conjunction with this instrument other than those supplied by Hologic for use with the instrument.

1.4.3 Caution

Caution

To protect the integrity of the system and the safety of the patient and operator, do not power the computer and accessories plugged into the outlet strip at the rear of the console from another source. Also, do not power any accessories or appliances other than those provided with the system from the outlet strip at the rear of the console.

1.5 Labels





Horizon System Main Label

The Main Label includes:

- QDR X-ray Bone Densitometer
- Horizon Model
- Manufacturer name and address
- Complies with FDA Radiation Performance Standards 21 CFR Sub-chapter J applicable on date of manufacture.
- Date of manufacture
- Serial Number
- IEC standards

Laser Exit Aperture Label

The Laser Exit Aperture Label includes:

- Avoid Exposure
- Laser Radiation emitted from this Aperture

INMETRO & ULBR Product Label

The INMETRO & ULBR Product Label includes:

- includes the INMETRO & ULBR symbols
- Segurança

7 P(AC OW	; Z er	Ŷ					
Line Voltage (V)	100	120	230					
Frequency (Hz)	50/60	50/60	50/60					
Maximum Current (A)	16	14	8					
Maximum Apparent Resistance (ohm)	0.32	0.32	1.28					
		280-037	9 Rev 004					
				60 kV 2				
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AC Power Label

The **AC Power Label** lists system power specifications including:

- Line Voltage (V)
- Frequency (Hz)
- Maximum Current (A)
- Maximum Apparent Resistance (ohm)

Tank Label

The Tank Label includes:

- Manufacturer name and address
- Model numbers
- Serial Numbers
- Nominal tube rating
- Focal spot
- Filtration Aluminum equivalence

HF X-ray Source Label

The HF X-ray Source Label includes:

- 21 CFR Compliance statement
- Date of manufacture
- Model number of the source assembly
- Type: Assembly, source
- Serial Number of the source assembly
- IEC standards

Drum Filtration Label

The Drum Filtration Label includes:

- 21 CFR Compliance statement
- Manufacturer name and address
- Model assembly number
- Serial number
- EC and IEC numbers



Computer Certification Label

The **Computer Certification Label** includes:

- 21 CFR Compliance statement
- Manufacturer name and address
- Computer assembly number
- Computer assembly serial number
- Manufacture date

System Computer Label

The System Computer Label includes:

- System Computer assembly number
- Manufacture date
- Manufacturer name and address
- Electrical rating

X-ray Warning Label

• Warning: This x-ray unit may be dangerous to patient and operator unless safe exposure factors, operating instructions and maintenance schedules are observed.

1.6 Symbols

	Table	1: Symbols	
	CSA listed device	C € 2797	CE mark
4	Dangerous voltage	\triangle	Caution
8	X-ray source assembly		X-ray source emitting
Â	Warning: Electricity	\sim	Alternating current
×	Type B applied part		Protective earth (ground)
	Date of Manufacture		Manufacturer
X	Equipment to be disposed in compliance with European Directive 2002/96/EC on Waste Electrical and Electronic Equipment.	EC REP	Authorized representatives in the European Community
REF	Catalog number	SN	Serial number
<u>}}} *</u>	Radiation filter	Segurança	INMETRO & ULBR Mark
	Follow instructions for use	PINCH POINT	Danger: Pinch Point
	Temperature Limit	<u>%</u>	Humidity Limit

1.7 Support Materials

1.7.1 QDR Reference Manual

Reference for system technology information.

1.7.2 Online Help

Click Help in the main window or on most dialogs, or press F1.

1.7.3 QDR Series Technical Specifications Manual

Reference for specification information.

1.7.4 QDR Cyber-Security Information

Log on for support information. For QDR cybe<u>r</u>-security information access: <u>https://www.hologic.com/package-inserts/breast-skeletal-health-products/horizon-dxa-system-package-insertsifus</u>.



1.8 Main Window

Figure 1: Horizon Main Window

Main Window Features

- 1. Menu Bar
- 2. Main Work Area
- 3. Patient and Scan Records
- 4. System Functions
- 5. System Messages
- 6. Daily Functions
- 7. Help

Chapter 2 System Startup and Shutdown

2.1 System Startup

1. Verify the Control Panel up and locked in horizontal position (on Horizon A Models only).

Note

On Horizon A models, when the table is in the Patient On/Off position, the Control Panel swings down vertically to facilitate patient transfer from a stretcher to the Horizon table.

When the Control Panel is in the vertical down position, the system automatically shuts down table communication with the application as a normal safety function. When the Control Panel is returned to the horizontal position, after a three-second delay, table communication is restored for normal operation.

- 2. On the **Control Panel**, verify the **E-stop** button is up.
- 3. Turn the computer on.

The monitor and printer should already be on during a normal system startup.

4. Log on to the **QDR**.

2.2 System Shutdown

- 1. Select **Exit** in the main window.
- 2. Select Exit QDR with shutdown?
- 3. Select OK.

Caution

Do not turn off the circuit breaker.

Do not turn off the monitor or printer.

Chapter 3 Quality Control Procedure

- 1. Select **Daily QC** in the main window.
- 2. Place the spine phantom on the table with the registration mark (**A**) to the left, foot end.
- 3. Position the phantom parallel to the back of the table.
- 4. Align the laser cross hair (**B**) with the registration mark.
- 5. Select **Continue**.



Figure 2: Spine Phantom Position

3.1 System Test

If the system test fails, follow the instructions on screen to resolve the problem, and repeat QC.

3.2 Auto QC

When Auto QC passes, click OK to start scanning patients.

If Auto QC fails, follow the instructions to resolve the problem.

3.3 Automatic Body Composition Calibration

QDR systems with **APEX or QDR for Windows XP Version 12.4.2 and higher** (except upgraded systems), incorporate an automatic calibration for Body Composition. The system monitors when the calibration was last performed and, if a week has elapsed, automatically performs the calibration when QC is run. This process adds only a few seconds to the QC procedure.

When the calibration is complete, the system prompts you to remove the QC phantom.

- 1. Click **OK** to perform the Radiographic Uniformity test for Adult WB. If Infant WB is installed, it will run immediately after the Adult WB test.
- 2. When this test is complete, click **OK** to return to the main screen.

Chapter 4 Patient Records

Note

Note

If the patient is a woman under the T-score reporting age, and postmenopausal, the menopause age must be entered in the biography or a T-score will not be generated.



The T-score reporting age is configurable (age 50 is the default).

4.1 Retrieving a Patient Record

- 1. Select **Patients** in the main window.
- 2. Select the **patient's name** to select.



To use search criteria, click the heading and type the search criterion into the text box.

4.2 Creating a Patient Record

- 1. Select **Patients** in the main window.
- 2. Select New Patient.
- 3. Select the **Biography** tab.
 - a. Enter the patient information.
 - b. Select OK.
- 4. Select the **Insurance** tab.
 - a. Enter the patient information.
 - b. Select OK.

4.3 Editing a Patient Record

- 1. Select **Patients** in the main window.
- 2. Select Edit Patient.
- 3. Select the **Biography** tab and change information as needed.
- 4. Select the **Insurance** tab and change information as needed.
- 5. Select **OK**.

4.4 Using Worklist to Retrieve a Patient Record



Worklist is an option on QDR systems

- 1. Select **Perform Exam** in the main window.
- 2. Select the **Worklist** tab.
- 3. Select Query.
- 4. Highlight the desired patient's name.
- 5. Select OK.

Note

- 6. Confirm the patient information
 - a. Select **OK** or
 - b. Select Edit Patient to edit the patient record, as described above.

4.5 Confirming Patient Information

The Patient Confirmation window is used to confirm, or enter, exam related information, FRAX[®] limiting criteria and FRAX risk factors.

4.5.1 Biographical Information

Ensure the patient name, ID, birth date, current weight and current height are correct.

4.5.2 Exam Information

Confirm or edit the exam information, as necessary.

- 1. Enter or change patient information.
- 2. Answer Questionnaire data.
- 3. Complete all assessment data.

Exam information includes:

- operator
- accession number that uniquely identifies a patient visit,
- referring physician
- user defined fields

Chapter 5 Performing an Exam

5.1 Patient Interview

The following is a list of questions to ask the patient (some may not apply).

Is there any chance of pregnancy?

If a female patient is (or may be) pregnant, postpone the scan until pregnancy is ruled out.

Has the patient had any radiological procedure using the following contrast agents within the last 7 days:

- Iodine
- Barium

Radiological contrast agents used for X-ray and CT can interfere with DXA scans. In particular, oral contrasts can remain in the gastrointestinal tract for several days affecting DXA results. Intravenous iodine normally clears within 72 hours for those patients with normal kidney function.

Hologic DXA measurements have been shown in several studies to be unaffected by nuclear isotope studies, so DXA measurements can be done immediately after nuclear isotope studies as long as the studies do not also include radiological contrast agents (such as iodine and barium).

Is the patient wearing any objects in the scan area such as an ostomy device, metal buttons or snaps, or jewelry?

This may interfere in the scanning of the patient.

Has the patient had any surgery in the area being scanned?

If so, consider whether to perform the examination. For example, any of the following internal objects could interfere with the scan:

- Pacemaker leads
- Radioactive seeds
- Metal implants
- Surgical staples
- Foreign bodies; e.g., shrapnel
- Radio-opaque catheters or tubes

If the patient had surgery on a hip or forearm, then the uninjured hip or forearm should be scanned.

5.2 Patient Preparation

To prepare the patient for the examination:

- Ensure that there is no metal (e.g., zipper, snap, belt, etc.) in the scan field. If necessary, have the patient change into a gown for the examination.
- For AP lumbar spine, hip, or whole body examinations, instruct the patient to remove their shoes.
- The subject weight limit is 227 kg (500 lbs.). For patients over this limit, scan the forearm.

5.3 Patient Selection

- 1. Select **Perform Exam** in the main window.
- 2. Create or retrieve a patient record.
- 3. Select OK.
- 4. Confirm the patient information.
- 5. Select OK.

5.4 Choosing the Scan Type

- 1. From the **Select Scan Type** window, select the type of exam to be performed.
- 2. Select Next >>.

5.5 Performing a Scan

For instructions on how to perform a specific scan type refer to the appropriate section of this manual.

Chapter 6 Manual Scan Analysis

To analyze a scan use the analysis step buttons, toolbox and brightness/contrast control found at the left side of the analysis window.

6.1 Analysis Step Buttons

Each of the analysis step buttons is used to perform one step of the analysis. To analyze a scan, click each button in order, starting at the top button, and perform the operation required until all steps are completed.

Button	Function
Global ROI (Region of Interest)	The defined boundaries of the image that is being analyzed. The ROI appears on the image as a box.
Bone Map	An illustration, created by the system, of the area of bone defined by the ROI. The map is superimposed, in yellow, on the image.
Vertebral Lines (Spine)	Used to mark the intervertebral spaces within the spine region of interest.
Neck (Hip)	Allows positioning of the hip femoral neck box. The neck box should not include the ischium.
MID/UD (Forearm)	Allows positioning of the middle (MID) and ultra- distal (UD) regions of the forearm.
Regions (Whole Body)	Used to define regions of interest on a whole body scan.
A/G Regions (Whole Body)	Used to delineate the Android and Gynoid regions on the Whole Body image. For Body Composition results the A/G Regions are used to calculate fat and lean (includes BMC) content and percent fat of the Android and Gynoid regions of the body.
VAT Regions	On APEX 4.0 and above the A/G Regions analysis step button also displays VAT Regions. VAT Regions are used to delineate the Visceral Adipose Tissue (VAT) regions on the Whole Body image. The VAT regions include the skin on the edge of the abdominal region and the visceral cavity.

Table 2: Analysis Step Buttons

Button		Function
Sub Regi	ions (Whole Body)	Used to delineate one or more areas within the Whole Body scan. There can be up to seven SubRegions and they can have irregular shapes and overlap. For Body Composition results, the fat and lean (includes BMC) content, and percent fat, of each SubRegion is calculated along with the net average (NETAVE) for all SubRegions.
\mathbf{k}	Note If SubRegions overlap the individual regions.	nen the net average will be the mathematical union of
Sub Regi Body)	ions Results (Whole	Displays results of the SubRegion analysis on the analysis window. For Body Composition results, click BCA.
Vertebra Spine BN	l Boundaries (Lateral MD)	Used to identify the anterior boundary of the vertebral bodies, as a dashed yellow line, on a lateral spine image.
Vertebra BMD)	l Bodies (Lateral Spine	Used to identify the boundaries of the vertebral bodies, as boxes, on a lateral spine image.
Mid Reg BMD)	ions (Lateral Spine	Used to adjust the regions in the middle of the vertebral bodies on a lateral spine image (rarely necessary).
Results		Displays results of the analysis on the analysis window.

Table 2: Analysis Step Buttons
6.2 Toolboxes

Toolboxes provide the tools used at each step of the analysis. The tools that are available depend upon scan type being analyzed and the step button in use.

Table 3: Global ROI Toolbox		
Tool	Function	
Whole Mode	Allows the entire ROI box to be moved over the image. The box is shown as dashed yellow lines.	
Line Mode	Allows one line on the ROI box to be moved. Click any line to select it. The active line is shown as yellow dashes.	
Point Mode	Allows one point on the ROI box to be moved. Points are shown on the box as plus signs. The active point is yellow.	
1/3 Distal (Forearm only)	Used to adjust the size of the 1/3 distal region of the forearm (rarely necessary).	

6.2.1 Bone Map Toolbox

Table 4: Bone Map Toolbox		
Tool	Function	
Add Bone	Used to connect outside edges on an incomplete bone map and fill in the missing area (rarely necessary).	
Delete Bone	Used to erase an area on the bone map (rarely necessary).	
Undo	Negates the last action performed. Note Undo allows the operator to view the unprocessed bone map. Viewing the bone map before islands are sunk and holes in the bone are filled by the analysis algorithm can reveal how well the automatic bone-finding algorithm worked and can be used with problematic scans. The Undo feature is active and available for all non-whole body scan types in the Bone Map stage.	

6.2.2 Lines Toolbox (Spine)

Used to mark the intervertebral spaces within the spine region of interest.

6.2.3 Neck Toolbox (Hip)

Table 5: Neck Toolbox (Hip)	
Tool	Function
Neck Box	Allows the neck box to be moved and/or resized (rarely necessary). The box is shown as dashed yellow lines.
Other Regions	Expands the Neck Toolbox to include Midline, Ward's Triangle and Trochanter tools.
Midline	Allows the midline to be moved or pivoted (rarely necessary).
Ward's Triangle	Allows Ward's Triangle to be adjusted (rarely necessary).
Trochanter	Allows the Trochanter area to be adjusted (rarely necessary).
Auto Position	Allows the system to automatically locate a region.

6.2.4 Results Toolbox (Whole Body)

Table 6: Neck Toolbox (Hip)	
Tool	Function
BMD	Display BMD results in the analysis window.
BCA	Display BCA results in the analysis window.
Rulers	Allows measurement of the patient's anatomy by placing rulers on the scan image. Rulers are displayed in the analysis window, on the image, when Display is checked.

6.2.5 Sub Region Toolbox

Table 7: Sub Region Toolbox	
Tool	Function
One Region	Manipulate SubRegions individually.
All Regions	Manipulate all SubRegions together as a unit.
Undo	Negate the last action performed.

6.2.6 Toolbox Controls

	Table 8: Toolbox Controls		
Control	Function		
*	Cursor used to move a SubRegion(s).		
¢ ¢	Cursor used to rotate a SubRegion(s). The hand is open until rotation is started; during rotation the hand is shown with pinched thumb and first finger.		
	Used in lumbar spine analysis (Vertebral Lines) to select the line above the currently selected line.		
	Used in lumbar spine analysis (Vertebral Lines) to select the line below the currently selected line.		
	Increases the size of the cursor used to add or delete bone from a bone map.		
	Decreases the size of the cursor used to add or delete bone from a bone map.		
E	Used to create a SubRegion.		
E	Used to delete a SubRegion.		
	Used to select the next SubRegion in a sequence of two or more SubRegions.		
	Used to select the previous SubRegion in a sequence.		
	Used to flip a SubRegion(s) vertically.		
	Used to flip a SubRegion(s) horizontally.		
	Used to move counterclockwise to select the next line or point in a SubRegion.		
Q	Used to move clockwise to select the next line or point in a SubRegion.		

6.3 Brightness/Contrast Control



Used to adjust brightness and contrast of the image to obtain best definition of anatomical features. Does not affect calculations.

Chapter 7 AP Lumbar Spine Examination

Start the exam as described in <u>Using Worklist to Retrieve a Patient Record</u> on page 16. In the **Select Scan Type** window, select **AP Lumbar Spine**.

7.1 **Positioning the Patient**

- 1. On the Control Panel, press Patient On/Off.
- 2. Place the patients on their back with their head at the right end of the table.
- 3. On the Control Panel, press Center.
- 4. Position the patient as shown in the following figure, *AP Lumbar Spine Positioning* and in the figure, *Leg Positions in Positioning the C-arm*.



Figure 3: AP Lumbar Spine Positioning

7.2 Positioning the C-arm

- 1. On the Control Panel, press **Laser**.
- 2. Use the **Arm** and Table controls to place the laser's cross hair 2.5 to 5 cm (1 to 2 inches) below the iliac crest and centered at the patient's mid-line.
- 3. Instruct the patient to remain still and breathe normally.



Figure 4: Leg Positions

7.3 Starting the AP Lumbar Spine Scan

Warning:

If the Control Panel X-ray indicator fails to shut off within 10 seconds after the end of the scan, press the red Emergency Stop button immediately. Call your Hologic service representative before resuming operation.

- 1. Select Start Scan. The X-rays On indicator flashes until the scan stops.
- 2. Inspect the image as it is generated.
 - If the spine is positioned correctly, when you see the ribs attached to T12 on the image, *AP Lumbar Spine*, click **Stop Scan**.
 - If the spine is not positioned correctly, click **Reposition Scan** to stop the scan for repositioning.



Figure 5: AP Lumbar Spine

7.3.1 Repositioning the Scan (if necessary)

- 1. Select **Reposition Scan** before the scan is completed.
- 2. Position the cursor over the spine image as shown in the figure, *Spine Repositioning*.



Figure 6: Spine Repositioning

- 3. To reposition the spine, click and drag the image so that:
 - The center of the lumbar spine is aligned with the blue vertical positioning line **[1]**.
 - The iliac crest is at or below the blue horizontal positioning line [2].



Note

You can also use the scroll bars to reposition the image.

- 4. When the spine is positioned correctly, click **Restart Scan**.
- 5. Select Start Scan.
- 6. When you see the ribs attached to T12 on the image, click Stop Scan.

7.4 Analyzing the Scan

- 1. Select Analyze Scan.
- 2. If there is a prior scan, click **Results**.

If there is no prior scan, click **Next** >>. A histogram appears to the right of the image in a window during all the analysis steps. It is intended to aide in the placement of intervertebral lines.

Select the >> button at the upper right, to expand and the << button to contract the window.

You can configure the histogram from the DXA*pro* configuration screen. Refer to Configuring the System in the *MAN-03648 Horizon Reference Manual*.

7.4.1 One-Time™ Auto Analysis

When auto-analysis is complete, results are displayed.



If Auto-Analysis is unsatisfactory, perform a manual analysis for correct analysis.



Figure 7: AP Lumbar Spine Analysis

7.5 Exiting Analysis

- 1. Select Close.
- 2. Select Report.

7.6 Generating and Printing Reports

See <u>*Reports*</u> on page 113.

Chapter 8 Hip Examination

- 1. Start the exam as described in <u>Using Worklist to Retrieve a Patient Record</u> on page 16.
- 2. In the Select Scan Type window, select Left Hip, Right Hip or Dual-Hip.

8.1 Positioning the Patient for Left, Right and Dual Hip Examinations

1. Position the patient as shown.

Use the foot positioner supplied by Hologic to maintain the correct femur position.



Figure 8: Left Hip Positioning without Hip Autopositioning

- 2. On the Control Panel press **Center**.
- 3. Place the foot positioner under the patient's legs
- 4. Align its center with the table and the patient's midline.
- 5. Rotate the entire leg (on the side to be scanned) 25° inward.
- Place the medial edge of the foot against the positioner.
 The foot should be flexed toward the ceiling as shown in the figures for *Left Hip Positioning without Hip Autopositioning* and *Right Hip Positioning without Hip Autopositioning*.
- 7. Adjust the strap to hold the foot in the correct position.



8. For Dual Hip scans rotate both legs 25° inward and adjust both straps to hold feet in the correct position.

Figure 9: Right Hip Positioning without Hip Autopositioning



Figure 10: Foot Positioning (Dual Hip)

8.2 Positioning the C-arm



Note

On hip scans of overweight or obese subjects: Abdominal fat overlying the hip region should be held back or otherwise retracted from the X-ray field of view because overlying fat (panniculus) may interfere with hip BMD results.

8.2.1 Tips to Identify the Greater Trochanter

- 1. Place your thumb on the iliac crest.
- 2. Spread your fingers.
- 3. Direct your little finger toward the knee.

The greater trochanter is located under your little finger.

If you are not able to feel the trochanter:

- 4. Have the patient bend the leg at the knee and lift.
- 5. Locate the crease formed at the top of the leg.

Use this as an approximate location of the greater trochanter.

8.2.2 Left Hip or Right Hip without Autopositioning for Hip Scans

- 1. On the **Control Panel** press **Laser**.
- 2. Use the **Arm** and **Table** controls on the Control Panel to place the laser's cross hair:
 - 7.6 cm (3 inches) below the greater trochanter
 - 2.5 cm (1 inch) medial to the shaft of the femur.

Refer to:

Figures, *Left Hip Positioning without Hip Autopositioning* or *Right Hip Positioning without Hip Autopositioning*, in *Positioning the Patient for Left, Right and Dual Hip Examinations* on page 31 and *Tips to Identify the Greater Trochanter* on page 33.

3. Instruct the patient to remain still and breathe normally.

8.2.3 Left Hip or Right Hip with Autopositioning for Hip Scans

- 1. Enable **Autopositioning** for Hip Scans.
- 2. From the **APEX Main Menu** select:
 - Utilities
 - System Configuration
 - Check Autopositioning for Hip Scans.
- 3. On the **Control Panel** press **Laser**.
- 4. Use the **Arm** and **Table** controls on the **Control Panel** to place the laser's cross hair at the outer edge of the greater trochanter (See the figure *Left Hip Positioning without Hip Autopositioning*). See. *Tips to Identify the Greater Trochanter* on page 33.



5. Instruct the patient to remain still and breathe normally.

Figure 11: Left Hip Positioning with Hip Autopositioning

8.2.4 Dual-Hip

- 1. Turn on the laser and position the C-arm over the patient's midline.
- 2. Select Continue.
- Palpate the patient's left leg to identify the greater trochanter. See <u>*Tips to Identify the Greater Trochanter*</u> on page 33.



For Dual-Hip, the left hip is always scanned first.

4. Use the **Arm** and **Table** controls to place the laser's cross hair 7.6 cm (3 inches) below the greater trochanter and 2.5 cm (1 inch) medial to the shaft of the femur.



Note

Note

If using Autopositioning, place the laser's cross hair at the outer edge of the greater trochanter.

5. Select Continue.

6. Instruct the patient to remain still and breathe normally.

8.3 Starting the Hip Scan

Warning:

If the Control Panel X-ray indicator fails to shut off within 10 seconds after the end of the scan, press the red Emergency Stop button immediately. Call your Hologic service representative before resuming operation.

- 1. Select Start Scan. The X-rays On indicator flashes until the scan stops.
 - a. Verify that the hip scan is acceptable as the image generates.
 - b. If the hip is not positioned correctly, select **Reposition Scan** to stop the scan for repositioning.

To reposition the image to include the entire femoral head (*<u>Repositioning the scan</u>* on page 35). If the shaft is not parallel, see <u>*Repositioning the patient*</u> on page 36.

8.3.1 Repositioning the scan

- 1. Select **Reposition Scan** before the scan is completed.
- 2. Position the cursor over the hip image.

Note

You can also use the scroll bars to reposition the image



Figure 12: Hip Repositioning

- 3. To position the hip, click and drag the image to the intersection of the two blue positioning lines, so the:
 - vertical positioning line [A] touches the outer edge of the Greater Trochanter [B]
 - horizontal positioning line **[C]** is centered on the Greater Trochanter **[B]**
- 4. Select Restart Scan.
- 5. Select Start Scan.

8.3.2 Repositioning the patient

- 1. Select **Reposition Scan** before the scan is completed.
- 2. Select Restart Scan.
- 3. Select Start Scan.
- 4. Evaluate the image as it displays. If the image is satisfactory, allow the scan to complete.

8.3.3 Additional Steps for the Dual-Hip Scan

In a Dual-Hip scan, the scanner moves to the right hip after finishing the scan of the left hip. If **SE Femur** is enabled, the system will prompt you to perform a SE Femur scan of the left femur prior to moving to the right hip. It will also prompt you to perform a SE Femur of the right femur after finishing the scan of the right hip. See <u>Positioning for the SE</u> Femur Scan after a Hip Scan on page 39 and <u>Starting the SE Femur Scan</u> on page 39.

- 1. Turn the laser on.
- 2. Check that the laser is positioned properly over the patient's right hip.
- 3. Select Continue.
- 4. Verify that the right hip scan is acceptable.
- 5. If it is not, reposition the image or patient and rescan.

8.4 Analyzing the Scan

- 1. Select Analyze Scan.
- 2. If there is a prior scan, click **Results**.
- 3. If there is no prior scan, click **Next** >>.

8.4.1 One-Time Auto Analysis

Note

When auto-analysis is complete, results are displayed.



If Auto-Analysis is unsatisfactory, perform a manual analysis. For correct analysis, see the following figure.



Figure 13: Properly Analyzed Hip Scan

8.5 SE (Single Energy) Femur Examination

Start the exam as described in <u>Using Worklist to Retrieve a Patient Record</u> on page 16. SE Femur scans can be selected directly from the **Select scan type** window or they can be selected at the end of a BMD hip scan. Selecting a SE Femur scan at the end of a hip scan is only possible if SE Femur scans are configured via a check-box on the **System Configuration** screen.



Figure 14: Left SE Femur Positioning

8.5.1 Positioning for the SE Femur scan

1. Position the patient as shown in the figure *Left SE Femur Positioning* in <u>SE (Single Energy) Femur Examination</u> on page 38.

Use the foot positioner supplied by Hologic to maintain the correct femur position.

- 2. On the **Control Panel** press **Center**.
- 3. Place the foot positioner under the patient's legs, and
- 4. Align it center with the table and the patient's midline.
- 5. Rotate the entire leg (on the side to be scanned) 25° inward and place the medial edge of the foot against the positioner.

The foot should be flexed toward the ceiling.

6. Adjust the strap to hold the foot in the correct position.

- 7. In the **Select Scan Type** window choose SE Femur.
- 8. Palpate the patient's leg to identify the greater trochanter. See <u>*Tips to Identify the Greater Trochanter*</u> on page 33.
- 9. On the Control Panel, press Laser.
- 10. Use the Arm and Table controls on the Control Panel to place the laser's cross hair:
 - Level of the greater trochanter
 - Middle of the shaft of the femur or slightly off set outward toward the outer edge of the femur to accommodate the entire length of the femur.
- 11. Instruct the patient to remain still and breathe normally.

8.5.2 Positioning for the SE Femur Scan after a Hip Scan

- 1. After performing a single hip scan, you may choose to perform an SE Femur scan.
- 2. Select the button labeled **SE Femur Scan**.

The **Positioning Femur Scan** Screen appears which shows you where the system will automatically position the C-arm. The screen provides the operator with the ability to reposition the scan image before the femur scan.

- 3. Verify that the image will start at the level of the greater trochanter and that the shaft of the femur is centered or slightly off set toward the inner edge of the femur to accommodate the entire length of the femur.
- 4. Select Next.

Warning:

8.5.3 Starting the SE Femur Scan

If the Control Panel X-ray indicator fails to shut off within 10 seconds after the end of the scan, press the red Emergency Stop button immediately. Call your Hologic service representative before resuming operation.

- 1. Select Start Scan. The X-rays On indicator flashes until the scan stops.
- 2. Verify that the SE femur scan is acceptable as the image generates. If the image is satisfactory, allow the scan to complete.
- 3. If the SE Femur scan is not positioned correctly, click **Reposition Scan** to stop the scan for repositioning.

8.5.4 Analysis of SE Femur Scan

The image appears in the viewer window.

The Viewer allows the operator to view the SE Femur image. The controls on the Viewer allow the operator to:

- change viewing modes
- zoom
- adjust the brightness and contrast of the image
- add annotations
- add comments
- place Rulers.

8.5.5 Visual Assessment

- 1. Visually inspect the image for deformity¹²³ particularly along the lateral cortex from the lesser trochanter to the supracondyal flare.
- 2. Look for focal reaction or thickening along the lateral cortex, which may be accompanied by a transverse radiolucent line.
- 3. Use the visual tools to increase magnification and adjust the contrast.

The changes in the lateral cortex may be subtle.



Note

The images should be read by a qualified medical professional

8.5.6 Image Toolbox

Table 9: Image Toolbox		
Tool	Function	
Multi View	When selected, places the viewer in Multiple View mode, displaying a second image in the right-hand panel.	
Visual Tools and Analysis Tools	will operate independently on either image. If both	
9 1	formed on the center image will be reflected on the	
image in the right panel.		
W-L Control	Select and drag the <i>ball</i> in the center of the triangle to adjust the contrast and brightness of the image.	
	For finer adjustment:	
	1. Right click on the image.	
	2. Select Adjust WL.	
	3. Select and drag the cursor on the image.	
Invert	Inverts the grayscale value of each pixel creating a negative of the image.	
Zoom	Is used to shrink or enlarge the scan image.	

8.5.7 Analysis Tools Control Tab

The Analysis Tools control tab provides tools to assess any deformities seen in the image. The tools in this section provide tools to:

- Control the method of adding Annotations and/or Rulers
- Control display of Annotations and Rulers on image
- Change Annotations

8.5.8 Assessment

Two radio buttons that determine which tool will be used when a user clicks in the image.

	Table 10: Assessment
Tool	Function
Annotations	When selected and you left-click on the image an annotation window will appear allowing for text to be entered and saved. Up to two annotations can be created.
Rulers	When selected and you left-click and drag on the image a ruler will be created. Rulers is a tool that allows the operator to measure cortical thickness and/or other features by placing rulers on the scan image. Up to six rulers may be added.

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Tool	Function	
Add Ruler	The cursor changes to a cross when placed over the scan image.	
	The cursor changes to a cross when placed over the scan image.	
	 Select and drag a line to place the ruler. The length of the new ruler (in centimeters) is displayed below the Rulers button. 	
	2. To add rulers repeat the above.	
Select Ruler	The cursor changes to a hand when placed directly on a ruler.	
	With the hand cursor displayed, click to select a ruler.	
Move Ruler	3. Select the ruler.	
	4. With the hand cursor displayed, click and drag the ruler to the desired location,	
	or	
	 Use the keyboard arrow keys to move the ruler up, down, left or right. 	

Tool	Function	
Select Endpoint of Ruler	The cursor changes to an arrow at the endpoint of a ruler. With the arrow cursor displayed, click to select the endpoint.	
Change the Size of Ruler	 With the arrow cursor displayed, 6. Select and drag the endpoint to the desired length and location, or 7. Use the keyboard arrow keys to move the endpoint up, down, left or right. 	
Delete a Ruler	 Select the ruler and click the keyboard Delete key, or, With the hand cursor displayed, right-click and select Delete. 	

Table 11: Adding and Modifying Rulers

Display Options

Two check boxes that specify what assessments are displayed on the image.

Tool	Function
Annotations	When selected, locations of any annotations that have been created on the image will be displayed. They are identified as 1 and 2.
Rulers	When selected, any rulers that have been created on the image will be displayed.

Table 12: Display Options

Change Annotations

Three buttons that provides the user with a means to add/change annotations. These buttons are enabled when an Annotation window is displayed on the image.

	Table 13: Change Annotations	
Tool	Function	
Focal Thickening	Selecting this button will display the annotation edit window and populate the edit field with the following text: "Focal reaction or thickening along the lateral cortex of the femoral shaft is present."	
Radiolucent Line	Selecting this button will display the annotation edit window and populate the edit field with the following text: "A transverse radiolucent line is present."	
Free Text	Selecting this button will display the annotation edit window where free text can be entered.	

Atypical Femoral Fracture Assessment

The text for all the annotations and lengths of all rulers placed on the image will be displayed in this section. There is also a comment section where free text can be entered as well as two buttons with predefined text that when clicked will populate in the Comments box.

Tool	Function
Correlation Advised	Selecting this button will insert the following text into the comment section: "Clinical correlation is advised as these features may be consistent with an incomplete atypical femur fracture."
Drug Complication	Selecting this button will insert the following text into the comment section: "Atypical femur fractures are a complication associated with long term use of antiresorptive therapy."

Table 14: Atypical Femoral Fracture Assessment

8.6 References

These references include further information about atypical femur fractures and the accompanying radiographic appearance and risk factors:

- Elizabeth Shane, David Burr, Peter R Ebeling, Bo Abrahamsen, Robert Adler, et al. "Atypical Subtrochanteric and Diaphyseal Femoral Fractures: Report of a Task Force of the American Society for Bone and Mineral Research." Journal of Bone and Mineral Research, Vol. 25, No. 11, November 2010
- 2. Elizabeth Shane (Co-Chair)*, David Burr, Bo Abrahamsen, Robert A. Adler, Thomas D. Brown, Angela M. Cheung, Felicia Cosman, Jeffrey R. Curtis, Richard Dell, David W. Dempster, Peter R. Ebeling, Thomas A. Einhorn, Harry K. Genant, Piet Geusens, Klaus Klaushofer, Joseph M. Lane, Fergus McKiernan, Ross McKinney, Alvin Ng, Jeri Nieves, Regis O'Keefe, Socrates Papapoulos, Tet Sen Howe, Marjolein C.H. van der Meulen, Robert S. Weinstein, Michael Whyte. Atypical subtrochanteric and diaphyseal femoral fractures: Second report of a task force of the American society for bone and mineral research J Bone Miner Res. DOI: 10.1002/jbmr.1998
- 3. Zehava Sadka Rosenberg, Renata La Rocca Vieira, Sarah S. Chan, James Babb, Yakup Akyol, et al. "Bisphosphonate-Related Complete Atypical Subtrochanteric Femoral Fractures: Diagnostic Utility of Radiography." AJR:197, October 2011
- FDA Drug Safety Communication: Safety update for osteoporosis drugs, bisphosphonates, and atypical fractures; <u>http://www.fda.gov/Drugs/DrugSafety/ucm229009.htm</u>

8.6.1 Dual-Hip Scans

- 1. After analysis of the right hip, click Close.
- 2. To analyze the left hip, click Analyze Another Scan.

8.7 Generating and Printing Reports

See <u>Reports</u> on page 113.

Chapter 9 Forearm Examination

- 1. Start the exam as described in <u>Using Worklist to Retrieve a Patient Record</u> on page 16.
- 2. In the Select Scan Type window, select Left Forearm or Right Forearm.

9.1 Measuring the Patient's Forearm

- 1. Measure the forearm up to the ulna styloid **[1]** in centimeters.
- 2. Record the measurement.



Figure 15: Measuring the Left Forearm

9.2 Positioning the Patient

- 1. On the Control Panel press Center.
- 2. Position the patient as shown in:
 - See *Laser Position on Left Forearm* for left forearm.
 - See Laser Position on Right Forearm for right forearm.
 - See the figure *Patient On/Off Position for Left Supine Forearm* in <u>*Positioning the*</u> <u>*Patient for Supine Forearm Examinations*</u> on page 50 and the following figure Laser *Position on Left Forearm* for left supine forearm.
 - The figure *Patient Position for Right Supine Forearm* in *Positioning the Patient for Supine Forearm Examinations* on page 50 for right supine forearm.



Figure 16: Left Forearm Positioning



Figure 17: Laser Position on Left Arm



Figure 18: Right Forearm Positioning



Figure 19: Laser Position on Right Arm

9.2.1 Positioning the Patient for Supine Forearm Examinations

- 1. Position the patient hip at the outer edge of the table.
- 2. For right forearm, place the patient's left arm across the chest. For left forearm, place the patient's right arm across the chest.
- 3. Bend the elbow so that the forearm is parallel to the long side of the table.
- 4. Align the long bones of the forearm with the long axis of the laser.
- 5. Ensure that no obstruction is within 1 cm (.5 inch) of the radius and ulna sides of the forearm.
- 1. Move C-Arm to far right
- 2. Extend table to outer limit



Figure 20: Patient On/Off Position for Left Supine Forearm

Allow no obstruction within 1 cm (.5 inch) on each side of the arm.



Figure 21: Patient Position for Left Supine Forearm



Figure 22: Patient Position for Right Supine Forearm

9.3 Positioning the C-arm

- 1. On the Control Panel, press Laser.
- 2. Use the Arm Controls on the Control Panel to position the starting point on the forearm:
 - Left Forearm Scan: Starts at the mid-forearm. Verify that the first row of carpal bones is within 15 cm (6 inches) of the starting point (see the figure, *Laser Position on Left Forearm* in *Positioning the Patient for Supine Forearm Examinations* on page 50).
 - **Right Forearm Scan**: Starts at the first row of carpal bones (see the figure, *Laser Position on Right Forearm* in *Positioning the Patient for Supine Forearm Examinations* on page 50).
- 3. Instruct the patient to remain still and breathe normally.

9.4 Starting the Forearm Scan



Warning:

If the Control Panel X-ray indicator fails to shut off within 10 seconds after the end of the scan, press the red Emergency Stop button immediately. Call your Hologic service representative before resuming operation.

- 1. Select **Start Scan**. The *X*-rays On indicator flashes until the scan stops.
- 2. Inspect the image.
 - The radius and ulna must appear straight and centered.
 - The image shows at least the first row of carpal bones near the bottom for a right forearm scan, or near the top for a left forearm scan.
 - The ulna side contains at least enough air to equal the ulna's shaft width.
- 3. If the image is not acceptable, click **Reposition Scan**.
- 4. If the image is acceptable, allow the scanning to complete.

9.4.1 Repositioning the Scan (if necessary)

- 1. Select **Reposition Scan** before the scan is completed.
- 2. Position the cursor over the forearm image.



Note

You can also use the scroll bars to reposition the image



Figure 23: Forearm Repositioning

- 3. To position the forearm:
 - Move the first row of carpal bones within the horizontal blue positioning line and the outer limit of the scan field.
 - The radius and the ulna should be parallel between the two blue vertical positioning lines.
- 4. When the forearm is positioned correctly, select **Restart Scan**.
- 5. Select Start Scan.

9.4.2 Repositioning the Patient (if necessary)

1. Adjust the forearm so that it is straight.

Note

The patient's forearm should be moved to include or exclude more of the carpal bones

- 2. Select Restart Scan.
- 3. Select Start Scan.
- 4. Evaluate the image as it displays. If the image is satisfactory, allow the scan to complete.

9.5 Analyzing the Scan

- 1. Select Analyze Scan.
- 2. Select **Next** >>.

Note

For descriptions of Analysis Step Buttons and Toolboxes, see <u>Analysis Step Buttons</u> on page 19.

9.5.1 Entering the Forearm Length

- 1. Select Length. The forearm length must be between 4.0 and 42.0 centimeters.
- 2. Enter the length in centimeters.

9.5.2 Defining the Global ROI

- 1. Select Global ROI.
- 2. Using the Whole Mode and Line Mode tools, adjust the ROI as shown.



Figure 24: Forearm ROI

Table	15:	Ulnar	Styloid	Process

Item	Description
1	Ulnar Styloid Process



Note Using Whole Mode, the top line of the Global ROI box should be placed at the tip of the Ulnar styloid.

Using Line Mode, move the ROI line on the radius side in until the blue dashed line is just touching the lateral edge of the radius.

Repeat this process on the ulna side, making sure approximately 1 - 2 cm of air is included in the Global ROI on the ulna side.

In subjects with large forearms, the dashed blue line may have to be moved outward from the lateral edge of the ulna to include the required 1 - 2 cm of air background points.

9.5.3 Viewing the Bone Map

- 1. Select Bone Map.
- 2. In most cases it will not be necessary to edit the bone map. If it is required, use the toolbox tools to edit the bone map as shown in the following figure.



Figure 25: Forearm Bone Map



Note

The Global ROI must contain adequate air points to ensure proper bone mapping and analysis results. In some subjects, the blue dashed line on the ulna side may need to be adjusted outward to include adequate air points

9.5.4 Checking the MID/UD Region

1. Select MID/UD.

In most cases it will not be necessary to adjust the MID/UD region or reposition the ulna/radius divider.

2. If it is required, use the toolbox tools to adjust as shown in the figure.



Figure 26: Forearm MID/UD

9.6 Viewing Results

Select Results.

9.7 Exiting Analysis

- 1. Select Close.
- 2. Select Report.

9.8 Generating and Printing Reports

See <u>Reports</u> on page 113.
Chapter 10 Whole Body Examination

Start the exam as described in <u>Using Worklist to Retrieve a Patient Record</u> on page 16. In the Select **Scan Type** window, select **Whole Body**.

10.1 Body Composition Analysis

Hologic's Body Composition Analysis provides the ability to analyze the soft tissue composition of the entire body and to follow the changes in soft tissue composition over time in response to interventions such as diet and exercise. Analysis of fat mass, lean mass, and % fat mass can be reported for the entire body and head, arms, trunk, pelvis, and legs.

QDR whole body systems provide body composition results:

- Fat
- Lean combined with bone mineral content (BMC)
- Lean, (not including bone mineral content)
- %Fat. The % Fat is the fat mass divided by the total of the fat mass plus lean mass plus BMC.

The Hologic Visceral Fat Software estimates the visceral adipose tissue (visceral fat) content in adult males and non-pregnant females. The results are reported as:

- Visceral Fat Area
- Visceral Fat Mass
- Visceral Fat Volume

Due to the sensitivity of the soft tissue analysis, *the patient should wear only a hospital cloth or paper gown for the scan*. To the extent possible, tuck any clothing under the patient. If a sheet is used to cover the patient, the entire patient from the chin down including all extremities, are covered. A pillow must not be in the scan, since the material will affect the soft tissue measurement.

10.2 Positioning the Patient

1. Position the patient as shown the figure *Whole Body Positioning*.

Note

Warning:

Instruct large patients to place their hands vertically at their sides with the fifth finger on the table pad. Their hands should be next to the thighs to ensure the hands and arms are within the table limits

During the whole body exam, ensure that all patient body parts are at, or above, the surface of the examination table to avoid pinch points



2. Instruct the patient to remain still and breathe normally.

Figure 27: Whole Body Positioning

10.3 Starting the Whole Body Scan



Warning:

If the Control Panel X-ray indicator fails to shut off within 10 seconds after the end of the scan, press the red Emergency Stop button immediately. Call your Hologic service representative before resuming operation.

- 1. Click **Start Scan**. The *X*-rays On indicator flashes until the scan stops.
- 2. Make sure the patient's arms are included in the scan on the C-arm's first and last pass. The patient should remain still until the scan is complete.

10.4 Whole Body Fan Beam Analysis

The default QDR system Whole Body Fan Beam Analysis method employs an Auto Whole Body feature that automatically adjusts the analysis for patients weighing from 17.6 lbs (8 Kg) to 88 lbs (40 Kg). Above 88 lbs the Auto Whole Body analysis provides results identical to previous whole body analysis versions. The software has not been evaluated for patients weighing less than 17.6 lbs and use on patients below this weight is not recommended.

Because Auto Whole Body automatically provides an improved analysis for patients less than 88 lbs (40 Kg), the Legacy PWB analysis method is no longer recommended in humans. Patients previously analyzed with Legacy PWB should be re-analyzed using Auto Whole Body because the reference database comparisons and Z scores will not be valid.



Note

If you are performing Whole Body Exams in subjects less than 40 kg, e.g. children, it is important to make sure that Auto Whole Body is not disabled in the System Configuration for Analyze. The default setting should be used, i.e., Auto Whole Body should be enabled

10.5 Analyzing the Scan



Note

Body Composition Analysis is performed at the same time as the Whole Body analysis. The Analysis window initially displays with Regions and Line Mode active.

1. Select Analyze Scan.



Note

For descriptions of Analysis Step Buttons and Toolboxes, see <u>Analysis Step Buttons</u> on page 19.

2. Select Results.

10.5.1 Body Composition Analysis

The same anatomical regions are used for calculating soft tissue values and Bone Mineral values. To correctly report the soft tissue values, adjust the anatomical cut lines to include the soft tissue appropriate for that region. For example, adjust the leg regions such that the thigh tissue is located within the appropriate leg region, not the arm regions.

10.5.2 Default Placement of Whole Body Regions

When default placement is complete, the region lines should be checked for accuracy and adjusted if necessary. Refer to the following figures, *Whole Body Analysis with Horizontal and Lower Pelvic Divider Lines* and *Whole Body Analysis (Vertical Lines)* for correct analysis.

To check the horizontal and lower pelvic divider lines, see the following figure *Whole Body Analysis with Horizontal and Lower Pelvic Divider Lines.*

- 1. Verify that the Neck line is just under the patient's jaw.
- 2. Verify that the T12-L1 line in the spine is at the approximate level of T12-L1.
- 3. Verify that the Upper Pelvic line is just above the iliac crest.
- 4. Verify that the Lower Pelvic divider lines separate the legs and trunk.

If adjustment is required, use the tools in the Regions Toolbox to click and drag the lines to their correct position.



Figure 28: Whole Body Analysis (Horizontal and Lower Pelvic Divider Lines)

To Check the Vertical Lines (see figure Whole Body Analysis with Vertical Lines):

- 1. Verify that the chest lines are close to the chest.
- 2. Verify that the spine lines are close to the spine.
- 3. Verify that the leg lines are close to the leg.
- 4. Verify that the leg divider line evenly separates the legs and feet.

If adjustment is required, use the tools in the Regions Toolbox to click and drag the lines to their correct position.



Figure 29: Whole Body Analysis (Vertical Lines)

To Fine Tune the Vertical Lines (if necessary)

If necessary, use the Point Mode controls in the Regions Toolbox to adjust (fine tune) the vertical lines as follows:

- 1. Drag the point on the left shoulder so that it is positioned between the head of the humerus and scapula at the glenoid fossa. Repeat this step for the right shoulder.
- 2. Drag the three points along the left side of the spine close to the spine, matching the curvature if possible. Repeat this step for the three points along the right side of the spine.
- 3. If necessary, drag the left point above the iliac crest out at the sides to include soft tissue of the chest and thighs. Repeat for the right point above the iliac crest.
- 4. Drag the lower point of the triangle below the pelvis to bisect both femoral necks.
- 5. Use the scroll bar to scroll the image to the bottom of the scan. Drag the left and right points near the feet to include as much of the soft tissue in the thighs as possible without including the patient's hand and fingers.

10.5.3 Adjusting A/G Regions (if necessary)

Select A/G Region.

Note



Adjustment of the A/G regions is usually not required. Only adjust the A/G regions if necessary

Refer to the figure *A*/*G Regions* and check the Android and Gynoid regions as follows:

Android ROI

- 1. The Android region height should equal 20% of the distance from the pelvic horizontal line to the neckline.
- 2. Ensure the lower boundary of the Android region coincides with the pelvic horizontal line.
- 3. Ensure the lateral boundaries of the Android region coincide with the arm lines.

Gynoid ROI

- 1. The Gynoid region height should equal twice the height of the Android region.
- 2. Ensure the upper boundary of the Gynoid region is below the pelvic horizontal line by 1.5 times the height of the Android region.
- 3. Ensure the lateral boundaries of the Gynoid region coincide with the arm lines.

If adjustment is required, use the tools in the Android/Gynoid SubRegion Toolbox to click and drag the lines to their correct position.



Figure 30: A/G Regions

10.5.4 Visceral Adipose Tissue

Visceral Adipose Tissue (VAT) is the fat inside the abdominal cavity, inside the abdominal muscle wall. Horizon VAT area results are calibrated to, and highly correlated with, VAT area results provided by a computed tomography slice at the L4-L5 level (see figure *VAT Regions Pictured as CT Slice*).

The VAT regions occupy a band crossing the patient's abdominal cavity between the pelvis and the rib cage. One region covers the entire width of this band, from one side of the patient's body to the other. The other region includes only the interior of the abdominal cavity, from the inner edge of the abdominal muscle wall on one side of the body to the inner edge of the abdominal muscle wall on the other side. See the figure in <u>Adjusting VAT Regions (if necessary)</u> on page 65.

VAT regions are available only in APEX 4.0, and higher, on Horizon A, W and Wi systems.



1. Subcutaneous Fat

- 2. Abdominal Organs
- 3. Abdominal Muscle Wall
- 4. VAT

Figure 31: VAT Regions Pictured as CT Slice

10.5.5 Adjusting VAT Regions (if necessary)



Manual adjustment is usually not required. Avoid minor adjustments

1. Select A/G Regions.

- 2. Refer to the figure *Abdominal Tissue Features* and adjust the image contrast and brightness so that you can see:
 - Abdominal muscles on either side of the visceral cavity.
 - Dark subcutaneous fat on the outer edge of the abdomen.



- 1. Visceral Cavity
- 2. Inner Abdominal Muscle Wall
- 3. Subcutaneous Fat

Figure 32: Abdominal Tissue Features

3. Ensure that the VAT regions do not include any pelvic bone. The pelvic cut line determines the vertical placement of the VAT regions. To adjust the pelvic cut line, drag to just above the iliac crest.

- 4. Ensure that the abdominal region extends from the outer skin line of one side of the body to the outer skin line of the other side. The larger rectangle defines the abdominal region. See the following figure, *VAT Regions*.
- 5. Ensure that the next set of lateral lines in the abdominal region is positioned at the edge of the laterally evident subcutaneous fat just outside the abdominal muscle wall. See the following figure, *VAT Regions*.
- 6. Ensure that the lateral lines that identify the visceral cavity are positioned at the inner edge of the abdominal muscle wall. See the following figure *VAT Regions*.



- 4. Abdominal Region
- 5. Edge of Subcutaneous Fat
- 6. Visceral Cavity

Figure 33: VAT Regions

You can select and move the VAT vertical lines. Whole and line modes are available for the VAT option. Point mode is unavailable for this option.

10.5.6 VAT References

References related to the clinical utility of visceral fat estimation include:

- Sam S, Haffner S, Davidson MH, D'Agostino Sr RB, Feinstein S, Kondos, et al. "Relationship of Abdominal Visceral and Subcutaneous Adipose Tissue With Lipoprotein Particle Number and Size in Type 2 Diabetes." *Diabetes*, Vol. 57, August 2008
- Pascot A, Lemieux I, Prud'homme D, Tremblay A, Nadeau A, Couillard C, et al. "Reduced HDL particle size as an additional feature of the atherogenic dyslipidemia of abdominal obesity." *Journal of Lipid Research*, Volume 42, 2001
- Fox C, Massaro JM, Hoffmann U, Pou KM, Maurovich-Horvat P, Liu C, et al. "Abdominal Visceral and Subcutaneous Adipose Tissue Compartments: Association With Metabolic Risk Factors in the Framingham Heart Study" *Circulation* 2007, 116:39-48

- Preis S, Massaro JM, Robins SJ, Hoffmann U, Vasan RS, Irlbeck T. "Abdominal Subcutaneous and Visceral Adipose Tissue and Insulin Resistance in the Framingham Heart Study." *Obesity* (Silver Spring). 2010 November; 18(11): 2191– 2198. doi:10.1038/oby.2010.59
- Nieves DJ, Cnop M, Retzlaff B, Walden CE, Brunzell JD, Knopp RH, Kahn SE. "The Atherogenic Lipoprotein Profile Associated With Obesity and Insulin Resistance Is Largely Attributable to Intra-Abdominal Fat." *Diabetes*, VOL. 52, January 2003
- Goodpaster BH, Krishnaswami S, Harris TB, Katsiaras A, Kritchevsky SB, Simonsick EM, et al. "Obesity, Regional Body Fat Distribution, and the Metabolic Syndrome in Older Men and Women." *ARCH INTERN MED*, Vol 165, Apr 11, 2005. <u>WWW.ARCHINTERNMED.COM</u>

10.6 Viewing Results

Select Results and BMD to obtain Bone Mineral Density results.

Select **BCA** to obtain Body Composition results that include the regions plus the android and gynoid SubRegions.

10.6.1 Rulers

Rulers are placed on the Whole Body scan image to measure the patient's anatomy. Up to six rulers may be added.



Note

Rulers require a special scan image not included in some earlier versions of APEX. Rulers are not available for those scans.

To add a ruler:

- 1. Ensure that BMD is selected in the Results Toolbox and click **Rulers**. *The cursor changes to a cross when placed over the scan image.*
- 2. Place the cross on the image where the ruler will start and click and drag a line to place the ruler.

To select a ruler:

The cursor changes to a hand when placed directly on a ruler. With the hand cursor displayed, click to select a ruler.

To move a ruler:

Select the ruler. With the hand cursor displayed, click and drag the ruler to the desired location, or use the keyboard arrow keys to move the ruler up, down, left or right.

To select the endpoint of a ruler:

The cursor changes to an arrow at the endpoint of a ruler. With the arrow cursor displayed, click to select the endpoint.

To change the size of a ruler:

With the arrow cursor displayed, click and drag the endpoint to the desired length and location, or use the keyboard arrow keys to move the endpoint up, down, left or right.

To delete a ruler:

Select the ruler and click the keyboard Delete key, or, with the hand cursor displayed, right click and click **Delete**.

To zoom the image:

With Rulers selected, select the **Sun/Moon** button and select the desired zoom factor using the Zoom control. The image can be sized to 100%, 144%, 200%, 288% or 400%. Select the **Sun/Moon** button again to return to ruler placement.

Display Checkbox

With Rulers selected, this checkbox will be checked by default. With Rulers unselected, check to show rulers on the image. If unchecked, rulers do not appear on the image (rulers are not deleted, but not shown).

10.7 Exiting Analysis

If your analysis is complete, click **Close** to print a report or to analyze another scan. To create user-defined SubRegions, click **Sub Regions** and continue to <u>User-Defined</u>. <u>SubRegions</u> on page 68.

10.8 User-Defined SubRegions

SubRegion analysis mode is a research tool developed to allow investigators to measure several user-defined regions of bone and soft tissue. Any area within the Whole Body scan can be analyzed. There can be up to seven SubRegions and they can overlap and have irregular shapes. If regions overlap then the net average will be the mathematical union of the individual regions.



Note

For descriptions of Analysis Step Buttons and Toolboxes, see <u>Analysis Step Buttons</u> on page 19.

10.9 Resolving Asymmetric Results with Reflection

The asymmetric results warning message is displayed if a significant difference in mass is detected between the right and left arm (25%), or the right and left leg (15%).

Warnin Asymmetric Results	
A difference in mass between the has been detected. To copy the actions:	
Copy results from right arm to left arm	
Copy results from right leg to left leg	
Сору	Do Not Copy

Figure 34: Asymmetric Results Warning

The possible causes of detection and resolutions include:

- The whole body analysis lines that separate the arms and legs are asymmetrically placed. Examine the lines used in the analysis and click Regions to adjust the arm and leg lines so they are symmetrical.
- Part of an arm or leg (usually at the hip) was outside of the scan field. In the warning screen you can select to copy the complete arm or leg to the arm or leg with missing information.
- The patient is asymmetrical (e.g. amputation, polio, etc.). In the case of a significantly asymmetrical patient, consult the physician to determine how results should be reported. Not copying provides the most accurate measurement of the patient, but copying may allow more accurate comparison to reference data for whole body measurements.

The system automatically determines which limbs are smaller. In the warning message dialog box you can select to copy the larger limb results to the smaller limb by checking one or both of the message check boxes.

In the warning message dialog box example, the mass of the right arm is significantly larger than the mass of the left, but the difference in the mass of the right and left leg was not large enough for the system to suggest copying. However, this box can be checked if you also want the system to copy the leg results.

Select **Copy** to perform the action indicated by the check boxes. Select **Do Not Copy** to close without changing any results. Results that are copied from one arm or leg to the other are indicated on the report. Accuracy and precision may be affected by copying from one side to the other. However, depending on body habitus of the subject, copying may give the most accurate results.

10.10 Enable NHANES BCA

To enable NHANES BCA go to the APEX Main Menu, select **Utilities**, **System Configuration**, **Analyze** tab and select the **Enable NHANES BCA** check box. Select this option to apply the calibration recommended by Schoeller *et al.*¹ When enabled, NHANES BCA will be noted in the BCA results section.

¹ Schoeller DA, Tylavsky FA, Baer DJ, Chumlea WC, Earthman CP, Fuerst T, Harris TB, Heymsfield SB, Horlick M, Lohman TG, Lukaski HC, Shepherd J, Siervogel RM, Borrud LG "QDR 4500A dual-energy X-ray absorptiometer underestimates fat mass in comparison with criterion methods in adults." Am J Clin Nutr. 2005;81(5):1018-25.

10.11 Generating and Printing Reports

See <u>Reports</u> on page 113.

Chapter 11 Supine AP/Lateral Spine BMD Examination (Horizon A)

11.1 Table Safety Feature

C-arm motion stops if you or the patient touch the safety strips along either edge of the table. If this happens, press **Enable Lateral** to complete C-arm rotation.

11.2 Positioning for the AP/Lateral Scan

Start the exam as described in <u>Using Worklist to Retrieve a Patient Record</u> on page 16. In the Select **Scan Type** window, select **AP/Lateral**.

- 1. On the Control Panel, press Patient On/Off.
- 2. Place the patient on their back with their head at the right end of the table.
- 3. On the Control Panel, press **Center**.
- 4. Position the patient as shown in the figure AP/Lateral Spine Positioning.

Note

The patient must not move between the AP and lateral scans.



Figure 35: AP/Lateral Spine Positioning

11.3 Starting the AP Scan

- 1. On the Control Panel, press **Enable Lateral**.
- 2. Select Continue.
- 3. Refer to *<u>Starting the AP Lumbar Spine Scan</u>* on page 27 and complete the scan.

11.4 Analyzing the AP Scan

- 1. Select **Next** >>.
- 2. Select Close.
- 3. Continuously press **Enable Lateral** on the Control Panel until the C-arm has rotated fully to the lateral scan position.

11.5 Starting the Lateral Scan

Warning:

If the Control Panel X-ray indicator fails to shut off within 10 seconds after the end of the scan, press the red Emergency Stop button immediately. Call your Hologic service representative before resuming operation.

- 1. Select Start Scan. The X-rays On indicator flashes until the scan stops.
- 2. When L2 to L4 is fully displayed (see the figure *Lateral Spine*), select **Stop Scan**.



Figure 36: Lateral Spine

3. Continuously press **Enable Lateral** on the Control Panel until the C-arm has rotated to its original position.

11.6 Analyzing the Lateral Scan

- 1. Select Analyze Scan.
- 2. Select **Next** >>.

Note

For descriptions of Analysis Step Buttons and Toolboxes, see <u>Analysis Step Buttons</u> on page 19.

11.6.1 Defining the Global ROI

- 1. Select Global ROI.
- 2. Using the **Whole Mode** and **Line Mode** tools, adjust the ROI as shown in the figure *Lateral ROI*.



Figure 37: Lateral ROI

11.6.2 Adjusting the Vertebral Boundaries

- 1. Select Vertebral Boundaries and Ant. Boundary.
- 2. Using **Line Mode** and **Point Mode** tools, adjust the Vertebral Boundaries as shown in the figure *Vertebral Boundaries*.



Figure 38: Vertebral Boundaries

11.6.3 Adjusting the Vertebral Bodies

- 1. Select Vertebral Bodies.
- 2. Using **Line Mode** and **Point Mode** tools, adjust the Vertebral Bodies as shown in the figure *Vertebral Bodies*.



Figure 39: Vertebral Bodies

11.6.4 Viewing the Bone Map

- 1. Select Bone Map.
- 2. If the bone map is incomplete for the vertebral bodies, adjust the boxes as shown in the figure *Vertebral Bodies in <u>Adjusting the Vertebral Bodies</u> on page 76. In most cases it will not be necessary to edit the bone map.*



Figure 40: Lateral Bone Map

11.6.5 Adjusting the Mid Regions

If activated, use the **Mid Regions** button and **Whole Mode**, **Line Mode** and **Point Mode** tools to adjust the regions that report bone density in the middle of the vertebral bodies (See the figure *Vertebral Bodies* in <u>Adjusting the Vertebral Bodies</u> on page 76).



Note

Mid Regions can be activated in the Utilities Menu, System Configuration, Analyze Tabs.

11.7 Viewing Results

- 1. Select **Results**.
- 2. Select **BMD** to display BMD results or **WA-BMD** to display width-adjusted BMD results.

11.8 Exiting Analysis

- 1. Select **Close**.
- 2. Select Report.

11.9 Generating and Printing Reports

See <u>Reports</u> on page 113.

Chapter 12 Decubitus Lateral Spine BMD Examination

Start the exam as described in <u>Using Worklist to Retrieve a Patient Record</u> on page 16. In the Select **Scan Type** window, select **AP/Decubitus**.

12.1 Performing and Analyzing the AP Scan

The AP part of the decubitus lateral spine scan is performed the same as an AP lumbar spine scan. Refer to <u>AP Lumbar Spine Examination</u> on page 25 and follow the procedure until the AP analysis is complete.

12.2 Positioning the Patient for the Decubitus Lateral Scan

- 1. Position the patient as shown in the following figures, *Decubitus Lateral Positioning* and *Spine Position*.
- 2. Place a pillow under the head to square the shoulders and to keep the spine parallel to the table.
- 3. Flex the patient's knees about 90 degrees.
- 4. Extend the patient's upper arms 90 degrees from mid-coronal plane.
- 5. Adjust the body to a true lateral position.



Figure 41: Decubitus Lateral Positioning



Figure 42: Spine Position

12.3 Positioning the C-arm for the Decubitus Lateral Scan

- 1. Position the C-arm as shown in figure *Decubitus Lateral Positioning* in <u>Positioning the</u> <u>Patient for the Decubitus Lateral Scan</u> on page 79.
- 2. Turn on the laser.
- 3. Position the cross hairs of the laser 5 cm (2 inches) below the iliac crest and centered over the spine.
- 4. Move the laser cross hairs 2.5 cm (1 inch) toward the back of the patient.
- 5. Select **Continue**.

12.4 Starting the Decubitus Lateral Scan



Warning:

If the Control Panel X-ray indicator fails to shut off within 10 seconds after the end of the scan, press the red Emergency Stop button immediately. Call your Hologic service representative before resuming operation.

- 1. Select Start Scan. The X-rays On indicator flashes until the scan stops.
- 2. Inspect the image as it is generated. When L2 to L4 is fully displayed, click **Stop Scan**.
- 3. Upon completion of an acceptable lateral image (see the following figure, *Decubitus Spine*), assist the patient off the table.



Figure 43: Decubitus Spine

12.4.1 Repositioning the Scan (if necessary)

- 1. Select **Reposition Scan**.
- 2. Position the cursor over the spine image (see the figure *Decubitus Repositioning*).



Figure 44: Decubitus Repositioning

- 3. To reposition, select and drag the image so that:
 - The iliac crest is at or below the blue horizontal positioning line **[1]** and within the lower portion of the scan field.
 - The center of the spine is aligned with the blue vertical positioning line [2].
- 4. When the spine is positioned correctly, select **Restart Scan** and repeat the scan.

12.5 Analyzing the Scan

- 1. Select Analyze Scan.
- 2. Select **Next** >>.

Note



For descriptions of Analysis Step Buttons and Toolboxes, see <u>Analysis Step Buttons</u> on page 19.

12.5.1 Defining the Global ROI

- 1. Select Global ROI.
- 2. Using the **Whole Mode**, **Line Mode** and **Point Mode** tools, adjust the ROI (see the figure *Lateral ROI*).



Figure 45: Lateral ROI

12.5.2 Adjusting the Vertebral Boundaries

- 1. Select Vertebral Boundaries and Ant. Boundary.
- 2. Using **Line Mode** and **Point Mode** tools, adjust the **Vertebral Boundaries** as shown in the figure *Adjusting Vertebral Boundaries*.



Figure 46: Adjusting Vertebral Boundaries

12.5.3 Adjusting the Vertebral Bodies

- 1. Click Vertebral Bodies.
- 2. Using **Whole Mode**, **Line Mode** and **Point Mode** tools, adjust the Vertebral Bodies (see figure *Vertebral Bodies*).



Figure 47: Vertebral Bodies

12.5.4 Viewing the Bone Map

- 1. Select Bone Map.
- 2. In most cases it will not be necessary to edit the bone map. If it is required, select **Vertebral Bodies** and re-adjust **L2** or **L3**, so they are just inside the bone edges.



Figure 48: Lateral Bone Map

12.6 Viewing Results

Select Results.

12.7 Exiting Analysis

- 1. Select Close.
- 2. Select Report.

12.8 Generating and Printing Reports

See <u>Reports</u> on page 113.

Chapter 13 IVA, IVA HD Imaging on Horizon C, W; IVA Imaging on Horizon Ci, Wi

Start the exam as described in <u>Using Worklist to Retrieve a Patient Record</u> on page 16. In the **Select Scan Type** window, select **IVA Imaging**.

13.1 Choosing the Scan Type

- 1. In the **Select Scan Type** window, remove the check mark in the **Use Default Scan Mode** box.
- 2. Select **Next** >>.
- 3. In the Include SE AP Spine Scan in IVA Exam? window, select SE AP Image.
- 4. Select **Next** >>.
- 5. In the Include SE Lateral Spine Scan in IVA Exam? window, select SE Lateral Image.
- 6. Select **Next** >>. The **Scan Parameters** window for the AP IVA scan displays.

13.2 Positioning the Patient for the AP IVA Scan

Position the patient and C-arm as described in <u>AP Lumbar Spine Examination</u> on page 25 with the following exception: the patient's shoulders should be positioned below the upper scan limit (see figure AP IVA Positioning).



Figure 49: AP IVA Positioning

13.3 Starting the AP IVA Scan

Warning:

If the Control Panel X-ray indicator fails to shut off within 10 seconds after the end of the scan, press the red Emergency Stop button immediately. Call your Hologic service representative before resuming operation.

- 1. Ask the patient to hold their breath during the scan.
- 2. Select Start Scan. The X-rays On indicator flashes until the scan stops.
- 3. Inspect the image.
- 4. When you see L4 through T4 on the image, select the **Stop Scan** button and tell the patient to breathe normally.
- 5. Select **Close**.

13.4 Positioning the Patient and C-arm for the Lateral IVA Scan

- 1. Position the patient as shown in the following figures, *Lateral IVA Positioning* and *Spine Position*.
- 2. Flex the patient's knees about 90 degrees.
- 3. Extend the patient's upper arms 90 degrees from mid-coronal plane.
- 4. Adjust the body to a true lateral position.



Figure 50: Lateral IVA Positioning



Figure 51: Spine Position

13.5 Starting the Lateral IVA Scan



Warning:

If the Control Panel X-ray indicator fails to shut off within 10 seconds after the end of the scan, press the red Emergency Stop button immediately. Call your Hologic service representative before resuming operation.

- 1. Ask the patient to hold their breath during the ten second scan.
- 2. Select **Start Scan**. The *X*-*rays On indicator* flashes until the scan stops.

13.6 IVA Analysis for Lateral IVA Scan

See IVA Analysis on page 97.

Chapter 14 IVA and IVA HD Imaging on a Horizon A System

Start the exam as described in <u>Using Worklist to Retrieve a Patient Record</u> on page 16. In the **Select Scan Type** window, select **IVA Imaging.**

14.1 Positioning the Patient

Position the patient as described in *AP Lumbar Spine Positioning in <u>Positioning the Patient</u>* on page 25 *with the following exception*: the patient's shoulders should be positioned below the upper scan limit line.



Figure 52: AP IVA Positioning

14.2 Choosing the Scan Type

- 1. In the **Select Scan Type** window, remove the check mark in the **Use Default Scan Mode** box.
- 2. Select Next >>.
- 3. In the Include SE AP Spine Scan in IVA Exam? window, select SE AP Image.
- 4. Select **Next** >>.
- 5. In the Include SE Lateral Spine Scan in IVA Exam? window, select SE Lateral Image.
- 6. Select **Next** >>. The *Continuously Press ENABLE...* message appears.

Once **Enable Lateral** is pressed the table is locked. Only the C-arm can move. If necessary, move the patient to the long axis of the laser cross hair.

14.3 Starting the AP IVA Scan



Warning:

If the Control Panel X-ray indicator fails to shut off within 10 seconds after the end of the scan, press the red Emergency Stop button immediately. Call your Hologic service representative before resuming operation.

- 1. Ask the patient to hold their breath during the scan.
- 2. Select Start Scan. The X-rays On indicator flashes until the scan stops.
- 3. Inspect the image.
- 4. When you see L4 through T4 on the image, select the **Stop Scan** button and then tell the patient to breathe normally.
- 5. Select Close. The Continuously Press ENABLE ... message displays.
- 6. Continuously press **Enable Lateral** on the Control Panel until the C-arm has rotated fully to the lateral scan position.


Figure 53: Lateral IVA Positioning

14.4 Starting the Lateral IVA Scan



Warning:

If the Control Panel X-ray indicator fails to shut off within 10 seconds after the end of the scan, press the red Emergency Stop button immediately. Call your Hologic service representative before resuming operation.

- 1. Ask the patient to hold their breath during the ten second scan.
- 2. Select **Start Scan**. The *X*-rays On indicator flashes until the scan stops.
- 3. Inspect the image.
- 4. When you see L4 through T4 on the image, click the **Stop Scan** button and then tell the patient to breathe normally.
- 5. Continuously press **Enable Lateral** on the Control Panel until the C-arm has rotated to its original position.
- 6. Select Exit.
- 7. Press **Patient On/Off** on the Control Panel and, when motion stops, assist the patient off the table.

14.5 IVA Analysis for Lateral IVA Scan

See <u>IVA Analysis</u> on page 97.

Chapter 15 IVA with BMD Examination

Start the exam as described in <u>Using Worklist to Retrieve a Patient Record</u> on page 16. In the Select Scan Type window, select IVA with BMD.

Hologic recommends performing scans in the following order:

- 1. an AP IVA scan
- 2. an AP BMD scan
- 3. a Lateral BMD scan, and
- 4. a Lateral IVA scan

15.1 Positioning the Patient

Position the patient as described *AP Lumbar Spine Positioning in <u>Positioning the Patient</u> on page 25 <i>with the following exception*: the patient's shoulders should be positioned below the upper scan limit line.

15.2 Choosing the Scan Type

- 1. In the **Select Scan Type** window, remove the check mark in the **Use Default Scan Mode** box.
- 2. Select Next >>.
- 3. In the Include SE AP Spine Scan in IVA Exam? window, select SE AP Image.
- 4. Select Next >>.
- 5. In the **Select AP Lumbar Spine Scan Mode** window, select **Express (x)**.
- 6. Select Next >>.
- 7. In the Select Lateral Scan Mode for AP/Lateral Exam window select Fast Array (f).
- 8. Select Next >>.
- 9. In the Include SE Lateral Spine Scan in IVA Exam? window, select SE Lateral Image.
- 10. Select Next >>. The Continuously Press ENABLE... message appears.

15.3 Perform the AP IVA Scan

Perform the AP IVA scan as described in <u>IVA and IVA HD Imaging on a Horizon A System</u> on page 91, starting with <u>Positioning the Patient</u> on page 91.

Once **Enable Lateral** is pressed the table is locked. Only the C-arm can move. If necessary, move the patient to the long axis of the laser cross hair.

15.4 Perform and Analyze the AP BMD Scan

Perform and analyze the AP BMD scan as described in *<u>Starting the AP Lumbar Spine Scan</u>* on page 27 and *<u>Analyzing the Lateral Scan</u>* on page 73.

15.5 Perform the Lateral BMD Scan

Perform the Lateral BMD scan as described in *<u>Starting the Lateral IVA Scan</u>* on page 94.

15.6 Perform the Lateral IVA Scan

Perform the Lateral IVA scan as described in *<u>Starting the Lateral IVA Scan</u>* on page 94.

15.7 IVA Analysis

See *IVA Analysis* on page 97.

Chapter 16 IVA Analysis

The Viewer window (see figure *IVA Viewer Window*) is described in <u>Viewer Window, Left</u> <u>Panel</u> on page 97.



Figure 54: IVA Viewer Window

16.1 Viewer Window, Left Panel

Table 16:	Viewer	Window,	Left Panel
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Control	Description
DE Scan	Used on Lateral IVA Scan only (<u><i>DE Scan</i></u> on page 101).
Multi View Button	Select to toggle between Multi View and Single View.
Visual Tools Tab	Select to display the visual tools (<i>Viewer Window, Left Panel, Visual Tools</i> <u><i>Tab</i></u> on page 98).
Analysis Tools Tab	Select to display analysis tools (see <u>Viewer Window, Left Panel Analysis</u> <u>Tools Tab</u> on page 99)
Print Report	Prints the report to the selected printer.
Print Image	Prints the image to the selected printer.
Close	Exits the analysis window and returns to the main window, saving any changes made to the scan.

16.2 Viewer Window, Middle Panel

Table 17. Vienna	Min down	Middle David	
Table 17: Viewer	ννιπιοω,	<i>Iviluule</i> Funel	

Control	Description
Image Display Area	Displays the image from the selected scan. Right click on the image to display the image control menus (<i>Image Controls on page 100</i>).
Scan ID	Appears above the image on the left.
Scan Type	Appears above the image on the right.

16.3 Viewer Window, Right Panel

Table 18:	Viewer	Window.	Right	Panel
10000 10.	1 10001	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	10,200	I WINCI

Control	Description
Patient Data Tab	Select to display patient data.
Deformity Tools Tab	Select to display deformity identification reference images and the results for each vertebra analyzed in the image.
Multi View Enabled	The viewer displays an image in both the middle and right panels.

16.4 Viewer Window, Left Panel, Visual Tools Tab

Control	Description
W-L	 Select and drag the ball in the center of the triangle to adjust the contrast and brightness of the image. For finer adjustment: 1. Right click on the image. 2. Select Adjust WL. 3. Select and drag the cursor on the image.
Revert	Returns all viewing parameters to their initial values.
Invert	Inverts the grayscale value of each pixel creating a negative of the image.
Flip	Flips the image about the center vertical axis.
÷.	Select to increase the magnification of the image.
Q	Select to decrease the magnification of the image.

 Table 19:
 Viewer Window, Left Panel, Visual Tools Tab

16.5 Viewer Window, Left Panel Analysis Tools Tab

Table 20: Assessment Area		
Control	Description	
Radio button control the following descri	ls that determine how vertebral annotations will be placed on the image. See ptions:	
Labeling Only	Position the cursor and select to place a vertebral label. Select and drag to move the label, right click to change it or add an assessment.	
Manual	Position the cursor in the center of a vertebra, and select to place a vertebral label and markers. Select between markers and drag to move the label and markers together. Select a marker and drag to move it individually. Right click to change the label or add an assessment.	
МХАрго	Position the cursor in the center of a vertebra, and select to place a vertebral label, markers, a vertebral outline (in green), and a deformity assessment based on the ratio calculated. Select between markers and drag to move everything together. Select on a marker and drag to move it until it is in the proper position (<i>Markers</i> on page 105). Right click to change the label or assessment. The asterisk indicates that the assessment is based on the ratios calculated from the vertebral heights. The vertebral deformity assessment is at the sole discretion of the physician or trained health care professional. Before printing or reporting, assessments must be changed or accepted by the physician. See <i>Interpreting the IVA Image</i> on page 103 for assessment guidelines.	

Table 21: Display	Options Area
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Control	Description
Controls that determine what vertebral annotations display on the images (see the following descriptions).	
Labels	Check to display all labels on the image.
Display Markers	Check to display all markers on the image.
Display Deformity	Check to display all deformity assessments on the image.
Display MXA <i>pro</i> Outline	Check to display all MXA <i>pro</i> outlines on the image. The outline is the software's estimation of the vertebral edges that it used for marker placement.
Refresh Image Button	Select to apply the selected display options to the last analyzed vertebrae (shows how the image will appear when viewed or printed).

Control	Description
Controls to change vertebra following descriptions).	l labels and assessments on selected vertebral annotations (see the
Label Selection Drop Down List	Select the down arrow to change the label of the selected vertebral analysis.
Assessment Selection Drop Down List	Select the down arrow to change the deformity assessment of the selected vertebral analysis.
Delete Button	Select to delete one or more selected vertebral analyses.
Reset Button	Select to remove all the new vertebral analyses and display the original analyses (if there were any) from the DICOM file.

Table 22: Change Annotations Area

16.6 Image Controls

Place the mouse pointer in the image area and right-click.

Table 23: Image Controls

Control	Description
Zoom and W-L Selection Menu (upper section)	Select one of the choices to activate: Drag Zoom Drag Pan Adjust WL
Drag Zoom	Drag the magnifying glass over the area of the image to be enlarged and release.
Drag Pan	Select and drag the image to place it anywhere in the window.
W-L Adjust	Allows fine tuning of W-L. Move in any direction to change the W and L values.

Control	Description
Zoom Control (lower section)	Select zoom control settings to enlarge or reduce the image size: Fit to Window
	25%
	50%
	100%
	200%
	400%

Table 23: Image Controls

16.7 DE Scan

If a dual energy scan is desired of one or two vertebrae on the IVA scan, use DE Scan.

The patient must remain on the table in the same position as during the Lateral IVA scan. If the patient has moved, another Lateral IVA scan must be acquired and the DE scan taken immediately after with the patient in the same position.

- 1. Select DE Scan.
- 2. Select Analysis Tools tab.
- 3. Place the mouse pointer over the upper green line (at the top of the image) and drag it to the top of the desired region.



Note

If the image is magnified, the green lines may be outside the viewing area. To see the lines, select Fit to Window.



Figure 55: DE Scan 1 (Top of Region)

- 4. Place the mouse pointer over the lower green line at the bottom of the image and drag it to the bottom of the desired region.

Figure 56: DE Scan 2 (Top to Bottom of Region)

- 5. Select Close.
- 6. Start the new scan. The patient may breathe normally during the Dual Energy scan.

Chapter 17 Interpreting the IVA Image

IVA images are to be interpreted by a physician or properly licensed practitioner. IVA images are not intended for general radiological diagnosis, but are intended to be used for evaluation of vertebral deformities.

The anatomy of the spine is shown in the figure *The Human Spine*, including vertebral level labels. IVA images typically include levels **T4** to **L4**. Following the classification scheme of Genant., the figure, *Vertebrae Deformities*, shows examples of a typical vertebral body shape and examples of deformed vertebral shapes.



Figure 57: The Human Spine

The human spine usually consists of:

- 1. 7 Cervical Vertebrae
- 2. 12 Thoracic Vertebrae
- 3. 5 Lumbar Vertebrae
- 4. Ossacrum



- 1. Typical Vertebral Body
- 2. Severe Wedge Deformity

3. Severe Biconcavity Deformity Typical vertebral body shown with

severely deformed vertebrae. Adapted from Genant, H.K., C.Y. Wu, et al. (1993). "Vertebral fracture assessment using a semiquantitative technique." J. Bone Miner Res 8(9): 1137-48.

Figure 58: Vertebrae Deformities

For detailed information regarding vertebral deformity evaluation and the utility of vertebral deformity assessment in clinical osteoporosis evaluation, see the references and organizations listed in <u>Interpreting Results</u> on page 123.

Chapter 18 Markers

To define the shape of one or more vertebrae, markers are placed on the posterior, anterior and midpoints, as shown in the following figure.



Figure 59: Marker Placement

The proper placement of these six markers is found in "*The Appendix to Chapter 20: Point Placement in Vertebral Morphometric X-ray Absorptiometry*" by Jacqueline A. Rea in '*The Evaluation of Osteoporosis: Dual Energy Absorptiometry and Ultrasound in Clinical Practice,* Second Edition', pages 456-457.

18.1 Using Markers

18.1.1 Flipping the Image

The IVA scan image should initially display the vertebrae on the left. Before adding Markers make sure the spine is on the left. Select **Flip** if necessary.



Note

The image can be flipped after Markers are added.

18.1.2 Adding Markers

- 1. Select Markers.
- 2. Right click on the image and select **Add Marker**.



Figure 60: Adding a Marker



Note

The software attempts to determine what vertebral label based on the location of the pointer when the user clicks. After adding the first label to the image, if the assumption made by the software is wrong, change the label before adding another. As additional labels are added, if any label is wrong, change that label before adding another

18.1.3 Selecting Markers

1. Place the mouse pointer in the middle of the six markers and click,



Figure 61: Selecting a Marker

or select the marker data in the Result Block.

L3	22.6	21.1	22.9	1.01	0.93
L4	21.3	19.3	22.4	1.05	0.91
	(2.)	r 1 7	<u> </u>	1 D	1. D1 1

Figure 62: Marker Data in the Result Block

The selected markers appear yellow (red if the image is inverted). The unselected markers appear red (cyan if the image is inverted).

18.2 Result Block

The result box for the Lateral and the AP IVA scans must be interpreted differently because of the positions of the spine in the scan.

18.2.1 Result Block for Lateral IVA Scans

he Vertebral Assessment panel is displayed in the lower right corner of the window when the viewer is in Single View mode. This panel lists the results in tabular form for each vertebra analyzed in the image displaying in the image area.

There are two lines of data for each possible vertebral analysis.

- The first line contains the vertebra's label, Post, Mid, and Ant Height in mm and the percentage deformation for Wedge, Biconcave, and Crush.
- The second line contains the deformity type along with its severity. Some of this data may be blank based on which kind of analysis of the vertebra was performed. Two blank lines appear for unanalyzed vertebrae.
- The last line of data in this table shows the Standard Deviation values for each of the displayed types of data. For example, the Post Hght (mm) has a standard deviation of ± 1 mm; Wedge percent deformation has a standard deviation of ± 5%.

	Heigh	it(mm)		Perce	ent Deforma	tion
Label	Post	Mid	Ant	Wedge	Biconcave	Crush
	Defor	mity (G	rade)			
T7	18.0	16.0	18.0	0.0%	11.1%	0.0%
T8	18.0	16.0	18.0	0.0%	11.1%	0.0%
Т9	18.0	16.0	18.0	0.0%	11.1%	0.0%
T10	18.0	16.0	18.0	0.0%	11.1%	0.0%
T11	18.0	16.0	18.0	0.0%	11.1%	0.0%
T12	18.0	16.0	18.0	0.0%	11.1%	0.0%
L1	18.0	16.0	18.0	0.0%	11.1%	0.0%
L2	18.0	16.0	18.0	0.0%	11.1%	0.0%
L3	18.0	16.0	18.0	0.0%	11.1%	0.0%
L4	18.0	16.0	18.0	0.0%	11.1%	0.0%
Std De	v 1.0	1.0	1.0	5.0%	5.0%	5.0%

Figure 63: Result Block

Table 24: Result Block Labels

Label	Description	
Post Hght (mm)	The height of the vertebrae between the superior and inferior posterior markers in millimeters.	
Mid Hght (mm)	The height of the vertebrae between the superior and inferior midpoint markers in millimeters.	
Ant Hght (mm)	The height of the vertebrae between the superior and inferior anterior markers in millimeters.	
Wedge	The distance in millimeters when the Ant Hght is divided by the Post Hght .	
Mid Wedge	The distance in millimeters when the Mid Hght is divided by the Post Hght .	

18.2.2 Result Block for AP IVA Scans



Note

Hologic does not recommend placing Markers on the AP IVA scan.

The Result Block for the AP IVA scan will appear the same as the above Lateral IVA scan but the labels are used differently because of the position of the spine. The **Mid Hght**, **Wedge** and **Mid Wedge** are the same but the **Post Hght** must be interpreted as the **Right Side** of the vertebrae and the **Ant Hght** must be interpreted as the **Left Side** of the vertebrae according to the following:

Lateral IVA Scan	AP IVA Scan
Post Hght (mm)	Right Side (mm)
Ant Hght (mm)	Left Side (mm)

The **Left Side** and **Right Side** labels will not appear in the AP IVA scan Result Block report or be printed on any reports.

18.3 Printing

The viewer provides two modes for printing what is viewed on the screen: Print Report and Print Image.

Printing is integrated into the standard Windows print architecture. That is, you can select the output device, number of copies, change properties, and so forth. You can also select a print preview to view what will be printed on your screen.



Note

If any new *CADfx* analyses have been placed on the image, you will have to accept the analyses before the results can be printed or viewed on the monitor screen.

18.3.1 Print Report

Print Report can be used only in Single View mode.

18.3.2 Print Image

Print Image can be used in both Single and Multiple View modes.

For additional Report types, refer to <u>*Reports*</u> on page 113.

Chapter 19 Compare and Follow-up

19.1 Restore a Baseline or Prior Scan

If the patient's baseline or prior scan is not currently on the system, locate and restore it (*Locate Scans* on page 129 and *Restore Scans* on page 129).

19.2 Evaluate the Baseline or Prior Scan

Ensure that the baseline or prior scan is analyzed correctly. If it is not, re-analyze and archive it and all subsequent scans.

19.3 Perform the Follow-Up Scan

For the procedure to perform the follow-up scan refer to the following:

Table 26: Follow-up Scan Procedures			
Scan	Section		
AP Lumbar	<u>AP Lumbar Spine Examination</u> on page 25		
Нір	<u>Hip Examination</u> on page 31		
Forearm	Forearm Examination on page 47		
Whole Body	Whole Body Examination on page 57		
AP/Lateral	<i>Supine AP/Lateral Spine BMD Examination (Horizon A)</i> on page 71		
Decubitus Lateral	Decubitus Lateral Spine BMD Examination on page 79		

Then, from the Exit Exam screen, select Analyze Scan.

19.4 Analyze the Follow-up Scan Using Compare Analysis

- 1. Auto compare to the baseline, or prior, scan depending upon analysis configuration.
- 2. Select Results.
- 3. Select Close.
- 4. If necessary, use the Analysis Step Buttons and Toolbox tools to compare the followup scan. Match the analysis of the follow-up scan to the baseline or prior scan.



Note

Use Compare Analysis for best rate of change results.

19.5 Generate the Rate of Change Report

1. Select **Report** in the **Exit Analysis** window.



If the regions for prior scans do not exactly match but partially match the current scan, a report is generated for only those regions that match.

- 2. Select the **Rate of Change** check box.
- 3. Select **Print**.

Note

Chapter 20 Reports

- 1. Select **Report** in the main window.
- 2. Select the name of the patient, then click **Next** >>.
- 3. Select the desired scans, then click **Next** >>. For creating dual hip scan pairs (*<u>Create</u>* <u>*Hip Pairs for Dual Hip Rate of Change Reports*</u> on page 115).
- 4. For comments on the printed report, select **Edit comment...** (see the figure *Report Blocks* in *Report Information Blocks* on page 113).
- 5. Select one of the following:
 - Choose the report type by clicking its box. (*DICOM Report* on page 120).
 - Select DICOM / IVA report. (DICOM Report on page 120).
 - Select **Generate DxReport**. See *DxReport User Guide* (MAN-02331). You can configure the DxReport from the Report Tab under the System Configuration screen.
- 6. Select **Print**.

20.1 Report Information Blocks

Reports contain *blocks* of information which vary slightly depending upon the type of report you choose. Refer to the following figure and the following text for an explanation of report blocks.



Figure 64: Report Blocks

20.2 Edit Comments

Note

1. In the *Print* window, select **Edit comment...**

- To select from the list of predefined comments, select the down arrow.
- To enter a new comment, click in the *Comment* text box.



New comments are not added to the list of predefined comments.

2. Select **Update**.

20.3 Rate of Change Report

The Rate of Change Report tracks changes in results over time and includes:

- Detailed patient and scan information
- Scan date, patient age, BMD, and T-score results for each visit
- Changes in results provided in percent (%) and/or as an absolute difference (gm/cm²) vs. Baseline and previous scans
- Scan image with ROI and bone map outline for current scan
- BMD vs. Age reference curve graph for each baseline and subsequent scans
- 10-year Fracture Risk (Hip scans only)
- Body Composition Rate of Change Results (Whole Body scans only)

For more information, see *Interpreting Results* on page 123.

20.3.1 Remove Asterisks (*) and Pound Signs (#) from Reports

Reports may include asterisks (*) and a pound signs (#) to indicate that scan types and analysis methods do not match. To prevent asterisks (*) and pound signs (#) from appearing in reports:

- 1. Select **System Configuration > Report tab**. The General tab is displayed.
- 2. Select Rate of Change.
- 3. Select the **Configure** button. The Configure Rate of Change dialog box is displayed.
- 4. Select the **Results Block tab**.
- 5. Uncheck Indicate Different Scan Types or Analysis Methods.
- 6. Select **OK**, then **OK** again.

20.3.2 Create Hip Pairs for Dual Hip Rate of Change Reports

Dual Hip Rate of Change report provides information about result changes in hip "pairs". A *hip pair* includes a right hip scan and a left hip scan performed within 14 days of each other.

- 1. Access the patient scan list as you would for any report (*<u>Reports</u>* on page 113).
- 2. Select a left and right scan—one scan is the most recent. The Match Pairs of Scans dialog box is displayed.
- 3. Select a right hip scan from the left list box.
- 4. Select a left hip scan from the right list box. The down arrow is enabled.
- 5. Select the hip pair from the Dual Hip Pairs list.
- 6. Select **OK**.

20.4 Body Composition Reports

APEX software can display the DXA measures along with a representative color image mapping of "fat" and "lean" tissue. See figure *Advanced Body Composition Report*, in <u>BCA</u><u>*Results*</u> on page 116.

A Rate-of-Change report can also be generated to display the trend of serial DXA Body Composition measurements over time. See figure *Advanced Body Composition Rate of Change Report* in <u>BCA Rate of Change Results</u> on page 117.



Note

The images on these reports should not be used for diagnosis.

20.4.1 BCA Results

The report blocks and graphs for BCA results (see the following figure) are listed in the following tables. For descriptions of images, see *DICOM Report* on page 120.



Figure 65: Advanced Body Composition Report

Report Block	Description
Body Composition Results	Results for the standard SubRegions (arms, trunk, legs and head), subtotal (excludes head), total (includes head) and Android and Gynoid regions.
Adipose Indices	Results and indices for the subject's adipose tissues.
Lean Indices	Results and indices for the subject's lean mass tissues.

Graph	Description
Age vs. Total Body % Fat Graph ¹	Graph of the subject's age vs. Total Body % fat.
WHO BMI Classification	Scalar representation of the subject's WHO Body Mass Index Classification.

1. User Configurable

20.4.2 BCA Rate of Change Results

The Advanced Body Composition[™] report blocks and graphs for BCA Rate of Change results (see the following figure) are listed in the following tables.



Figure 66: Advanced Body Composition Rate of Change Report

Report Block	Description
Total Body % Fat Results ¹	Results, indices and comparison data for the subject's % fat.
Total Fat Mass Results*	Results, indices and comparison data for the subject's total fat.
Total Lean Mass Results*	Results, indices and comparison data for the subject's lean plus BMC mass.
Total Mass Results*	Results, indices and comparison data for the subject's total mass.

Table 29: Advanced Body Composition Rate of Change Re	port Fields
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1. User Configurable

Table 30: Advanced Body Composition Rate of Change Graph Fields

Graph	Description
Age vs. Total Body % Fat Graph ¹	Graph of the subject's age vs. Total Body % fat.
Compartmental Trending*	Graph of the changes in Total Body Fat Mass and Total Body Lean Mass

1. User Configurable

20.4.3 Body Composition Reports and Reference Database Comparisons

In 2008 NHANES released a population-based DXA whole body dataset acquired on Hologic scanners. Selected DXA measures can be compared to gender, ethnicity, and age-specific reference databases developed from the NHANES whole body dataset released in 2008.⁴

The software can also display the DXA measures along with a representative color image mapping of "fat" and "lean" tissue (see the figure in <u>BCA Results</u> on page 116). The color image displays the relative amounts of fat and lean tissue in the DXA image, with yellow regions representing regions with higher %Fat and orange and red regions indicating progressively lower %Fat. Bone containing regions are indicated in blue. Beside the color image is an image that is brighter in regions of greater tissue thickness and darker in thinner tissue. It is used to display the region of interest lines placed by the operator during analysis. Beneath the images the phrase "Image not for diagnostic use" appears, informing the user that the image should not be used for diagnosis. The color image displays the relative distribution of fat and lean tissue in the image and does not contain diagnostic or quantitative information.

A reference curve is generated for Total Body % Fat versus Age that is matched to the patient's gender and ethnicity. The graph provides a graphical representation of the patient's measurement relative to age-matched peers. The midline of the graph represents the median reference value and the upper and lower shaded regions define the 95% confidence interval for the plot. Note that the upper and lower shaded regions of the reference curve may not be exactly equal in size; this is an indication that the underlying reference data are not normally distributed. An algorithm that adjusts for skewness in the underlying reference data has been implemented to provide accurate T-scores, Z-scores, and percentiles.

A Body Mass Index (BMI) scale appears on the report to display the patient's calculated BMI based upon the patient's height and weight as entered by the operator. Proper height and weight should always be verified before interpreting the results displayed in the BMI scale. Above the scale the WHO BMI classification appears along with an explanation of the health risks associated with a high BMI. Beneath the graph a paragraph appears that summarizes the U.S. Surgeon General's Health Consequences for overweight and obesity

(<u>http://www.surgeongeneral.gov/topics/obesity/calltoaction/fact_advice.htm</u>). For more information, see <u>http://www.surgeongeneral.gov/topics/obesity/calltoaction/CalltoAction.pdf</u>.

Patient results can be compared to values in the Hologic Whole Body DXA Reference Database both graphically and quantitatively (see the figure in <u>BCA Results</u> on page 116). The graphical plot displays the reference values along with the subject's measured DXA value. In adults, the quantitative comparison provides a Z-score or an Age-matched (AM) Percentile value and a T-score or a Young Normal Percentile value (YN). For subjects less than 20, only a Z-score or an Age-matched (AM) Percentile value is provided. A simple mathematical transformation is used to convert Z-scores and T-scores to Age-matched and Young Normal Percentile values, respectively, depending upon whether the user configures the software to display Z-and T-scores (standard deviation scores) or percentiles.

A Rate-of-Change report can also be generated to display the trend of serial DXA Body Composition measurements over time (see the figure in <u>BCA Rate of Change Results</u> on page 117). The Total Body %Fat curve at the top left of the report displays the trend of the Total Body %Fat results over time. These measurements are displayed on an age, gender, and ethnicity-matched reference curve from the Hologic Whole Body DXA Reference Database.

Immediately below the Total Body %Fat curve is another plot labeled "Compartmental Trending." This plot provides a graphical display of the changes in Total Body Fat Mass (yellow shaded region) and Total Body Lean Mass (blue shaded region). Total Mass, i.e. the sum of yellow Fat Mass region plus the blue Lean Mass region, is indicated by the uppermost line of the plot.

20.5 Pediatric Reports

The following figure displays a graph of the subject's measurement plotted on a gender and ethnicity-matched reference curve. Below the plot are corresponding results based upon the available measures selected in the system configuration for this report. Each DXA measure is plotted on a percentile scale and the Z-score and centile for the subject's measurement relative to gender and ethnicity-matched peers is provided at the far right of the scale. Reference data from Hologic, the Bone Mineral Density in Childhood Study, and NHANES is used for Z-scores and percentiles.



Figure 67: Pediatric Spine Report

20.6 DICOM Report

Create and Send a DICOM Report.

20.6.1 Select a DICOM BMD Report Type

- 1. Select the desired scans.
- 2. Select the **DICOM BMD** report type.

20.6.2 View Scan Details and Enter Patient Biography Fields

- 1. Select the scan on the **DICOM Report** window.
- 2. Select Scan Details.
- 3. Select the **Details** tab.
- 4. Edits are allowed on the following fields:
 - Accession Number 16 characters maximum
 - Study Instance UID 28 characters maximum
 - HL7 Field 1 64 characters maximum
 - HL7 Field 2 64 characters maximum
 - HL7 Field 3 64 characters maximum



Note

The HL7 fields are user definable and provide additional information.

- Operator 5 characters maximum
- Height 5 characters maximum
- Weight 5 characters maximum
- Scan Comment 100 characters maximum
- 5. Select the **Identification** tab to view the scan information.
- 6. Select **OK** to save edits; select **Cancel** to close without saving.

20.6.3 Enter the Accession Number and User Defined Entries

- 1. Select a scan on the *DICOM Report window*.
- 2. Select **Save As** or **Send**.
- 3. If the selected scan does not have an accession number enter one, then press **Enter** or **OK**.
- 4. Select **Cancel** if the accession number is unknown or will be entered later.
- 5. If prompted to add additional user defined entries, enter and select **OK** for each dialog box.

20.6.4 Preview a DICOM Report

Select the **Preview** button to view the *DICOM* report before you save or send the report.

20.6.5 Print a DICOM Report

Click the **Print** button on the *DICOM Preview* screen to print the *DICOM* report to the local default printer.

20.6.6 Save a DICOM Report

Select the **Save As** button to save a *DICOM* report as a file to your desired location.

20.6.7 Send a DICOM Report

1. Select the scans on the *DICOM Report window*.

Assign the same Accession Number to all scans associated with this patient's visit.

2. Select Send.

For each selected scan, a *DICOM* report is generated, placed in the queue, and sent in the order the report was placed in the queue.

To view send status, see *View the Queue* on page 122.

20.6.8 Sort the Scan List

Select any heading to sort the scan list by ascending or descending order.

20.6.9 View the Queue

Select the View Queue button to view scans in the queue waiting to be sent.

View a History of Sent Reports

Select the View Log button on the View Queue dialog box.

Update the status of DICOM Reports in the Queue

Select the **Refresh** button on the *View Queue* dialog box.

Delete a DICOM Report from the Queue

Select the **Delete** button on the *View Queue* dialog box.

20.6.10 Close a DICOM Report

Select the **Cancel** button or the **<<Back** button on the *DICOM Report window*.

20.7 DxReport

20.7.1 Create a DxReport

- 1. Select Interpreting Physician.
- 2. Select or clear the check box for Include rate of Change.
- 3. Select Generate DxReport.

A Word report will be generated in accordance with the configuration settings see *DxReport Users Guide* MAN-02331.



Caution

A qualified medical professional must review each patient report that DxReport generates before the report is released

Chapter 21 Interpreting Results

Websites

- <u>www.iscd.org</u> Particularly, the ISCD Official Positions
- <u>www.nof.org</u> Particularly, the NOF Physician's Guide
- <u>www.iofbonehealth.org</u> Particularly, Health Professionals, including Educational Tools and Slide kits.
- <u>http://www.aace.com</u> American Association of Clinical Endocrinologists

Publications

- U.S. Department of Health and Human Services. Bone Health and Osteoporosis: A Report of the Surgeon General. Rockville, MD: U.S. Department of Health and Human Services, Office of the Surgeon General, 2004.
- Kanis, JA on behalf of the World Health Organization Scientific Group (2007), Assessment of osteoporosis at the primary healthcare level. Technical Report. World Health Organization Collaborating Centre for Metabolic Bone Diseases, University of Sheffield, UK. 2007:Printed by the University of Sheffield.
- The Evaluation of Osteoporosis: Dual Energy Absorptiometry and Ultrasound in Clinical Practice, Second Edition; Blake, G. M., Walgner, H. W., Fogelman, I., [©] Martin Duritz Ltd 1999
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21.1 About FRAX

Fracture Risk Assessment as calculated by FRAX has specific age, weight, and height limits. The age range is between 40 years and 90 years. If you enter an age between 20 and 40 years, FRAX will calculate the probability of fracture at the age of 40 years. If you enter an age above 90 years, FRAX will calculate the probability of fracture at the age of 90 years. The weight range is between 25 kg (55 lbs) and 125 kg (276 lbs); the height range is between 100 cm (39 in) and 220 cm (86 in). If you enter a weight or height outside of those ranges, FRAX will calculate the probability of fracture at these limits.

- BMI is calculated by the software using patient's weight and height data.
- Femoral Neck BMD value is obtained from the patient's most recent Hip scan analysis.



Note

The FRAX assessment does not tell you who to treat which remains a matter of clinical judgment. In many countries, guidelines are provided that are based on expert opinion and/or on health economic grounds.

21.2 FRAX Limiting Criteria

NOF/ISCD recommends the use of the FRAX limiting criteria for US configurations. However, you can configure FRAX to remove the limiting criteria. For more information see <u>*Configuring FRAX*</u> on page 125.

Choose Yes or No for the FRAX limiting criteria as follows.

21.2.1 Previous hip or vertebral fracture

Choose **Yes** if the patient had a prior hip or vertebral fracture (clinical or morphometric). If yes, FRAX will not be calculated.

21.2.2 Treatment for osteoporosis

Choose **Yes** if the patient is currently being treated for osteoporosis. If yes, FRAX will not be calculated.

Examples of "untreated" patients include:

- No ET/HT or SERM for the past one year
- No calcitonin for the past one year
- No PTH for the past one year
- No denosumab for the past one year
- No bisphosphonate for the past two years (unless it is an oral taken for less than 2 months)



Calcium and vitamin D do NOT constitute "treatment" in this context

21.2.3 Premenopausal woman

Choose **Yes** if the woman had menses in the last year or is breast feeding. If yes, FRAX will not be calculated.

21.2.4 Configuring FRAX

Note

To remove the FRAX limiting criteria:

- 1. From the **Utilities** menu select **System Configuration** > **Report tab**.
- 2. Ensure the **General** tab is selected and in the **Ten Year Fracture Risk** section select **Configure**.
- 3. In the **Display Settings** section, select **Use IOF configurations**.
- 4. Select OK.

21.3 About 10-year Fracture Risk - All Countries

The following was adapted from the WHO Collaborating Centre for Metabolic Bone Diseases, University of Sheffield, UK website, January 2008, and used with permission.

The FRAX tool has been developed by WHO to evaluate fracture risk of patients. It is based on individual patient models that integrate the risks associated with clinical risk factors as well as bone mineral density (BMD) at the femoral neck.

The FRAX models have been developed from studying population-based cohorts from Europe, North America, Asia and Australia.

The FRAX algorithms give the 10-year probability of fracture. The output is a 10-year probability of hip fracture and the 10-year probability of a major osteoporotic fracture (clinical spine, forearm, hip or shoulder fracture).

For answers to frequently asked questions about FRAX, see <u>FRAX FAQs</u> on page 171.

21.4 FRAX Risk Factors

See the following table to determine appropriate response for FRAX risk factors.

Risk Factor	Response
Country Code	<i>Select the desired Country (Ethnicity) by clicking the down arrow and choosing from the list.</i>
Previous fracture	Check Yes if the patient sustained a broken bone after age 40 excluding fractures of the skull, hands and feet.
	A previous fracture denotes more accurately a previous fracture in adult life occurring spontaneously, or a fracture arising from trauma which, in a healthy individual, would not have resulted in a fracture.
	Note Note: A fracture detected as a radiographic observation alone, i.e. seen with IVA, counts as a previous fracture
Parental fractured hip	Check Yes for a history of hip fracture in the patient's mother or father.
Current smoking	Check Yes if the patient currently smokes tobacco.
Glucocorticoids	Check Yes if the patient is exposed to oral glucocorticoids or has been exposed to oral glucocorticoids for more than 3 months at a dose of prednisolone of 5mg daily or more (or equivalent doses of other glucocorticoids).
Rheumatoid arthritis (RA)	Check Yes if the patient has a confirmed diagnosis of rheumatoid arthritis by a physician, (i.e., not a self-diagnosis of RA).
Secondary osteoporosis	Check Yes if the patient has a disorder strongly associated with osteoporosis. These include type I (insulin dependent) diabetes, osteogenesis imperfecta in adults, untreated long- standing hyperthyroidism, hypogonadism or premature menopause (<45 years), chronic malnutrition, or malabsorption and chronic liver disease.
Alcohol 3 or more units per day	Check Yes if the patient takes 3 or more units of alcohol daily. A unit of alcohol varies slightly in different countries from 8-10g of alcohol. This is equivalent to a standard glass of beer (285ml), a single measure of spirits (30ml), a medium-sized glass of wine (120ml), or 1 measure of an aperitif (60ml).

Whenever there is uncertainty by the patient as to an answer, mark it as **No**.

21.5 References

The development of the models for fracture risk assessment has been based on a program of work undertaken at the WHO Collaborating Centre for Metabolic Bone Diseases at Sheffield University. Further details are provided in the QDR Reference Manual. These include papers on the modeling approach, meta-analyses to evaluate bone mineral density and other risk factors, and recent reviews.
Chapter 22 Scans

22.1 Archive Scans

- 1. Select Archive Scans in the main window.
- 2. Select the scans to be archived.
- 3. Select Archive Scans. The Transfer Results window is displayed.
- 4. Select OK.

Hologic recommends an immediate second archive of the same scans to another cartridge or disk. Creating the second archive protects against scan loss in case of damage to the first cartridge or disk.

22.2 Locate Scans

Locate scans archived to a PACS server using Query/Retrieve Scans. See <u>Query/Retrieve</u> <u>Scans</u> on page 130.

- 1. Select **Locate Scans** in the main window.
- 2. Select the patient's name, then select Locate Scans.
- 3. Select scans from the Primary Archive tab.



Note

If you cannot restore scans from the Primary Archive media screen, contact your Hologic service representative before using the Secondary Archive media.

- 4. Place the cartridge or disk with the correct Label into the disk drive.
- 5. Select **Restore Scans**.
- 6. Select OK.

22.3 Restore Scans

- 1. Select the **Archive** dropdown menu in the main window, then select **Restore Scans**.
- 2. Select the scans to be restored and select **Restore Scans**.
- 3. Select OK.

22.4 Copy Scans

- 1. Select the Archive dropdown menu in the main window, then select Copy Scans.
- 2. Select the scans to be copied to the specified location:
- 3. Select Copy Scans.
- 4. Select OK.

22.5 Query/Retrieve Scans

Use Query/Retrieve to locate and retrieve scans from a configured PACS server into the QDR system.

- 1. Select **Query/Retrieve** from the **Archive** drop-down menu on the main screen.
- 2. Complete **Query** parameters as desired.
- 3. Select **Optional Filters** to add study level filters to the query or go to Step 5.
- 4. Complete **Study level filters** as desired.
- 5. If more than one active location is configured, select archive location (**Destination**).
- 6. Select Query.
- 7. In the **Retrieve** section, select the study or studies to retrieve.
- 8. Select **Retrieve**.

Chapter 23 Perform System Backup

Perform a System Back to copy the system database to removable media or to a directory on a computer network.

- 1. Select **System Backup** in the main window.
- 2. Enter the backup location (or accept the default location).
- 3. Accept the default backup filename, or enter a different filename (not recommended).



Caution

Changing the backup filename makes restoring the correct file difficult.

4. Select OK.

Chapter 24 Clean the System

24.1 Clean the QDR and Computer Components

- 1. Turn off power at the main breaker.
- 2. Use a soft, damp cloth to wipe surfaces. If necessary, use a mild detergent to remove dirt or debris.
- 3. Turn on power at the main breaker.

24.2 Clean the Table Pad

Use a simple solution of neutral soap and lukewarm water. Let dry thoroughly before scanning.



Note

Do not remove the cover from the table pad to clean or disinfect.

If cleaning does not produce satisfactory results, contact your Hologic representative to order a replacement table pad.

24.3 Disinfect the Table Pad

1. Use a phenolic, or quaternary type disinfectant in the concentration recommended by the manufacturer.



Note

Disinfectants applied at full concentration, or in highly concentrated solutions may damage the fabric.

Idophor (i.e., iodine bearing) type disinfectants may result in staining if these are not treated with a diluted (10 to 1) bleach solution within 20 minutes of application or spillage.

2. Let dry thoroughly before scanning.

24.4 Clean Accidental Spills

Discourage the presence of liquids in the vicinity of the Horizon system.

1. Wipe the spill immediately with a slightly dampened sponge. If the spill penetrates the system's interior, turn off power at the main breaker immediately.



Contact your Hologic service representative if you need assistance.

2. Let the table pad dry thoroughly before scanning.



Note

Note

Moisture on the table pad can distort X-ray transmission and produce erroneous analysis results.

3. Turn on power at the main breaker when the unit is thoroughly dry.

Chapter 25 Emergency Procedures

25.1 Power Failure

Turn off all equipment. When the power returns, it may be unstable. Wait a few minutes before turning on the equipment.

25.1.1 Shutting Down

- 1. If the Horizon was operating when the power failure occurred, assist the patient from the table.
- 2. Turn off the computer.
- 3. Turn off the circuit breaker (see the following figure).



Figure 68: Circuit Breaker and Indicator

Horizon W Rear View

- 1. Circuit Breaker
 - a. Indicator
 - b. Switch
- 2. Circuit Breaker Location

After power is restored

- 1. Wait a few minutes for power to stabilize, then turn on the circuit breaker. The green indicator goes on.
- 2. Perform System Startup and Shutdown (*System Startup and Shutdown* on page 11).

25.2 Failure During Operation

- 1. On the Control Panel, press the red Emergency Stop Button. The table and C-arm immediately stop moving, and the X-rays and laser turn off.
- 2. Assist the patient off the table.
- 3. Turn off the circuit breaker (see the figure in *Shutting Down* on page 135).
- 4. Disconnect the power cord from the AC outlet (if possible).
- 5. Call your Hologic service representative.

25.3 Loss of Power

If the circuit breaker (see the figure in <u>Shutting Down</u> on page 135) has been turned off (not due to equipment failure), or the system has been unplugged from the wall outlet, restore power as follows:

- 1. If necessary, place the power cord into the AC outlet.
- 2. Turn on the circuit breaker. The green indicator goes on.
- 3. Perform System Startup (*System Startup* on page 11).
- 4. If the system does not turn on, call your Hologic service representative.

Chapter 26 Dose Area Product Meter

The Dose Area Product (DAP) Meter measures the amount of radiation a patient receives during an examination. The measurement is displayed when the examination is exited.

26.1 Turning the DAP Meter On and Off

- 1. Select **Utilities** in the main window Menu bar.
- 2. Select **System Configuration** in the drop-down list.
- 3. Select the System Tab and place a checkmark in the **Report Dose Area Product** box.
- 4. Select **OK**.

Chapter 27 Utilities

Use Utilities to find, move, store, and edit patient biographies, patient scans, patient data, and system information. Select **Utilities** in the main window Menu Bar to access Utilities. Select Help within each utility for more information about a specific utility.

27.1 System Configuration

Use to change the configuration settings in functional areas of the system. Select the tab for the function desired.

27.2 Usage

Use to display and print billing information for leased systems. Select Usage.

27.3 Database Tools

Use to move patient, reference, and QC data to and from other databases.

27.3.1 Patient Management

Use to erase patient and scan data. You must erase all scans listed for a patient before you can erase the patient. Also use Patient Management to select a new baseline scan.

27.3.2 Export

Use to move data into a new or existing database on another system. Select Export.

27.3.3 Import

Use to move data from another system into Horizon. Click Import.

27.3.4 Reconcile

Compares the system database with the scan files in the system directory and automatically corrects discrepancies.

27.3.5 Patient Callback

Provides a list of patients based upon selected Last Exam Date and T-score values. Select **Callback List**.

27.3.6 Auto Baseline Utility

Sets the baseline scan of all restored scans (patients and scan types) to the oldest scan.

27.4 Scan File Look

Lists records in the scan files. Select Scan File Look.

27.5 Scan File Plot

Displays a plot of records in the scan files. Select **Scan File Plot**.

27.6 Emergency Motion

Use only when directed to do so by an authorized Hologic representative.

27.7 AP Reposition

Use only when directed to do so by an authorized Hologic representative.

27.8 Factory Utilities

Hologic use only.

27.9 Service Utilities

Only used by an authorized Hologic representative.

27.10 Reference Curve

Use to set up and manage custom reference curves.

27.10.1 Editor

Functions available under Editor include: **New, Edit, Copy, View and Delete**. Select the desired function.

27.10.2 Add Ethnicity

Use to add a new ethnicity name to the Ethnicity selection list used in reference curve descriptions.

27.10.3 Restore

Use to restore the reference curve database to the original state as supplied by Hologic.

27.11 Rebuild Archive Index

Rebuilds the archived scans index file. Use if unable to see scans on archive media known to contain scans. Select **Rebuild Archive Index**.

27.12 Install Options

To perform the Install Options function:

- 1. Obtain a license key from Hologic for the option you want to install.
- 2. Select **Utilities > Install Options** from the main window **Menu Bar**.
- 3. Type the license key obtained from Hologic in the License Key field.
- 4. Select Install Option.
- 5. Follow instructions on the screen.
- 6. Select another option to install or select **Close.**

Chapter 28 Reference Curve

Standard reference curves are provided by Hologic based on studies performed on Hologic QDR bone densitometers. References curves are sets of data points for a given sex, ethnicity, and scan type/region and specify standard deviation and skew value for the point.

Reference Curve allows users to setup and manipulate custom reference curve data.

Using Reference Curve you can:

- view reference curve record data
- create new reference curve records
- modify reference curve records (Hologic supplied reference curve records cannot be modified)
- delete reference curve records (Hologic supplied reference curve records cannot be deleted)
- create new ethnicity groups
- restore database to Hologic-supplied reference curves

Hologic-supplied reference curves cannot be edited or deleted. However, Hologicsupplied curves can be marked as current or non-current and they can be copied and edited to create a new reference curve.

28.1 Starting Reference Curve Editor

1. Select **Utilities > Reference Curve > Editor** from the Menu Bar of the main window.



Modifying the contents of the Hologic supplied Reference Curve Database may change the T-scores, Z-scores, Peak Reference and Age Matched reference results.

2. Select OK to display the Reference Curve Editor dialog box.



Note

An *H* in the Hologic field indicates a Hologic provided reference curve record that cannot be modified or deleted

28.2 Viewing Reference Curve Data

- 1. Start the Reference Curve Editor (*Starting Reference Curve Editor* on page 143).
- 2. Locate and select the reference curve record line to view.
- 3. Select View.

The upper section of the View Reference Curve dialog box contains the reference curve description information. The lower section contains the reference curve point data. No changes can be made on this dialog box.



Note

Refer to <u>Creating New Reference Curve Records</u> on page 144 for descriptions of the fields on this screen.

- 4. Select **Close** to return to the Reference Curve Editor dialog box.
- 5. Select **Close** to return to the main screen.

28.3 Creating New Reference Curve Records



Note

New reference curve records can be easily created by copying an existing reference curve record where most of the data is to be the same as that of the existing record. Refer to <u>Copying a Reference Curve Record</u> on page 146.

- 1. Start the Reference Curve Editor.
- 2. Select New. A curve is added to the database.
- 3. Select OK to display the New Reference Curve dialog box.

The upper section of the New Reference Curve dialog box contains the reference curve description information. The lower section lists the reference curve point data as it is added.



Note

See the table *Reference Curve Description Fields* for descriptions of the fields on this screen.

- 4. Complete the reference curve description information in the upper section. Use the drop-down lists where available. Use the tab key to move between fields.
- 5. Select **X**, **Y** Labels.
- 6. Expand the label selection trees as required.
- 7. Select one label in both the **X** Label and **Y** Label section.
- 8. Select OK.
- 9. Select **Input** to add a new set of points to the reference curve.

- On the *Input Data* dialog box, complete the information fields, S.D. (standard deviation), and L (skew value for the point) fields and select OK.
 The point set appears in the lower section sorted by the X-axis selection.
- 11. Repeat Step 10 as required to add additional point sets. Or select **Cancel** to close the Input Record dialog box and continue.
- 12. If necessary, edit a point set by clicking on the desired point set and selecting the **Edit** button to display the *Edit Data* dialog box.
- 13. Change the information fields as required and select **OK**.

The point set appears in the lower section sorted by the X-axis selection.

14. If necessary, delete a point set by clicking on the desired point set and selecting the **Delete** button.



Note

You are about to delete the selected record! Are you sure you want to continue?

- 15. Select **Yes** to continue. *The point set is removed from the lower section*.
- 16. Repeat Steps 14 and 15 as required to delete additional point sets.
- 17. When you have completed adding the reference curve, select **Close** to record the curve data. Select **OK** to return to the *Reference Curve Editor* dialog
- 18. Select **Close** to return to the main screen.

Field	Description
Sex	Select from drop-down list.
Ethnicity	Select from drop-down list.
Date	Set by the system when a curve is created or modified. Cannot be edited.
Author	Identifier for person creating or modifying curve. Enter up to five characters.
Source	Identifier for supplier providing the reference curve data. Enter up to 61 characters.
Comment	Comments pertaining to the reference curve.
Select X, Y Labels	This button brings up the Select X, Y Labels window.
X-Axis	
Label	X-axis label to display on reports.

Table 32: Reference Curve Description Fields

Field	Description
Display from - to	Range of X-axis data to display on reports between which the reference curve is considered to be valid. This does not necessarily correspond to the low and high points that define the curve.
Y-Axis	
Label	Y-axis label to display on reports.
Display from - to	Range used for the Y-axis in the graphical display. Does not affect the operation of Normals.
Is this curve current?	Lets more than one reference curve (for the same sex, ethnic group, scan type, and bone region) on the system at the same time. Only one of these curves can be marked as current. Only current curves are used by Normals.
Method	Analysis method for the curve. Select from drop-down list.
Scan Type	Set by the system depending on the X, Y label selection when a curve is created or modified. Cannot be edited.
Age Peak BMD	Age of maximum bone density used to compute T-score. Visible when X-axis label is selected as "Age" and Y-axis label is selected as "BMD".

Table 32: Reference Curve Description Fields

28.4 Copying a Reference Curve Record

Copying an existing reference curve record makes it easy to create a new curve record where most of the data is to be the same as that of the existing record.

- 1. Start the Reference Curve Editor.
- 2. Locate and select on the reference curve record line to copy.
- 3. Select **Copy**. A curve is added to the database.
- 4. Select OK.

The upper section of the *Copy Reference Curve* dialog box contains fields to enter or change the reference curve description information. The lower section lists the current reference curve point data.

- 5. Change the reference curve description information in the upper section as required. Use the drop-down lists where available. Use the tab key to move between fields.
- Please refer to <u>Creating New Reference Curve Records</u> on page 144 Steps 5 through 8 – for selecting X, Y labels.

- Please refer to <u>Creating New Reference Curve Records</u> on page 144 Steps 9 through 16 for adding, editing, and/or deleting reference curve point data. Then continue with the following steps.
- 8. When you have completed changing the copied reference curve, select **Close** to record the curve data.
- 9. Select **OK** to return to the *Reference Curve Editor* dialog box.
- 10. Select **Close** to return to the main screen.

28.5 Editing Reference Curve Records

Note

With the exception of the *Is the curve current*? field in the reference curve description section, Hologic-provided reference curve records cannot be modified.

- 1. Start the Reference Curve Editor.
- 2. Locate and select the reference curve record line to edit. *The line is highlighted*.
- 3. Select Edit.

The upper section of the *Edit Reference Curve* dialog box contains fields to enter or change the reference curve description information. The lower section lists the current reference curve point data.



Note

See the table in <u>Creating New Reference Curve Records</u> on page 144 for descriptions of the fields on this screen.

- 4. Enter or modify the reference curve description information in the upper section. Use the drop-down lists where available. Use the tab key to move between fields.
- 5. Please refer to <u>*Creating New Reference Curve Records*</u> on page 144, Steps 5 through 8 for selecting X, Y labels.
- 6. Please refer to <u>*Creating New Reference Curve Records*</u> on page 144, Steps 9 through 16 for adding, editing, and/or deleting reference curve point data. Then continue with the following steps.
- 7. When you have completed changing the copied reference curve, select **Close** to record the curve data.
- 8. Select **OK** to return to the *Reference Curve Editor* dialog box.
- 9. Select **Close** to return to the main screen.

28.6 Deleting Reference Curve Records

Note

Hologic-provided reference curve records cannot be deleted.

- 1. Start the Reference Curve Editor.
- 2. Locate and select on the reference curve record line to edit.
- 3. Select **Delete**.

Note



You are about to delete the selected record. This data and all results will be permanently LOST! Are you sure you want to continue?

- 4. Select **Yes** to delete the selected record and return to the *Reference Curve Editor* dialog box.
- 5. Repeat Steps 2 through 4 to delete additional records, or select **Close** to return to the main screen.

28.7 Adding New Ethnic Groups

 Select Utilities > Reference Curve > Add Ethnicity from the Menu Bar of the main window.



Note

If you plan to exchange data with other users, make sure your new ethnicity code does NOT match any of their ethnicity codes unless you are actually using the same reference curves for that ethnicity

- 2. Select **OK** to display the *Add New Ethnicity* dialog box.
- 3. Type name and code (two alphanumeric characters) for the new ethnicity group into their respective fields and select **OK** to add the group and return to the **Main Screen**.

28.8 Restoring Reference Curve Database



Caution

This option restores the reference curve database to the original state as supplied by Hologic. Any changes that may have been made are lost.

1. Select **Utilities > Reference Curve > Restore** from the Menu Bar of the main window.



This action will restore the reference curve database to the original state as supplied by Hologic, Inc. Any changes that may have been made will be lost. Proceed with Restore?

2. Select **Yes** to restore the database. Or select **No** to stop the restore. The system returns to the **Main Screen**.

Chapter 29 DICOM Option

Digital Imaging and Communications in Medicine (DICOM) is a powerful tool that provides:

- Interpreting physicians with the ability to view electronic QDR bone density scan and analysis results on a Picture Archiving and Communications System (PACS) viewer. The DICOM option allows results to be transmitted automatically over a facility's network directly to a physician's DICOM viewing station for interpretation and report dictation. The results can also be archived on the PACS, making them available for future reference and for distribution to others on the PACS network.
- The QDR system with the ability to retrieve schedule and patient demographic information when the Modality Worklist option is installed on the system.
- Locating and retrieving of scans that have been previously archived to a remote storage system (PACS) when the Query/Retrieve option is installed on the system.

29.1 Configure DICOM Option

29.1.1 DICOM Configuration Tabs

The following sections describe how to configure the Modality Worklist; add, edit, and delete DICOM Report Send remote destinations; add, edit, and delete DICOM Report Storage Commitment remote destinations; add, edit, and delete Query/Retrieve remote destinations; and configure the Host Machine (local system).

Settings for DICOM functions are controlled using the **System Configuration – DICOM** tab found under the **Utilities** pull-down menu in the main window.

The DICOM tab contains five tabs used to configure:

- Modality Worklist option (when installed)
- DICOM report send destinations (when installed)
- DICOM report storage commitment destinations (when installed)
- Query/Retrieve option (when installed)
- Host machine

29.1.2 Modality Worklist

The Modality Worklist option adds two tabs to the APEX software:

- A Worklist tab is added to the System Configuration DICOM window to allow the configuration of the modality Worklist.
- A Worklist tab is added to the Select Patient for Exam window to allow the operator to receive schedules from the HIS/RIS to perform tasks on the QDR system.

Configuring Modality Worklist



Caution

Changing information that configures the modality Worklist can cause serious communication disruptions with the HIS/RIS. Only authorized personnel should change settings.

The Modality Worklist is configured by selecting **System Configuration** – **DICOM** tab – **Worklist** tab found under the **Utilities** pull down menu in the main window.

The Worklist tab is divided into seven areas used to control communication to and from the HIS/RIS and one area providing an option for obtaining Worklist data from an input file rather than from a remote Worklist provider.

- Query Parameters
- Auto Query Interval
- Query Retry Parameters
- Purge Interval
- Input from File
- Configure Worklist Provider
- Map Worklist Fields
- Local Ping (confirm network and PACS connection)

Each of these areas is explained in the sections that follow.

29.1.3 Query Parameters

The user controls in this area perform the following:

- Filter the query by modality and AE Title.
- Determine a limit of entries to the Worklist database over a specified period.
- Determine if **Detailed Query** and **Extended Details** buttons will appear in the *Select a Patient for this Exam Worklist* dialog.

Table	33:	Query	y Parameters	

Parameter	Function
Days Back and Forward	This provides the valid time range of the query. There is a pair of drop- down menus labeled Back and Forward . If both are set to 0 (zero) then the valid range is for today's date only. If the Back drop-down menu is set to 7 and the Forward drop-down menu is set to 0 (zero) then the valid range of the query is one week, starting with today's date going back seven (7) days. The range for the drop-down menu for Back is 0 to 9 and the range for the drop-down menu for Forward is 0 to 8. Both Back and Forward drop-down menus can manually be set from 0 to 99.

Parameter	Function
Modality	Modality is the type of system recognized by the HIS/RIS. The default modality for the QDR System is "OT."
AE Title	AE stands for Application Entity. This is a text entry box that provides a unique name for the QDR system. Every QDR system will (or should) have an AE name that uniquely identifies that specific system.
Maximum Hits Per Query	This is an entry box that accepts a numeric value only. The number is the maximum number of query hits that will be passed to the QDR system specified in the Days Back and Forward . If there are more hits than the maximum, only the number specified in this box are passed to the QDR system.
Enable Detailed Query	A check box determining if a Detailed Query button and an Extended Details button will appear in the <i>Select a Patient for this Exam - Worklist</i> dialog when performing an exam. When checked both buttons will appear.

Table 33: Query Parameters

29.1.4 Auto Query Interval

Controls in this area are used set up a specific time interval during which the QDR system queries the provider to update the Worklist.

Auto Query Interval area contains three radio buttons, only one of which may be selected.



Note

These controls remain enabled when Input From file is enabled

Table 34: Auto	Query	Interval
----------------	-------	----------

Parameter	Function
Every Day At	This entry provides a control that allows the user to select the specific time each day that the QDR system will query the provider to update the Worklist.
Every	This entry provides two drop-down menus labeled HR and Min that specify a query to be made at the chosen time intervals (every n hours <i>a</i> nd <i>n</i> minutes).
Never	If Never is selected, the QDR system will not automatically query the provider to update the Worklist. With this selected, queries must be done manually by the operator.

29.1.5 Query Retry Parameters

If the provider did not respond to a query to update the Worklist for some reason (e.g., busy, off-line), controls in this area will determine how long the QDR system will wait for a response and give a specific time before trying the query again.

This entry contains a checkbox and three drop-down menus that control how long the QDR system will wait for the provider to respond to a query.



Note These controls remain enabled when Input From file is enabled

Parameter	Function
Query Retry Checkbox	In order for the QDR system to perform a retry after a time-out period, they must be a check mark in this box. If there is no check mark in this box then the QDR system will continue to wait until the provider responds to the query to update the Worklist. To place a check mark in the box, click in the box.
Query Timeout	This is a drop-down menu labeled Min. The value in this drop-down menu tells the QDR system how long it can wait before retrying the query. Min has a range of 0 to 60 minutes in 5 minute intervals. The operator can enter a number from $0 - 99$ manually.
Retry Number	This is a drop-down menu labeled Times that tells the QDR system how many times it may issue a retry. This drop-down menu allows a selection of 0, 1, 2, 3, 4, 5, 6, 7, 8, 9 or 10 for the number of retries. The operator can enter a number from 0 – 99 manually.
Retry Interval	This is a drop-down menu labeled Min that tells the QDR system how long to wait between retries.This drop-down menu has a range of 10 to 90 minutes in 10 minute intervals. The operator can enter a number from 0- 99 manually.

29.1.6 Purge Interval

Each time the provider responds to a query from the QDR system the Worklist entries are stored in a database on the QDR system. Use controls in this area to allow the database to be automatically purged (data removed) at a specific time.

These controls remain enabled when Input From file is enabled

Parameter	Function
Used Entries	This is a drop-down menu labeled Days . The value in this drop-down menu provides a limit for storing those studies that have already been performed. The studies are purged (deleted) after the specified number of days. The range for Used Entries drop-down menu is 0 to 9 days. The Used Entries drop-down menu can be set manually to any number of days from 0 to 999.
Unused Entries	This is a drop-down menu labeled Days . The value in this drop-down menu provides a limit for storing those studies that have not been performed. These are purged (deleted) after the specified number of days. The range for the drop-down menu for Unused Entries is 0 to 9 days. The Unused Entries drop-down menu can be set manually to any number of days from 0 to 999.

Table 36: Purge Interval

29.1.7 Input From File

Provides an option for obtaining Worklist data from an input file generated by an electronic Medical Reporting System rather than from a remote Worklist provider.

Parameter	Function
Enable	A check box determining whether or not the Input from File feature is enabled. When checked, Worklist data is obtained from an input file.
	All controls on the Worklist tab not used for the Input from File feature are disabled when this box is checked.
Input File Name	Displays a full path to the current Worklist file. This field is populated or changed using Browse to select a file path.
(Browse)	Displays a "File Open" dialog allowing the user to locate a Worklist input file on the local system or network.

Table 37: Input From File

29.1.8 Worklist Provider

Worklist Provider Interface

The Worklist provider interface provides Worklist entries for the QDR system.

Refer to Worklist Provider Controls on page 154 for details.

Parameter	Function
Map Worklist Fields	Different hospitals and clinics may use the same DICOM attributes in various ways to identify their patients. Map Worklist Fields is used to ensure that data in the QDR system and in the HIS/RIS are addressing the same patient.
	Select Map Worklist Fields , on the Worklist Tab, to display a window called Map Worklist Keys that allows fields to be selected for patient verification by the QDR system and the HIS/RIS database. Once these entries are made, they will be checked on each study to verify that the QDR system and the HIS/RIS have identified the same patient. Refer to <u>Map Worklist Fields</u> on page 156 for details.
Local Ping	Confirms if the local system is connected to a network.

Table 38: Worklist Provider Features

Worklist Provider Controls

Selecting **Configure Worklist Provider** displays a window used to define the Worklist provider.

The Worklist provider supplies Worklist entries for the QDR system.

Worklist Provider

Use controls in this area to define the Worklist provider.

Parameter	Function
AE Title	This is the Application Entity title of the Worklist provider.
Remote Host	This is the host name or IP address of the Worklist provider. The host name or IP address must be on the same network as the QDR workstation. Accepts up to 120 alphanumeric characters.
Remote Port Number	This is the port number of the Worklist provider.

Table	20.	Worklist	Dromida

Performed Procedure Step

This area provides an option to automatically respond to a performed procedure provider once a specific task is performed in the Worklist.

Control	Function
Use Performed Procedure Step	When selected, each time a study is completed a response is sent to the provider indicating that the task has been completed. To place a check mark in the box, click in the box.

Provider

Use controls in this area to define the performed procedure step provider. The performed procedure provider may be the same as the Worklist provider or different.

Table 41: Provider			
Control	Function		
Use Worklist Provider Data	When selected, the performed procedure step provider is the same as the Worklist provider.		
AE Title	This is the Application Entity title of the performed procedure step provider when <i>Use Worklist Provider Data</i> is unchecked.		
Remote Host	This is the host name or IP address of the performed procedure step provider when <i>Use Worklist Provider Data</i> is unchecked.		
	The host name or IP address must be on the same network as the QDR workstation.		
	Accepts up to 120 alphanumeric characters.		
Remote Port Number	This is the port number of the performed procedure step provider when <i>Use Worklist Provider Data</i> is unchecked.		
Remote Ping	Confirms if the configured Worklist provider or Performed Procedure Step provider system is connected to the same network as the local system.		
С-ЕСНО	Confirms if the configured Worklist provider or Performed Procedure Step provider system is a Picture Archival and Communications system (PACS)		

29.1.9 Map Worklist Fields

Selecting Map Worklist Fields displays the Map Worklist Keys dialog box.



Caution

Do not change settings in the Map Worklist Fields without specific directions from your HIS/RIS department.

This dialog box consists of 15 different drop-down menus with specific labels on the left. The labels indicate data that is in the Patient Biography on the QDR systems (some information, such as HL7 fields, may not apply to each QDR system). The information in the drop-down menus is information that may appear in the HIS/RIS database for the patient. The goal is to find information in the Patient Biography that matches information in the HIS/RIS database to be used as a key to verify that the patient in the Patient Biography is the same as the patient in the HIS/RIS database.

This task of matching information must be performed by someone with knowledge of both the QDR system and the operations of the HIS/RIS.

There are two sets of drop-down menus:

- Patient Matching Keys six drop-down menus for Patient Matching
- Other Keys nine drop-down menus for Key Mapping

The **Patient Matching** drop-down menus allow specific data in the Patient Biography of the QDR system to be mapped to data in the HIS/RIS database. These drop-down menus have check boxes to the left of the label. A check mark in any of these boxes means that the information in the Patient Biography must match the information in the HIS/RIS database to identify the same patient.

The **Key Mapping** drop-down menus provide data from the HIS/RIS database that can be mapped to specific information in the Patient Biography.

Not Mapped in any drop-down menu indicates that the field in the Patient Biography will not be mapped with any field in the HIS/RIS database.

29.1.10 DICOM Send Destinations

Send destinations define destinations to which the DICOM reports are transmitted when the **Send** function is used.

This section describes how to configure, add, edit, and delete destination remote nodes and how to configure all Send destinations.

Send destinations are configured by selecting System Configuration — DICOM tab — Send tab found under the Utilities pull down menu in the main window.

Parameter	Function
Configure DICOM Send Destinations	Selection list of existing DICOM Send destinations configured on the local system
Add Destination	Displays a dialog that allows configuring a new Send destination
Edit Destination	<i>Enabled when one destination is selected.</i> Displays a dialog that allows changing the selected Send destination's configuration.
Delete Destination	<i>Enabled when one or more destinations are selected.</i> Deletes the selected Send destination(s)
Local Ping	Confirms whether the local system is connected to a network.
Remote Ping	<i>Enabled when one destination is selected.</i> Confirms whether the configured Send destination system is connected to the same network as the local system
C-ECHO	<i>Enabled when one destination is selected.</i> Confirms whether the configured Send destination system is a Picture Archival and Communications system (PACS).
Configure Parameters	Select Configure Parameters to define the Send destination auto retry parameters and when to purge log entries. Refer to <u>Configure DICOM Send</u> on page 161 for details.

Table 42: DICOM Send Destinations

- 1. To **add** a new destination:
 - a. Select Add Destination,
 - b. Complete the *Add / Edit DICOM Send Destination* dialog fields (refer to <u>Add / Edit</u> <u>DICOM Send Destination Dialog</u> on page 159 for details).
 - c. Select OK.

- 2. To **reconfigure** an existing destination:
 - a. Select the destination in the selection list.
 - b. Select Edit Destination,
 - c. Edit the *Add / Edit DICOM Send Destination* dialog box as required (refer to <u>Add /</u> <u>Edit DICOM Send Destination Dialog</u> on page 159 for details).
 - d. Select OK.
- 3. To **enable/disable** a destination for use:
 - a. Select the check box in the Active column of the selection list for the destination to check/uncheck (check to enable).
- 4. To **verify the local system is connected** to a local network:
 - a. Select Local Ping.
 - b. Select **OK** to close the resulting message.
- 5. To **verify the destination is connected** to the same network as the local system:
 - a. Select the destination in the selection list
 - b. Select **Remote Ping**. Select **OK** to close the resulting message.
- 6. To verify the destination is a PACS:
 - a. Select the destination in the selection list.
 - b. Select C-ECHO.
 - c. Select **OK** to close the resulting message.
- 7. To **delete** a destination:
 - a. Select the destination in the selection list.
 - b. Select **Delete Destination**.
- 8. To define auto query parameters for *all* configured Send destinations:
 - a. Select Configure Parameters.
 - b. Edit the **Configure DICOM Send** dialog box as required (refer to <u>Configure</u> <u>DICOM Send</u> on page 161 for details).
 - c. Select OK.
- 9. Select **OK** to return to the **Main Screen**.

Parameter	Function
AE Title	<i>Application Entity.</i> Provides a unique name for the destination system. Accepts up to 16 alphanumeric characters. Note: To add the QDR workstation as a destination node, type local in the AE Title box.
Host Name or IP Address	Name or IP address of destination. The host name or IP address must be on the same network as the QDR workstation. Accepts up to 120 alphanumeric characters. Note To add the QDR workstation as a destination node, type localhost in the Host Name or IP Address box
SCP Port	Port number on destination. The default for the Service Class Provider Port number is 104. Accepts up to 5 numeric characters in the range of 1 - 65535.
Destination Name	Provides an alias name used for selecting a destination. Accepts up to 120 alphanumeric characters.
Interpreting Physician	An optional entry that provides the name of the physician interpreting the scan. Accepts up to 120 characters.
Grayscale Only	When checked, DICOM reports are converted into grayscale when sending to the selected destination. When unchecked, images for all report types that may contain graphs will be created in color. Images for all other report types (IVA, questionnaire, etc) will be created in grayscale.
Presentation File	When checked, a DICOM GSPS file will be sent along with an image DICOM file if needed for IVA scan. If checked and the node is also marked for storage commitment, then the commitment request will be issued for both image and GSPS DICOM files.

Table 43: DICOM Send Destinations

Add / Edit DICOM Send Destination Dialog

Parameter	Function
IVA Results File	When checked, a DICOM IVA Results file will be sent along with an image DICOM file if needed for IVA scan.
	If checked and the node is also marked for storage commitment, then the commitment request will be issued for both image and IVA Results DICOM files.
Unicode	When checked, DICOM files are sent with Unicode coding.
	When using Unicode, an Extended Character Set attribute will be present in DICOM files. Text attributes will be populated with single byte strings converted from double byte strings using UTF-8 encoding.
	When not using Unicode, an Extended Character Set attribute will not be present in DICOM files. Text attributes will be populated with single byte strings converted from double byte strings using UTF-8 encoding.
Storage Commitment Provider	When checked, designates the destination is committed to storing information sent to it.
	When checked a storage commitment provider must be designated. To designate a storage commitment provider, select an existing provider from the drop-down list, or click Add New Provider to designate add a new storage commitment provider (refer to <u>Add / Edit DICOM</u> <u>Query/Retrieve Destination Dialog</u> on page 168 for details on adding a new provider). Storage commitment providers can also be added using the System Configuration - DICOM tab - Commit tab (refer to <u>DICOM</u> <u>Query/Retrieve Destinations</u> on page 166 for details).
Scan Archive Location	Displays if the Enterprise Data Management option is installed. When checked, designates the destination is a scan archiving location
	and will implement the inclusion of P&R files to the DICOM file. P files are QDR scan files that contain processing data for the scan.
	R files are QDR scan files that contain the raw data for the image.
	When checked, the Select Existing Provider field can be left blank to indicate the Send destination is also the scan archive location. Or, an existing scan archiving location can be selected from the drop-down list, or click Add New Provider to designate add a new scan archive location. Scan archive locations can also be added using the System Configuration - DICOM tab - Query/Retrieve tab (refer to <u>DICOM Query/Retrieve</u> <u>Destinations</u> on page 166 for details).
ОК	Validates data. If validation passes, the dialog is closed and the configure DICOM Send Destinations selection list is updated with new or changed data. If validation fails, a warning message displays.
Cancel	Ignores all edits and closes the dialog.

Table 43: DICOM Send Destinations

Table 44: DICOM Send Configuration			
Parameter	Function		
Auto Retry Parameters	If the Send destination did not respond to a request for some reason (e.g., busy, off-line), controls in this area will determine how long the QDR system will wait for a response and give a specific time before trying the query again.		
Retry Number	This is a entry box labeled Times that tells the QDR system how many times it may issue a retry. The Retry Number entry box can be set manually to any number of days from 0 to 99.		
Retry Interval	This is a entry box labeled Min that tells the QDR system how long to wait between retries. The Retry Interval entry box can be set manually to any number of minutes from 1 to 1440.		
Purge log entries after	This is a entry box labeled days . The value in this box provides a limit for storing DICOM Send log entries. The entries are purged (deleted) after the specified number of days. The Purge log entries after entry box can be set manually to any number of days from 0 to 99.		
Auto-accession number	When checked, an accession number is automatically generated in the format SSSSSSYYMMDDNNN, where SSSSSS is the QDR serial number, YYMMDD is the current date, and NNN is a number starting at 001 and going up to 999.		
Study Description	The contents of this edit box is used to populate the Study Description field in DICOM files if:		
	• the study is not a Worklist study, or the study is a Worklist study but the Study Description field is not mapped to any Worklist attribute		
	and,the edit box has a text entry (if left blank, the Study Description attribute is omitted)		

Configure DICOM Send

29.1.11 DICOM Storage Commitment Destinations

Storage commitment destinations define destinations to which the DICOM reports can be transmitted and stored when the **Send** function is used.

Storage Commitment destinations must be defined before defining DICOM Send destinations as storage commitment destinations.

This section describes how to configure, add, edit, and delete storage commitment destination remote nodes and how to configure all Storage Commitment destinations.

Send destinations are configured by selecting **System Configuration – DICOM** tab – **Commit** tab found under the **Utilities** pull down menu in the main window.

Configure DICOM Send Destinations

Selection list of existing DICOM Storage Commitment destinations configured on the local system.

Parameter	Function
Add Destination	Displays a dialog that allows configuring a new Storage Commitment destination
Edit Destination	Enabled when one destination is selected.
	Displays a dialog that allows changing the selected Storage Commitment destination's configuration.
Delete Destination	Enabled when one or more destinations are selected.
	Deletes the selected Storage Commitment destination(s).
Local Ping	Confirms whether the local system is connected to a network.
Remote Ping	Enabled when one destination is selected.
	Confirms whether the configured Storage Commitment destination is connected to the same network as the local system.
C-ECHO	Enabled when one destination is selected.
	Confirms whether the configured Storage Commitment destination is a Picture Archival and Communications system (PACS).
Configure Parameters	Select Configure Parameters to define the Storage Commitment destination auto retry parameters, when to purge log entries, and how DICOM storage commitment files are sent to a destination. Refer to <u>Configure Storage Commitment</u> on page 165 for details.

Table 45: DICOM Storage Commitment Destinations

- 1. To **add** a new destination:
 - a. Select Add Destination.
 - b. Complete the *Add / Edit Storage Commitment* dialog fields (refer to <u>Add / Edit</u> <u>Storage Commitment Destination Dialog</u> on page 164 for details).
 - c. Select OK.
- 2. To reconfigure an existing destination:
 - a. Select the destination in the selection list.
 - b. Select Edit Destination
 - c. Edit the *Add / Edit Storage Commitment* dialog box as required (refer to <u>Add / Edit</u> <u>Storage Commitment Destination Dialog</u> on page 164 for details).
 - d. Select OK.
- 3. To enable/disable a destination for use:
 - a. Select the check box in the Active column of the selection list for the destination to check/uncheck (check to enable).
- 4. To **verify the local system is connected** to a local network:
 - a. Select Local Ping.
 - b. Select **OK** to close the resulting message.
- 5. To **verify the destination is connected** to the same network as the local system:
 - a. Select the destination in the selection list.
 - b. Select Remote Ping.
 - c. Select **OK** to close the resulting message.
- 6. To **verify** the destination is a PACS:
 - a. Select the destination in the selection list.
 - b. Select C-ECHO.
 - c. Select **OK** to close the resulting message.
- 7. To **delete** a destination:
 - a. Select the destination in the selection list.
 - b. Select Delete Destination.
- 8. To **define auto query parameters** for *all* configured Storage Commitment destinations:
 - a. Select Configure Parameters.
 - b. Edit the *Configure Storage Commitment* dialog box as required (refer to <u>*Configure*</u> <u>Storage Commitment</u> on page 165 for details).
 - c. Select **OK**.
- 9. Select **OK** to return to the **Main Screen**.

	Table 46: Storage Commitment Destination Editing		
Parameter	Function		
AE Title	Application Entity Provides a unique name for the destination system. Accepts up to 16 alphanumeric characters.		
	Note To add the QDR workstation as a destination node, type local in the AE Title box		
Host Name or IP	Name or IP address of destination.		
Address	The host name or IP address must be on the same network as the QDR workstation.		
	Accepts up to 120 alphanumeric characters.		
	Note To add the QDR workstation as a destination node, type localhost in the Host Name or IP Address box.		
SCP Port	Port number on destination.		
	The default for the Service Class Provider Port number is 104. Accepts up to 5 numeric characters in the range of 1 - 65535.		
Destination Name	Provides an alias name used for selecting destinations. Accepts up to 120 alphanumeric characters.		
ОК	Validates data. If validation passes, the dialog is closed and the configure DICOM Storage Commitment Destinations selection list is updated with new or changed data. If validation fails, a warning message displays.		
Cancel	Ignores all edits and closes the dialog.		

Add / Edit Storage Commitment Destination Dialog

Configure Storage Commitment

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1 uble 47: 5	storage Con	nmitment (Configuration	

Parameter	Function	
Auto Retry Parameters	If the Storage Commitment destination did not respond to a request for some reason (e.g., busy, off-line), controls in this area will determine how long the QDR system will wait for a response and give a specific time before trying the query again.	
Retry Number	This entry box labeled Times tells the QDR system how many times it may issue a retry. The Retry Number entry box can be set manually to any number of days from 0 to 99.	
Retry Interval	This entry box labeled Min tells the QDR system how long to wait between retries. The Retry Interval entry box can be set manually to any number of minutes from 1 to 1440.	
Mode	DICOM files can be sent to a storage commitment destination as one file or all files per request.	
Image-by-image	This radio button, when selected, tells the QDR system to issue a single storage commitment request for all DICOM files to be sent.	
Batch	This radio button, when selected, tells the QDR system to issue a storage commitment request for each DICOM file to be sent.	
Purge log entries after	This is a entry box labeled days . The value in this box provides a limit for storing DICOM Send log entries. The entries are purged (deleted) after the specified number of days. The Purge log entries after entry box can be set manually to any number of days from 0 to 99.	

29.1.12 DICOM Query/Retrieve Destinations

Query/Retrieve allows the operator to query a remote location (PACS) for scans meeting given parameters and filters and to retrieve selective scans into the current computer. The scans must have been stored to that remote location prior to using the Query/Retrieve function.

This section describes how to configure, add, edit, and delete a destination remote node.

Query/Retrieve destinations are configured by selecting **System Configuration** – **DICOM** tab – **Query/Retrieve** tab found under the **Utilities** pull down menu in the main window.

Parameter	Function	
Configure DICOM Query/Retrieve Destinations	Selection list of existing DICOM Query/Retrieve destinations configured on the QDR system.	
Add Destination	Displays a dialog that allows configuring a new Query/Retrieve destination	
Edit Destination	Enabled when one destination is selected. <i>Displays a dialog that allows changing the</i> selected Query/Retrieve destination's configuration.	
Delete Destination	<i>Enabled when one or more destinations are selected.</i> Deletes the selected Query/Retrieve destination(s).	
Local Ping	Confirms whether the local system is connected to a network.	
Remote Ping	<i>Enabled when one destination is selected.</i> Confirms whether the configured Query/Retrieve system is connected to the same network as the local system.	
C-ECHO	<i>Enabled when one destination is selected.</i> Confirms whether the configured Query/Retrieve system is a Picture Archival and Communications system (PACS).	

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Table 48:	DICOM	Query/Retrieve	Destinations

- 1. To **add** a new destination:
 - a. Select Add Destination.
 - b. Complete the Add DICOM Query/Retrieve Destination dialog fields.
 - c. Select OK.

- 2. To **reconfigure** an existing destination:
 - a. Select the destination in the selection list.
 - b. Select Edit Destination.
 - c. Edit the Edit DICOM Query/Retrieve Destination dialog box as required.
 - d. Select OK.
- 3. To **enable/disable** a destination for use:
 - a. Select the check box in the Active column of the selection list for the destination to check/uncheck (select to enable).
- 4. To **verify the local system is connected** to a local network:
 - a. Select Local Ping.
 - b. Select **OK** to close the resulting message.
- 5. To **verify the destination is connected** to the same network as the local system:
 - a. Select the destination in the selection list.
 - b. Select Remote Ping.
 - c. Select **OK** to close the resulting message.
- 6. To **verify** the destination is a PACS:
 - a. Select the destination in the selection list.
 - b. Select C-ECHO.
 - c. Select **OK** to close the resulting message.
- 7. To **delete** a destination:
 - a. Select the destination in the selection list.
 - b. Select **Delete Destination**.
- 8. To configure **Query/Retrieve parameters:**
 - a. Select Configure Parameters. (Configure Query/Retrieve on page 168.
- 9. Select **OK** to return to the **Main Screen**.

Table 49: Add / Edit DICOM Query/Retrieve Destination		
Parameter	Function	
AE Title	Application Entity. Provides a unique name for the destination system.	
	Accepts up to 16 alphanumeric characters.	
Host Name or IP	Name or IP address of destination.	
Address	The host name or IP address must be on the same network as the QDR workstation.	
	Accepts up to 120 alphanumeric characters.	
SCP Port	Port number on destination.	
	The default for the Service Class Provider Port number is 104.	
	Accepts up to 5 numeric characters in the range of 1 - 65536.	
ОК	Validates data. If validation passes, the dialog is closed and the Configure DICOM Query/Retrieve Destinations selection list is updated with new or changed data. If validation fails, a warning message displays.	
Cancel	Ignores all edits and closes the dialog.	

Add / Edit DICOM Query/Retrieve Destination Dialog

Configure Query/Retrieve

Table 50: Query/Retrieve Configuration

Parameter	Function
Locate by Study	Select to perform retrieve operations on the study level, i.e. the entire set of scans belonging to the same DICOM study as the selected scan will be retrieved.
Locate by Series	Select to perform a retrieve operation on the series level, i.e. a single scan corresponding to the given DICOM series will be retrieved.

29.1.13 Host

Host Machine defines your system when DICOM functions are used. The Host Machine is configured by selecting **System Configuration – DICOM** tab **– Host** tab found under the **Utilities** pull down menu in the main window.

Important: If changes are made to either the AE Title or Listening Port data, the APEX application must be restarted for the changes to take effect

Parameter	Function
Host DICOM Configuration AE Title	Used to configure the Host.
	Application Entity Provides a unique name for the QDR system. Accepts up to 16 alphanumeric characters.
Station Name	Name of the QDR system. Accepts up to 120 alphanumeric characters.
Listening Port	Port number on which the QDR system listens. The default port number is 104. Accepts up to 5 numeric characters in the range of 1 – 65536.
DICOM Send Modality	Defines the value used to populate the Modality field in DICOM files for DICOM Send and Save operations.
Modality	This edit box defines the value entered into the Modality field in DICOM files. The default value is "OT" or the last saved value.
Worklist sets modality	When checked: If there is an entry in the Worklist database for a scan being saved or sent, then modality from the Worklist is used. If there is no entry in the Worklist database (either because the scan was not initiated using the Worklist, or because the Worklist entry has been already deleted), then the value from the Modality edit box is used.
ОК	Validates data. If validation passes, the System Configuration window closes returning to the main screen. If validation fails, a warning message displays.
Cancel	Ignores all edits, closes the System Configuration window, and returns to the main screen.

Table 51: Host Configuration

Chapter 30 FRAX FAQs

In clinical practice I would prefer to use one year probabilities — why use 10-year fracture probability?

In young healthy individuals (with a low mortality) the one year probability is approximately 10% of the 10-year probability. Thus, an individual with a 10-year fracture probability of 40% would have approximately a 1-year probability of 4%. Higher percentage figures are more readily understood by patients and clinicians.

The clinical risk factors demand a yes or no response. However, two prior clinical fractures carry a greater risk than a single previous fracture. Why is this not accommodated?

It is known that dose-responses exist for many of the clinical risk factors. In addition to the number of previous fractures, they include smoking, use of glucocorticoids and consumption of alcohol. The model is, however, based on information that is common to all the cohorts that participated in its creation and such detail is not available. This means that clinical judgment needs to be used when interpreting probabilities. A higher than average dose of glucocorticoids will carry a higher probability than that displayed. Conversely, a lower than average dose will signify a lower probability.

A prior vertebral fracture carries a higher risk than a prior forearm fracture. How is this accounted for in the algorithms?

It is not, for the reasons indicated in the question above. It should be noted, however, that a prior morphometric and asymptomatic vertebral fracture carries approximately the same risk as any previous fracture. A clinical vertebral fracture, however, carries a much higher risk (see reference list, Johnell et al 2006).

How is account taken of ethnic minorities?

It is not - with the exception of the United States where there is sufficient epidemiological information to make the appropriate adjustments.

Why can't I use the tool to predict fracture risk in a 30 year old patient?

The model is constructed from real data in population-based cohorts around the world that have a limited age range. If you enter an age below 40 years, the tool will calculate the probability of fracture at the age of 40 years. You must use your clinical judgment to interpret the risk.

For the clinical risk factors, there is no provision for missing values (i.e., a "do not know" category) in the program. What should I do?

Missing values are not provided for in our program. When calculating the 10-year probability it is assumed that every question (except BMD) can be answered. If you don't have information, for example on family history, you should answer no.

Why not report the probability of all osteoporotic fractures? This would give larger values.

Incorporating all osteoporotic fractures is problematic because of limited information on their epidemiology. From Swedish data, the inclusion of other major osteoporotic fractures (e.g. pelvis, other femoral fractures and tibial fractures) would increase the values by about 10 % (for example, in a patient with a calculated probability of major osteoporotic fractures of 5%, this might be uplifted to 5.5%). Including rib fractures would have a much larger effect. They are, however, difficult to diagnose.

Why not include falls which are a well established clinical risk factor for fracture?

Two reasons. The first is that the cohort data used to create the model reported falls in very different ways so that it was not possible to derive a standardized metric. Second, although plausible, pharmaceutical intervention has not been shown to reduce fracture risk in patients selected on the basis of a fall history. It is important that risk assessment models identify a risk that can be reduced by treatment.

Why have you ignored fractures diagnosed on X-ray and focused on clinical vertebral fracture?

A prior morphometric fracture has the same significance as any other prior fragility fracture and can be entered into the FRAX[®] model. The output does not, however, include the probability of a morphometric fracture. This is a conservative position, since their clinical significance is controversial (other than for risk prediction). Nevertheless, this does not affect who would be eligible for treatment.

How do I decide who to treat?

The FRAX assessment does not tell you who to treat which remains a matter of clinical judgment. In many countries, guidelines are provided that are based on expert opinion and/or on health economic grounds.

Additional information can be found at the WHO web site (<u>www.who.int</u>).

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