OmniTom Elite with PCD

User Manual

1-NL5100-060 Revision 01



1.1 Contents

Photon Counting Detector Overview	
Intended use of the system	
Clinical benefit	22
Consumer information	22
Proprietary rights	
Legal disclaimer	
Contact information	
NeuroLogica Corporation	
Damage in transportation	
User requirements	
Essential performance	25
About this user manual	25
Identified symbols and system classifications	25
Conventions used in this user manual	
Understanding the use of "you" in this user manual	
Active and inactive objects	
Use environment for the OmniTom Elite with PCD	
Chapter 1 Compliance and Safety Requirements	29
Chapter 1 Compliance and Safety Requirements	
Chapter 1 Compliance and Safety Requirements	29
Chapter 1 Compliance and Safety Requirements IEC classification and symbols Environmental specifications Site specification	29
Chapter 1 Compliance and Safety Requirements IEC classification and symbols Environmental specifications Site specification Hazardous substances	29
Chapter 1 Compliance and Safety Requirements	29 29 33 35 35 35 36
Chapter 1 Compliance and Safety Requirements	29
Chapter 1 Compliance and Safety Requirements	29 29 33 35 35 35 36 36 37 38
Chapter 1 Compliance and Safety Requirements	29 29 33 35 35 36 36 37 38 38
Chapter 1 Compliance and Safety Requirements	29 29 33 35 35 36 36 37 38 38 38
Chapter 1 Compliance and Safety Requirements IEC classification and symbols Environmental specifications Site specification Hazardous substances Part numbers and product-marking plates Class 1 Type B medical devices Focal spot Filtration Source to Detector Distance (SID) Compliance statement	29 29 33 35 35 36 37 38 38 38 38 38 38
Chapter 1 Compliance and Safety Requirements IEC classification and symbols Environmental specifications. Site specification. Hazardous substances Part numbers and product-marking plates. Class 1 Type B medical devices Focal spot Filtration Source to Detector Distance (SID) Compliance statement EMI/EMC terms	29 29 33 35 35 36 37 38 38 38 38 38 38 38 38 38 38 38 38 38
Chapter 1 Compliance and Safety Requirements IEC classification and symbols Environmental specifications Site specification Hazardous substances Part numbers and product-marking plates Class 1 Type B medical devices Focal spot Filtration Source to Detector Distance (SID) Compliance statement EMI/EMC terms Electromagnetic compatibility	29
Chapter 1 Compliance and Safety Requirements IEC classification and symbols Environmental specifications Site specification Hazardous substances Part numbers and product-marking plates Class 1 Type B medical devices Focal spot Focal spot Filtration Source to Detector Distance (SID) Compliance statement EMI/EMC terms Electromagnetic compatibility Radio frequency interference	29 29 33 35 35 36 37 38 38 38 38 38 38 38 39 39 40
Chapter 1 Compliance and Safety Requirements IEC classification and symbols Environmental specifications Site specification Hazardous substances Part numbers and product-marking plates Class 1 Type B medical devices Focal spot Focal spot Filtration Source to Detector Distance (SID) Compliance statement EMI/EMC terms Electromagnetic compatibility Radio frequency interference Susceptibility	29 29 33 35 35 36 37 38 38 38 38 38 38 38 39 39 40 40

Countermeasures against EMC related issues	44
Use recommendations	47
Installation recommendations	48
Cable shielding and grounding	48
Adjacent components and equipment	48
Static magnetic field limits	48
Electrostatic discharge environment and recommendations	49
Facility IT-NETWORK	49
Hazard Information	49
General safety considerations and statements	49
Laser safety	51
Scanner mobility safety	52
Electrical safety	53
Mechanical safety	54
Radiation safety	55
Fire and explosion safety	56
	ГC
EMERGENCY STOP button	
Battery system safety and information	
Battery system safety and information Battery component for the base power distribution assembly	
Battery system safety and information Battery component for the base power distribution assembly Maintenance and service	
EMERGENCY STOP button Battery system safety and information Battery component for the base power distribution assembly Maintenance and service Cleaning the system	
EMERGENCY STOP button Battery system safety and information Battery component for the base power distribution assembly Maintenance and service Cleaning the system Cybersecurity	
EMERGENCY STOP button Battery system safety and information Battery component for the base power distribution assembly Maintenance and service Cleaning the system Cybersecurity Contraindication(s)	
EMERGENCY STOP button	
EMERGENCY STOP button Battery system safety and information Battery component for the base power distribution assembly Maintenance and service Cleaning the system Cleaning the system Cybersecurity Contraindication(s) Personnel privileges and terminology Qualified operator Operator of record. Scanning privileges	
EMERGENCY STOP button Battery system safety and information Battery component for the base power distribution assembly Maintenance and service Cleaning the system Cybersecurity. Contraindication(s) Personnel privileges and terminology Qualified operator Operator of record. Scanning privileges Protocol privileges	
EMERGENCY STOP button Battery system safety and information Battery component for the base power distribution assembly Maintenance and service Cleaning the system. Cybersecurity. Contraindication(s) Personnel privileges and terminology Qualified operator Operator of record. Scanning privileges Protocol privileges Clinical operation.	
EMERGENCY STOP button	
EMERGENCY STOP button Battery system safety and information	

Chapter 2 System Overview	64
OmniTom Elite with PCD system	64
The LCD touch screen	65
The shielding curtains	65
The tablet	65
The silhouette scan board and universal transfer board	66
Parts that potentially come into contact with the patient	67
Chapter 3 Basic Scanner Operations	68
Powering on and off the OmniTom Elite with PCD system	68
Overview of the scanner's LCD touch screen	69
LCD tabs and icons	69
Drive bar calibration for transport	
Moving the scanner for transport	74
Using collision avoidance sensors	
Transporting with drive view	
Transporting with mood ring lights	80
Positioning the scanner before a scan	81
Positioning the patient	82
Using alignment camera	84
Positioning the scanner using the laser light	88
To activate the laser light on the touch screen	88
Using the curtains for shielding	88
Rebooting the system	89
Remote support feature	89
Enabling remote support	90
Chapter 4 Basic Tablet Operations	91
Understanding the types of users	91
Powering on the tablet	91
Logging in to the tablet	92
Locking the tablet with the LCD	93
Logging off the tablet	95
Powering off the tablet	96
Enabling the privacy screen on the tablet	

Navigating around the tablet's main screen	
Getting to know the scanner status screen	
System status orbs and mood ring lights	101
Getting to know the system screen	103
Getting to know the protocol screen	104
Getting to know the patient screen	105
The tablet tabs	106
Tablet buttons	106
Chapter 5 System Settings	109
System settings overview	109
Setting user accounts	110
Deleting a user	111
Applying dose configuration	112
Setting dose notifications	
Editing dose notifications on NL protocols	115
Setting dose alerts (system limits)	
Managing DICOM servers	
Setting up the audit trail viewer	119
Setting recon presets	121
Setting recon preset groups	122
Setting MPR presets	123
Setting windowing presets	
Selecting image orientation	125
Chapter 6 Protocol Manager	
Creating a new protocol	
Adding a New Protocol using Copy from an Existing Protocol	
Editing an existing protocol	135
Deleting a protocol	
Hiding protocols	139
Moving protocols	139
Chapter 7 Daily Calibration and Quality Assurance	141
Daily calibration	
Performing a daily calibration	

Tube seasoning	
The QA phantom overview	
Performing a quality assurance test	
Ensuring good image quality	
OmniTom Elite with PCD dose information (21 CFR 1020.33 c)	
CTDI ₁₀₀ measurements	
Note regarding detector collimation	
Calculation of CTDI ₁₀₀	
Sample calculation	
The scout dose	
Recommended dose verification procedure	
OmniTom Elite with PCD dose in air	
QA measurements	
Beam width	
Geometric efficiency in the Z axis direction	
Slice sensitivity	
Half-value layer	
Allowable variations	
Scatter radiation	
Scatter exposure	
Additional scatter measurements	
Dose linearity with tube voltage and current	
Chapter 8 Registration	
Navigating the Registration Screen	
Registering the patient	
Querying patient information	
Storing patients in the stored list	
Removing patients from the stored list	
Manually registering a patient using 'Manual Add'	
Viewing patient information	
Chapter 9 Patient Scanning	
Patient scanning overview	

Identifying protocol types	178
Axial	
Helical	179
Reference	
Scout	
Performing a PCD scan	
Re-using Scouts with a New Protocol	
Repeating an Image	
Scanning with special features	
Using the step & shoot option	
Performing a scan with Automatic Exposure Control (AEC)	
Adding Noise Reduction to Axial Reconstructions	
Performing a CT angiography scan with bolus tracking	
Performing test bolus	
Performing a CT perfusion scan	
Viewing images in the CTP viewer	
Examining the scanned image with tools	205
	205
Using tools on the acquisition tab	
Using tools on the acquisition tab	205
Chapter 10 Browser	
Chapter 10 Browser	
Chapter 10 Browser Browser overview Navigating the browser	
Chapter 10 Browser Browser overview Navigating the browser Selecting PCD Images in the Browser	
Chapter 10 Browser Browser overview Navigating the browser Selecting PCD Images in the Browser Compare Series	
Chapter 10 Browser Browser overview Navigating the browser Selecting PCD Images in the Browser Compare Series Loading a series to viewer from the browser	205 208 208 209 209 209 210 212
Chapter 10 Browser Browser overview Navigating the browser Selecting PCD Images in the Browser Compare Series Loading a series to viewer from the browser Deleting a series	205
Chapter 10 Browser Browser overview Navigating the browser Selecting PCD Images in the Browser Compare Series Loading a series to viewer from the browser Deleting a series Editing patient details	205 208 208 209 209 209 210 210 212 213 214
Chapter 10 Browser Browser overview Navigating the browser Selecting PCD Images in the Browser Compare Series Loading a series to viewer from the browser Deleting a series Editing patient details Archiving patient images	205
Chapter 10 Browser Browser overview Navigating the browser Selecting PCD Images in the Browser Compare Series Loading a series to viewer from the browser Deleting a series Editing patient details Archiving patient images Archiving to PACS	205
Chapter 10 Browser Browser overview Navigating the browser Selecting PCD Images in the Browser Compare Series Loading a series to viewer from the browser Deleting a series Editing patient details Archiving patient images Archiving to PACS	205
Chapter 10 Browser Browser overview Navigating the browser Selecting PCD Images in the Browser Compare Series Loading a series to viewer from the browser Deleting a series Editing patient details Archiving patient images Archiving to PACS	205
Using tools on the acquisition tab Chapter 10 Browser Browser overview Navigating the browser Selecting PCD Images in the Browser Compare Series Loading a series to viewer from the browser Deleting a series Editing patient details Archiving to PACS Chapter 11 Viewer Viewer overview	205
Using tools on the acquisition tab Chapter 10 Browser Browser overview Navigating the browser Selecting PCD Images in the Browser Compare Series Loading a series to viewer from the browser Deleting a series Editing patient details Archiving to PACS Chapter 11 Viewer Viewer overview Setting window width and center	205
Chapter 10 Browser	205

Viewing or crea	ting PCD lodine Overlay images	
Viewing images	in MPR	
Understanding	and using slab	
Viewing images	in 3D	
Viewing images	in CTP	
Chapter 12 Re	construction	
Reconstruction	overview	
Performing a N	laterial Decomposition Image reconstruction	
Chapter 13	Accessories and Options	241
Accessories and	options overview	
Using bed adap	ters	
Using bed adap	ter safety straps	
Attaching bed a	dapter with or without posts	
Using the remo	vable T-square handle with the bed adapter	
Transfer boards	for adult, pediatric, and neonate patients	
Using the unive	rsal transfer board and silhouette scan board	
Using the pedia	tric scan platform	
Using the infan	t and neonate scanning platform	
Positioning usir	ng the infant and neonate scanning platform	
Adjusting the ir	fant and neonate platform handle	
Chapter 14	Using the OmniTom Elite with PCD in the Operating Room	256
Using the opera	ating room table adapter	256
Using the unive	rsal transfer board in the OR	
Annondix A	Glossom	250
Appendix A		
Appendix B	Error Codes	266
Appendix C Em	ergency Service Kit Procedure	275
Introduction	275	
Purpose		275

Lifting the Omr	iTom	
In Transport Mo	ode	275
Jacking Up the	OmniTom	275
Positioning the	Dolly	279
Lowering the O	mniTom onto the dolly	
In Scan Mode		
Jacking up the (DmniTom	
Positioning the	Dollies	
Lowering the O	mniTom onto the Dolly	
Moving the On	iniTom	
Appendix D	Revision History	285
Appendix E	Varex Imaging Data	286

List of Figures

Figure 1: Scanner dimensions	36
Figure 2: Product marking plate	37
Figure 3: Laser safety symbol	51
Figure 4: Dangerous to patient and operator label	55
Figure 5: Tablet battery capacity icon	58
Figure 6: Acquisition, positioning, and transport tabs on the LCD touch screen	65
Figure 7: OmniTom Elite with PCD tablet	66
Figure 8: Tablet safe distance location (two views)	66
Figure 9: Silhouette scan board and universal transfer board	67
Figure 10: Acquisition tab	69
Figure 11: Positioning tab	69
Figure 12: Transport tab	70

Figure 13: Initial screen when drive bar calibration begins	73
Figure 14: Retry calibration screen	74
Figure 15: Drive bar calibration failed	74
Figure 16: Active transport tab on LCD	75
Figure 17: Transport mode	75
Figure 18: Drive bar motions	76
Figure 19: Enabled collision avoidance sensors	77
Figure 20: Disable collision sensor icon	77
Figure 21: Mute audible alerts icon	78
Figure 22: Optional drive view camera	78
Figure 23: System icon	78
Figure 24: System settings icon	78
Figure 25: System settings	79
Figure 26: Drive view	79
Figure 27: Entering drive view pop-up	80
Figure 28: Front of OmniTom Elite with PCD with mood ring light enabled	80
Figure 29: Mood ring light enable/disable feature	81
Figure 30: Active positioning tab on LCD	81
Figure 31: Bed adapter without posts insertion	82
Figure 32: Bed adapter with T-square	83
Figure 33: Attaching the scan board with bed adapter to patient's bed	83
Figure 34: Positioning the patient on the bed	83
Figure 35: Ensuring placement of patient's neck directly under laser light	84
Figure 36: OmniTom Elite with PCD alignment camera	84
Figure 37: QR code alignment indicators	85
Figure 38: Adult scan board	85
Figure 39: Align icon	86
Figure 40: Looking for alignment markers	86
Figure 41: Alignment found	86
Figure 42: Alignment not found	86
Figure 43: Alignment moving	87
Figure 44: Alignment complete	87
Figure 45: Power down pop-up	89

Figure 46: Enable remote support	90
Figure 47: Enable remote support prompt	
Figure 48: Disable remote support	90
Figure 49: Disable remote support prompt	90
Figure 50: Login panel	92
Figure 51: Tap to logIn prompt	92
Figure 52: Access denied prompt	92
Figure 53: Registration tab	93
Figure 54: Automatic lock out screen	
Figure 55: System lock button	
Figure 56: Login screen on LCD	95
Figure 57: System icon	95
Figure 58: Logout icon	95
Figure 59: Confirm logout prompt	
Figure 60: System icon	
Figure 61: Shutdown icon	
Figure 62: Confirm shutdown prompt	
Figure 63: System icon	
Figure 64: System screen	
Figure 65: Privacy screen	
Figure 66: Scanner status screen	
Figure 67: Mood ring light	102
Figure 68: System screen	103
Figure 69: Protocol screen	105
Figure 70: Patient screen	105
Figure 71: Tablet tabs to perform a patient examination	106
Figure 72: User accounts dialog box	111
Figure 73: Delete pop up	112
Figure 74: System icon	113
Figure 75: Protocol manager icon	113
Figure 76: Adult or pediatric selection	113
Figure 77: Body part selection	113
Figure 78: Protocol selection	113

Figure 79: Edit icon	113
Figure 80: Series selection	114
Figure 81: Max field selection	114
Figure 82: Update icon	114
Figure 83: Save icon	114
Figure 84: Close button	115
Figure 85: Dose notification message	115
Figure 86: Diagnostic reason dialog box	115
Figure 87: System icon	116
Figure 88: System setting icon	116
Figure 89: Dose configuration icon	116
Figure 90: Dose configuration dialog box	116
Figure 91: Entering mGy and mGy.cm values	117
Figure 92: Define dose alert	117
Figure 93: Dose alert	118
Figure 94: Dose alert dialog box	118
Figure 95: PACS configuration dialog box	119
Figure 96: Audit trail viewer dialog box	120
Figure 97: Calendar dialog box	120
Figure 98: Audit trail viewer options	120
Figure 99: Recon presets dialog box	121
Figure 100: New preset dialog box	122
Figure 101: Recon preset groups dialog box	123
Figure 102: New recon preset group dialog box	123
Figure 103: MPR presets dialog box	124
Figure 104: New MPR preset dialog box	124
Figure 105: Window preset options	125
Figure 106: Window preset option	125
Figure 107: Image orientation	126
Figure 108: Patient orientation toggle	127
Figure 109: Orientation verification dialog	127
Figure 110: Protocol manager	128
Figure 111: Protocol Manager New Protocol	129

Figure 112: New protocol dialog box	129
Figure 113: New series dialog box	130
Figure 114: Protocol manager	133
Figure 115: Copy Protocol	133
Figure 116: Copied Protocol	134
Figure 117: Save Modified Protocol	134
Figure 118: Copied Protocol	134
Figure 119: Save Copied Protocol	135
Figure 120: Adjust protocol order	135
Figure 121: Edit button illuminated	136
Figure 122: Edit protocol dialog box	137
Figure 123: Protocol manager with a protocol selected for deleting	138
Figure 124: Delete protocol icon	138
Figure 125: Delete protocol confirmation pop-up	138
Figure 126: Protocol manager hide protocol function	139
Figure 127: Protocol manager move protocol function	140
Figure 128: PCD Water/QA Phantom	142
Figure 129: Water phantom inserted in the scanner bore	142
Figure 130: Set the laser in the middle of water portion of the phantom	142
Figure 131: Daily Calibration Screen	143
Figure 132: Show Calibration Percentages	144
Figure 133: Daily Calibration Progress Bar	144
Figure 134: PCD Daily Calibration success	145
Figure 135: PCD daily calibration status	145
Figure 136: Tube seasoning prompt	146
Figure 137: Tube seasoning dialog box	146
Figure 138: OmniTom with PCD QA Phantom	147
Figure 139: Phantom's etchings appear on top and sides	148
Figure 140: PCD Quality Assurance dialog box	148
Figure 141: PCD Quality Assurance Continue	149
Figure 142: PCD Quality Assurance results	149
Figure 143: Images of ACR module 2 for each resolution and scan mode. The ROI's in the image show approximately where t ROI's were measured	the 155

Figure 144: Sample images of the ACR phantom for Axial scan (top) and Helical scan (bottom) with 308mm FOV	159
Figure 145: Sample images of the ACR phantom for Axial scan with 50mm FOV	159
Figure 146: Normalized Noise Power Spectrum magnitude plot for the three reconstruction kernels for Axial and Helical sca modes (Top: Axial scan, Bottom: Helical scan)	ın 160
Figure 147: The measured beam width of the OmniTom Elite with PCD scanner	161
Figure 148: OmniTom Elite with PCD dose profile	161
Figure 149: Coronal reformation of the ACR scan using SR mode	162
Figure 150: Slice sensitivity profile of SR mode axial scan of the ACR bead phantom	162
Figure 151: The scatter measurement setup	164
Figure 152: The isodose curves for 120 kV, 70 mAs using the CTDI 16cm phantom. The distances are in meters, and the sca measured in μR.	tter is 165
Figure 153: The pediatric body phantom	166
Figure 154: Active registration tab	169
Figure 155: Registration dialog box	169
Figure 156: Registration tab	171
Figure 157: Query information dialog box	171
Figure 158: Query fields	172
Figure 159: Registration query results table	173
Figure 160: Registration stored results table	174
Figure 161: Stored list	174
Figure 162: Manual Add button	175
Figure 163: Manually Add Pt Info Dialog Box	175
Figure 164: Active acquisition tab	177
Figure 165: Radiographic film of the 60mm scan coverage	179
Figure 166: Scan coverage and imaged region for a true coverage of 60mm	179
Figure 167: PCD Resolution options	181
Figure 168: PCD accept icon	182
Figure 169: PCD Exam planner dialog box	182
Figure 170: PCD protocol series dialog box	183
Figure 171: PCD Ready, press 'scan' on scanner	183
Figure 172: PCD initiate scan on LCD	184
Figure 173: PCD setting scout parameters	184
Figure 174: PCD ready. Press 'SCAN' on scanner	185
Figure 175: PCD Complete dialog box	186

Figure 176: Finalize before additional reconstructions start	186
Figure 177: Scan Complete	187
Figure 178: Start New	187
Figure 179: Reuse Scout	188
Figure 180: Accept new protocol	188
Figure 181: Begin	188
Figure 182: Reused Scout	189
Figure 183: Repeat scan pop up	189
Figure 184: Step & Shoot dialog box	190
Figure 185: Options dialog box with AEC enabled	191
Figure 186: Toggle AEC graph icon	192
Figure 187: AEC graph	192
Figure 188: mA modulation	193
Figure 189: Options	193
Figure 190: Noise Reduction	194
Figure 191: Noise Reduction Levels	194
Figure 192: Save	194
Figure 193: Dynamic-CTA options	195
Figure 194: Parameters on scout	196
Figure 195: Series boxes	196
Figure 196: Active scan regions on scout	197
Figure 197: Bolus ROI	197
Figure 198: Completed CTA	199
Figure 199: Parameters on scouts	200
Figure 200: Bolus timing graph	201
Figure 201: CT perfusion dynamic scan options	201
Figure 202: Set dynamic CTP scan location	202
Figure 203: Review completed scan	203
Figure 204: Active browser tab	208
Figure 205: Browser studies and series tables	209
Figure 206: Material Decomposition results	210
Figure 207: Calcium Maps	210
Figure 208: Compare Series Arrow	210

Figure 209: Compare Series Viewer	211
Figure 210: Series Selection Buttons	211
Figure 211: Series Selection Dropdowns	212
Figure 212: 2D viewer	213
Figure 213: Delete pop up	213
Figure 214: Edit patient details	214
Figure 215: Archive pop up	214
Figure 216: Select the PACS server	215
Figure 217: PACS queue dialog box	215
Figure 218: Active viewer tab	217
Figure 219: 2D, 3D, MPR, and CTP viewing tab	220
Figure 220: Windowing preset list, text boxes, and the apply button	221
Figure 221: 2D tools	221
Figure 222: PCD Series	222
Figure 223: PCD 2D Viewer	223
Figure 224: keV and lodine buttons	223
Figure 225: PCD keV dialog box	223
Figure 226: keV type in	224
Figure 227: PCD keV Scroll buttons	224
Figure 228: PCD Series	225
Figure 229: PCD 2D Viewer	225
Figure 230: keV and lodine buttons	225
Figure 231: PCD lodine dialog box	226
Figure 232: Iodine percentage type in	226
Figure 233: PCD Iodine Scroll buttons	227
Figure 234: MPR tools	227
Figure 235: Coronal, sagittal, and transverse plane options	228
Figure 236: Set the cyan lines	229
Figure 237: Slab option	229
Figure 238: Using the tilt tool	230
Figure 239: Capture dialog box	230
Figure 240: 3D viewing	231
Figure 241: Dynamic CTP viewer	232

Figure 242: Arterial and venous ROI's	. 234
Figure 243: Adjust position of ROI	. 234
Figure 244: Perfusion maps	. 235
Figure 245: Active reconstruction tab	. 236
Figure 246: PCD Reconstruction studies and series tables	. 238
Figure 247: PCD reconstruction viewing pane	. 238
Figure 248: Material Decomposition Options	. 239
Figure 249: PCD begin reconstruction	. 240
Figure 250: Bed adapter without posts (left) and bed adapter with posts (right)	. 242
Figure 251: Safety Straps	. 242
Figure 252: Hill rom bed adapter	. 243
Figure 253: Stryker In Touch bed adapter	. 243
Figure 254: Straps attached to bed adapter	. 243
Figure 255: Silhouette scan board attached to bed adapter	. 244
Figure 256: Patient properly immobilized on scan board	. 244
Figure 257: Bed adapter without posts insertion (setscrew showing under arrow)	. 245
Figure 258: Bed adapter T-square handle	. 245
Figure 259: Bed adapter without posts shown being inserted into scan board	. 245
Figure 260: Silhouette with cushion shown on head holder	. 246
Figure 261: Steps showing removing T-square handle	. 246
Figure 262: Steps showing reattaching T-square handle	. 247
Figure 263: Correct install depth	. 247
Figure 264: Universal transfer board and stiffeners	. 249
Figure 265: Four types of mattress stiffeners	. 249
Figure 266: Mattress stiffener in place	. 250
Figure 267: Universal transfer board properly positioned on the bed on a mattress stiffener	. 250
Figure 268: Universal transfer board with safety strap installed	. 251
Figure 269: Pediatric scan platform	. 251
Figure 270: Child placed on pediatric platform with safety strap (two views)	. 252
Figure 271: Platform brake	. 252
Figure 272: Infant and neonatal scan platform	. 253
Figure 273: Pediatric strap	. 253
Figure 274: Applying pediatric strap onto the platform (three views)	254

Figure 275: Proper position of neonate/infant for head scans (left) and body scans (right)	254
Figure 276: Neonate platform foot brake location	254
Figure 277: Locking pins for the handle	255
Figure 278: Three positions for the platform handle	255
Figure 279: OR table adapter	256
Figure 280: OR table adapter attached to mount	256
Figure 281: Jack Components	275
Figure 282: Closeup of Release Valve	276
Figure 283: Placement of space block	276
Figure 284: Using the release valve	277
Figure 285: Inserting the jack bar	278
Figure 286: Proper placement of Jack	278
Figure 287: Move the dollies into place	279
Figure 288: Correct placement of Omniwheel on dolly	279
Figure 289: Jack Components	280
Figure 290: Using the release valve	281
Figure 291: Inserting the jack bar	282
Figure 292: Jacking up the Scanner	282
Figure 293: Correct placement of Omniwheel on dolly	283
Figure 294: Proper placement of dolly	283

List of Tables

Table 32: Detector collimation	152
Table 33: kV vs. dose in air	
Table 34: CT values of the ACR inserts for each resolution mode	
Table 35: Results of low contrast resolution test	
Table 36: CT uniformity and noise level of ACR accreditation phantom for Axial scans	
Table 37: CT uniformity and noise level of ACR accreditation phantom for Helical scans	
Table 38: Axial image noise results	156
Table 39: Helical image noise results	157
Table 40: Cutoff frequencies of Axial scans for the kernels at different MTF responses in lp/cm	
Table 41: Cutoff frequencies of Helical scans for the kernels at different MTF responses in lp/cm	
Table 42: The measured slice thickness	158
Table 43: Half-value layer	
Table 44: The energy conversion rates	
Table 45: Scatter in vertical parallel to axis of rotation (µRem/s)	
Table 46: Scatter in vertical plane on patient side (mRem/s)	
Table 47: Exposure at two different mAs	
Table 48: Exposure at two different mAs	
Table 49: Linearity calculations (mGy)	
Table 50: Linearity calculations in accordance with IEC	
Table 51: Registration buttons	170
Table 52: Acquisition buttons	177
Table 53: PCD Slice Thickness Options	
Table 54: Bolus tracking icons	195
Table 55: Bolus tracking tool table	
Table 56: CTP Viewer Tools	203
Table 57: Image tools	205
Table 58: Command buttons	
Table 59: Store/print queue buttons	216
Table 60: Common tools	217
Table 61: MPR tools	219
Table 62: 3D tools	219
Table 63: MPR Options	229
Table 64: CTP Viewer Tools	
List of Tables	Page 20 of 289

Table 65: Reconstruction tools	236
Table 66: Scan boards and their weight-bearing restrictions	248
Table 67: Error code list	266

Photon Counting Detector Overview

The OmniTom Elite with Photon Counting Detectors (PCD) employs energy resolving detectors to register the arrival of individual photons, eliminating the issue of electronic noise in the images. PCD CT systems count each individual X-ray photon and measure their energy, therefore enabling dose efficient, high-spatial resolution multi-energy imaging.

OmniTom Elite with PCD is the same system as the OmniTom Elite with the only difference being the detector array system. Instead of using gadolinium oxysulfide energy integrating detectors (EID), it uses a cadmium telluride-based detector system.

OmniTom Elite with PCD provides the ability to capture CT data in multiple energy bands. The multiple sets of CT data are acquired at the same time with configurable energy thresholds.

Intended use of the system

The OmniTom Elite computed tomography (CT) system is intended to be used for X-ray computed tomography applications for anatomy that can be imaged in the 40cm aperture, primarily the head and neck.

OmniTom Elite with Photon Counting Detectors (PCD) configuration has multi-energy CT functionality with spectral capability for material decomposition and virtual monoenergetic images (VMI). OmniTom Elite with PCD is supported for adult imaging of the head and neck.



The OmniTom Elite with PCD only allows for a voltage parameter of 120kVp and scan current is selectable from 5 to 20mA.

Clinical benefit

Computed Tomography (CT) provides real time imaging of bone, soft tissue and blood vessels that can provide detailed information to diagnose, plan treatment for, and evaluate many conditions in adults and children. Additionally, the detailed images provided by CT scans may eliminate the need for exploratory surgery.

Consumer information

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Legal disclaimer

This user manual is intended as a guide for material supplied by NeuroLogica Corp. It provides the operator with the necessary information to carry out specific procedures and maintain NeuroLogica produced equipment. Use this manual in conjunction with instruction and training supplied by qualified NeuroLogica personnel.

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This user manual, though complete and accurate, may not provide answers to undocumented changes or unexpected results that could occur from system anomalies.

Contact information

Keep user information readily available to contact Customer Service about general assistance or reporting on serious incidents (should they occur).

In the case of a serious incident or adverse event, please notify NeuroLogica at the below contact information and establishments local competent authorities.

To provide any comments, suggestions, or corrections to this user manual, please write to and include chapter title and page number:

NeuroLogica Corporation

Customer Service	14 Electronics Avenue, Danvers, MA 01923 USA
USA and Canada	1-888-564-8561
International	1-978-564-8561
Email	support@neurologica.com

Note: If you have questions about the clinical use of your system, speak with the **Clinical Representative**.

If you have questions about the service or functional operation of the system, speak with the **Technical Representative**.

Winckels I	Medical [Devices Ex	kpertise
Europe Be	ergerweg	18 6085	AT Horn
The Nethe	erlands		
+31 (0)47	5 582285	5 Tel	
+31 (0)47	5 582278	3 Fax	
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2862			

Damage in transportation

Closely examine all packages at the time of delivery. If you see damage, notate **"damage in shipment"** on all copies of the freight bill **before** you accept or sign for delivery (by the hospital receiving agent).

Whether damage is noted immediately or concealed (noticed after delivery), damage **MUST** be reported to carrier **immediately** upon discovery, or within 14 days after receipt, and containers held for inspection by carrier.

Keep in mind – the transportation company *will not* pay a claim for damage if an inspection is not requested within the 14-day period.

User requirements

The equipment can *only* be operated by users who have received professional medical education and training, such as physicians, radiologists, and other medical specialists.

You, the user (the limited or restricted operator or the administrator), are a trained person who is certified to operate such systems **before** scanning or diagnosing patients. This training must include medical and x-ray education, and NeuroLogica applications training.

Everyone who uses this equipment must read, understand, and follow all instructions, precautions, and warnings.

Keep this user manual near the equipment. It is important to review the procedures and safety precautions periodically.

Essential performance

The OmniTom Elite with PCD offers the following essential performance factors for the system:

- Over radiation protection
- Rescan prevention
- Stray radiation exposure prevention
- Diagnostic performance

About this user manual

The instructions in this user manual describe how to use the NeuroLogica OmniTom Elite with PCD **Computed Tomography (CT)** system, manufactured by NeuroLogica Corp. OmniTom Elite with PCD is the trade name for the CT system and NL5100 is the device model.

This user manual *does not* provide medical explanations but does suggest potential applications for some of the software features. This user manual describes potential safety problems and how to avoid them.

Anyone who operates this system should have received training **before** attempting to scan or diagnose patients, to include medical and x-ray education, in addition to NeuroLogica applications training.

This manual is made available in electronic format to the customer as part of each product delivery. For electronic manuals, please go to: Forms.samsungneurologica.com. Click on "Downloads" and choose "CT Manuals". The site will ask for the serial number of your product and a password. The password can be provided by Field Service.

Translation of this manual is available for any country that does not allow for English labeling. Please reach out to NeuroLogica directly if translation is required.

Some statements and chapters in this user manual may refer to NL5000 (the original OmniTom Elite system with energy integrating detectors) and may not be applicable to OmniTom Elite with PCD.

Identified symbols and system classifications

The specifications and details of this user manual may change to improve the product or to enhance its performance.

Throughout this user manual, a yellow triangle with a black border and exclamation point is used to draw attention to those conditions or situations that fit one or more of the following criteria, which are definitions from ANSI Z535.5:

DANGER Indicates a hazardous situation, which if not avoided, will result in death or serious injury.



WARNING Indicates a hazardous situation, which if not avoided, could result in death or serious injury.

Conventions used in this user manual

Table 1: Conventions	used in	this	user	manual
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Convention	Use	
Commands to perform actions	To perform a string of commands, this user manual will present them as follows: Customize > System . This means click Customize and then click System .	
Bold	When content refers to commands, windows, screens, dialog boxes, pop-ups, tabs, buttons, options, keyboard keys, statuses, and modes, these items appear in bold for faster identification, especially in a procedure.	
Italic	Identifies a word that is emphasized for your attention.	
Numbered steps	Numbered paragraphs represent sequential steps that require you to take action in the sequence provided – unless otherwise instructed. Procedures that are numerical mean that the sequence is important to follow. You may perform some procedures out of the recommended sequence; however, the results may vary.	
Neter	The appearance of a note is as such:	
NOTES	Note: Indicates additional information to help you operate this product.	

CAUTION Indicates a hazardous situation, which if not avoided, could result in minor or moderate injury.

Convention	Use
	A cross reference appears in the electronic (.pdf) user manual as a hyperlink. To retrieve an electronic copy of this user manual (in .pdf), click Help > User Manual from the workstation.
Hyperlink (an electronic cross- reference)	A hyperlink is a quick way to go to another area of the user manual (the referred-to content) with a simple click. Hyperlinks appear like this: "Understanding the types of users" on page 91. In this case, hover the mouse pointer over the (gray) hyperlink text. The pointer changes to ⁽¹⁾ . Press the Ctrl key on your keypad and (simultaneously) left click the mouse button. After you left click the hyperlink, the hyperlink takes you to the referenced area in the user manual.

Understanding the use of "you" in this user manual

Unless specifically noted, the implied "you", in this user manual, is the user. It is assumed users/operators are certified and medically trained personnel, qualified to use these systems. If the user is not the (implied) operator, the user will be specifically identified as administrator.

The following identifies those actions each user is permitted to perform:

	Full access permission (rights) to the system and its configuration.
Administrator	Can create protocols, user names and passwords, and all
	functions of the system.
	Modified access permission (rights). Can modify protocols during
Limited operator	system use but cannot create and save protocols; has no access
	to system configuration.
Destricted energy	No access to create, modify, save, or delete protocols; has no
Restricted operator	access to system configuration.

Active and inactive objects

When a menu command is dim the item is not active (that is, not enabled). Menu commands can include options, buttons, tabs, etc. When the item is dim, it can mean additional and required tasks must be completed first or you do not have permission to complete the action. Inactive commands, buttons, and tabs are gray. An active menu command, option, button, tab, and field means you can use the item to perform an action. Active items are blue and/or highlighted, not dim.

Use environment for the OmniTom Elite with PCD

The OmniTom Elite with PCD is designed for use in the general hospital setting. The integrated drive system allows medical professionals to bring high quality imaging to the patient, be it in the ICU, Emergency Department, or Intraoperatively in the Operating Suite.

Chapter 1 Compliance and Safety Requirements

It is important that you are familiar with compliance and safety requirements to ensure you, the patient, and the systems are safe at *all* times.

IEC classification and symbols

T In accordance with International Safety Standard IEC 60601-1, the OmniTom Elite with PCD CT scanner is classified as Type B equipment; Class 1 equipment, internally powered equipment, and continuous connection to the supply remains in standby state and for specified loading.

Type B equipment provides an adequate degree of protection against shock, in particular regarding:

- Allowable leakage current.
- Reliability of the protective earth connection.
- In accordance with the International Safety Standard IEC 60601-1, the manufacturer is not responsible for any consequences caused by the unauthorized modification of this equipment.
- Earth leakage current.



WARNING Equipment is not suitable for use in the presence of a flammable anesthetic mixture with air or with oxygen or nitrous oxide.



WARNING Equipment is not suitable for use with oxygen or oxygen-enriched atmospheres.

Mode of operation is a continuous connection to the supply mains in standby state and for specified loading conditions.

The OmniTom Elite with PCD CT scanner is patient-environment equipment.

Symbol	Description
\sim	Alternating current
Ē	Protective earth (ground)
\checkmark	Functional Earth
	Caution: consult accompanying documents
<u>A</u>	Caution: risk of electrical shock
	Electrostatic sensitive devices
Ŕ	Type B equipment
A	X-ray warning
(InTriangle)	X-ray source assembly emitting
(0 ₂ 0)	Non-ionizing radiation
	Warning: laser in use
LASER RADIATION DO NOT STARE INTO BEAM CLASS 2 LASER PRODUCT Max Pewer Output. 1990 Wavelength 650ml Completes with IEC 60825-12014, 3id ed 0014-05). Socialis Journal	Warning: Laser Radiation Do Not Stare Into Beam Class 2 Laser Product Laser Output and Standards Information Label

Table 2: Applicable IEC symbols

Symbol	Description
Complies with 21 CFR 1040.10 and 1040.11 except for conformance with IEC 60825-1 Ed. 3., as described in Laser Notice No. 56, dated May 8, 2019.	Warning: FDA Laser Information
	Warning: high temperature
\square	Emergency switch
	Crush warning
	Foot/toe crush warning when lowering machine
↑	System up
+	System down
= 7.5 kg	Load limit for scan board
	Temperature limits
Ť	Keep away from rain for packaging
%	Humidity limit for packaging
	Warning: battery charging
-=	Fuse usage

Symbol	Description
	Refer to instruction in user manual/booklet
	Follow Instructions for Use
	Manufacturer
	Date of Manufacture
REF	Catalogue Number
SN	Serial Number
\triangle	Caution
	To indicate that separate collection for waste of electronic equipment
Rx Only	Prescription Device
MD	Medical Device Symbol

Symbol	Description
C E 2862	CE Mark or Conformité Européenne ; number below CE represent Notified Body number
	Intertek ETL (Edison Testing Laboratories) Mark
EC REP	European Authorized Representative Symbol



WARNING This x-ray unit may be dangerous to patient and operator unless safe exposure factors and operating instructions are observed.

Environmental specifications

CAUTION The specified environment must be constantly maintained: 24-hours a day, seven days a week.

Table 3: Operating environment

Operating	
Ambient temperature	10º C to 27º C (50º F to 80.6º F)
Relative humidity	20% to 85% (non-condensing)
Altitude	0-3048 m (0-10,000 ft.)
Storage	
Temperature (without batteries installed)	-20º C to 60º C (-4º F to 140º F)
Temperature (with batteries installed)	-20º C to 50º C (-4º F to 122º F)
Relative humidity	20% to 85% (non-condensing)
Powering system	
Time period prior to powering the system	24 hours

If the system is in a facility outside the noted operating temperature, it is recommended to allow the noted time for the system to acclimate to the environment.

Considerations when preparing gantry for use:



CAUTION Check for obstructions before moving and system setup.

CAUTION	Monitor scanner motion to prevent collision with surrounding environment and foreign objects.
CAUTION	Press red EMERGENCY STOP button immediately in case of abnormal or unexpected motion.
WARNING	Verify scanner is on its Translate system (fully down position) prior to positioning patient at scanner entrance.
WARNING	Make sure all extremities are clear of the scanner while lowering or raising it.
WARNING	Keep patient in view at all times. Ensure that the patient can be seen when the operator is near the LCD (touch screen) and EMERGENCY STOP button. Never leave the patient unattended when the patient is in the gantry.

NeuroLogica advises complying with local regulations and/or site recommendations as specified by the facility physicist or certified representative for the following:

- Use mobile x-ray protective-shielding devices. Technologists should be at the correct location and consider wearing personal radiation protective equipment.
- A radiation safety plan in the working-area boundaries, to include as needed, mobile x-ray protective shielding devices. Otherwise assign a larger, working area to avoid radiation to the public. Effective dose for people outside the working area should be less than 0.25mSv annually (equals to 5 uSv weekly). The air kerma rate 0.3 meters away from the working area will be smaller than 2.5 uGy/h. Have monitoring and personal dose management for occupational exposure and related public health care personnel.
- There should be a working plan before scanning. The plan should include CT condition, time, location, working area, scanning plan, and site-clearing method; clearly state the responsibilities of working, protection, and management personnel. Keep a good record of the whole process.
- Restrict the working control and monitor area. Place obvious warning signs at the control-area boundaries to prevent unauthorized personnel from entering. Installation of a working status indication light is recommended.
- In accordance with the safety plan, self-monitor during the scanning process. A certified radiation representative should monitor the working area and take measurements immediately if abnormal circumstances are detected. Additionally, this should be reported to the local environmental administrative and health departments. There should be a public notice at the working area, to include the nature of work, time, location, control area, name of the working department, person in charge of the project, contact telephone number, and radiation report telephone number.

Site specification

Table 4: Site specification

lssue	Comment
Receiving area	Secured
Packing material and	Near availability of a trash receptacle for
waste	dunnage
Boom dimonsions for uso	14ft x 14ft room with a finished, level floor;
Room dimensions for use	recommended the room be well lit
Power availability	120 VAC - 20 amp /240VAC – 16-amp wall
Fower availability	outlets
Floor flatness	< 0.120 in. (3mm) over 10 in. (250mm)

Note: Not all beds are compatible with this system. Please contact Customer Service for assistance.

Note: For good image quality, the recommended practice is to keep the system free from vibration and to maintain the flatness specification noted.

Table 5: System operating parameters

Operating voltage	100-240 VAC (±10%) (100-120 / 208-240 VAC)
Operating frequency	50Hz-60Hz (±5%)
Apparent resistance of supply mains at 120VAC	0.3 ohms
Operating current at 120VAC	16/8 amps
Heat dissipation	1672 watts

Table 6: Battery operating parameters

Operating voltage	57.5 - 83.95VDC per brick
Output current _(peak)	25 amps per brick

Hazardous substances

Table 7: Hazardous substances table

Substance/material	@ Weight/system
Lead	7.48kg (16.5lbs.)
Mercury	Okg (Olbs.)
Hexavalent chromium	Okg (Olbs.)
Polybrominated Biphenyls (PBB)	<0.45kg (<1lbs.)
Polybrominated Diphenyl Ethers (PBDE)	<0.45kg (<1lbs.)

Part numbers and product-marking plates

Table 8: Core-system-component part numbers and product-marking plate locations

Component	Part number	Product-marking plate locations
OmniTom Elite with	0-NI 5100-001	Near the main input plug or on the side of
PCD gantry	0-1112100-001	the system.
QA phantom	10-01573-001	On the front of the phantom.

Note: The applicable components making up the OmniTom Elite with PCD CT scanner is identified with the nameplate statement "This product complies with radiation performance standards, 21 CFR sub-chapter J."

Table 9: Core-system component dimensions and weight

Component	Size (inches)	Size (centimeters)	Weight	Weight
	(L x H x W)	(L x H x W)	(lbs)	(kg)
OmniTom Elite with PCD system	65.4 x 61.6 x 29.9	165.2 x 156.6 x 75.9	1700	726



Figure 1: Scanner dimensions


Figure 2: Product marking plate

Class 1 Type B medical devices

This equipment generates, uses, and can radiate radio-frequency energy. The equipment may cause radio-frequency interference to other medical and non-medical devices and to radio communications. To provide reasonable protection against such interference, this product complies with emission limits for Class 1 medical devices as stated in EN 60601-1-2.

However, there is no guarantee that interference will not occur in a particular installation. If this equipment is found to cause interference (which can be determined - by switching the equipment on and off), the user should attempt to correct the problem using one or more of the following measures:

Re-orient or relocate the affected device(s).

Increase the separating space between the equipment and the affected device.

Power the equipment from a source different from that of the affected device.

Consult the point of purchase or the service representative for further suggestions.

NeuroLogica Corp. is not responsible for any interference caused either by the use of interconnect cables other than those recommended or by unauthorized changes or modifications to this equipment. Unauthorized changes or modifications could void the user's authority to operate the equipment.

To comply with the regulations applicable to an electromagnetic interface for a **Group 1 Class A** medical device, note the following:

All interconnect cables to peripheral devices *must be* shielded and properly grounded.

Use of cables not properly shielded and grounded may result in the equipment causing radio-frequency interference in violation of the European Union's Medical Device Directive and FCC regulations.



CAUTION Ensure there is no potentially detrimental interaction of system's irradiation with a patient's active and implantable medical devices and/or body-worn and active medical devices.

CAUTION Do not use devices that intrinsically transmit radio waves (such as a cellular phone, radio transceiver, mobile radio transmitter, radio-controlled toy, etc.). Use of these devices near this equipment could cause this equipment to malfunction. Keep power of these devices turned off when near this equipment.

Medical staff in charge of this equipment are required to instruct technicians, patients, and other people who may be around this equipment to fully comply with the above regulations.

Focal spot

The x-ray tube has a single focal spot with nominal dimensions of 1.0mm wide by 1.0mm long, with a range of 1.0 to 1.4mm, as defined by IEC 336-601. The spot does not move by more than ± 0.2 mm due to thermal expansion in any direction. The tube does have a positional tolerance of ± 1 mm from the end of the anode and from the window.

Filtration

The x-ray tube's total filtration of the irremovable layers is 2.54mm of aluminum.

Source to Detector Distance (SID)

The SID value is 596mm.

Compliance statement

Note: All editions and years of revisions for standards noted in this chapter are static as of **Revision 00** of this *OmniTom Elite with PCD User Manual*.

The OmniTom Elite with PCD system complies with the regulatory requirements of the following:

- ISO 14971: Medical Devices Application of Risk Assessment to Medical Devices.
- CAN/CSA C22.2 No 60601-1:14 Medical electrical equipment Part 1: General Requirements for Basic Safety and Essential Performance.

- EN ISO 13485 Quality systems Medical devices Particular Requirements for the Application of Quality System.
- IEC 60601-1 Medical electrical equipment Part 1: General Requirements for Basic Safety and Essential Performance.
- IEC 60601-1-2 Medical Electrical Equipment Part 1-2: General Requirements for Basic Safety and Essential Performance - Collateral Standard: Electromagnetic Compatibility - Requirements and Tests.
- IEC 60601-1-3 Medical Electrical Equipment Section 1-3: General Requirements for Safety. Collateral Standard: General Requirements for Radiation Protection in Diagnostic X-ray Equipment.
- IEC 60601-1-6 Medical Electrical Equipment Part 1-6: General Requirements for Basic Safety and Essential Performance Collateral Standard: Usability.
- IEC 60601-2-44 Medical Electrical Equipment Part 2-44: Particular Requirements for the Safety of X-ray Equipment for Computed Tomography.
- IEC 60825-1:2014 Safety of Laser Products Part 1: Equipment Classification, and Requirements.
- IEC 60825-1:2007 Safety of Laser Products Part 1: Equipment Classification, and Requirements.
- IEC 62133 Secondary Cells and Batteries Containing Alkaline or Other Non-acid Electrolytes - Safety Requirements for Portable Sealed Secondary Cells, and for Batteries Made from them, for use in Portable Applications.
- IEC 62366 Application of Usability Engineering to Medical Devices.
- IEC 62304 Medical device software Software life cycle processes.
- JIS T0601-1: 2017 Medical Electrical Equipment General Requirements for Basic Safety and Essential Performance
- KS C IEC 60601-1:2013 Medical Electrical Equipment General Requirements for Basic Safety and Essential Performance
- AS / NZS IEC 60601.1:2015 Medical Electrical Equipment General Requirements for Basic Safety and Essential Performance
- NEMA XR-25 Specifies an Equipment Feature for CT Scanners to Produce Doserelated Notification and Alert Messages to Inform Operators Prior to Scanning if the Estimated Dose Would Exceed the Preset Levels.
- NEMA XR-29 Standard Attributes on CT Equipment Related to Dose Optimization and Manual.
- International Electrotechnical Commission (IEC) International Standards Organization, when applicable.
- NeuroLogica Corporation is ISO 13485:2016 and MDSAP certified.

EMI/EMC terms

Electromagnetic compatibility

Electromagnetic Compatibility (EMC) is the branch of electrical sciences that studies **Electromagnetic Interference (EMI)** which is the unintentional generation, propagation, and reception of electromagnetic energy with reference to the unwanted effects that such energy may induce. The goal of EMC is the correct operation, in the same electromagnetic environment, of different equipment, which use electromagnetic phenomena and the avoidance of any interference effects.

In order to achieve this, EMC pursues two different kinds of issues. Emission issues are related to the unwanted generation of electromagnetic energy, to the countermeasures that should be taken in order to reduce such generation, and to avoid the escape of any remaining energies into the external environment. Susceptibility or immunity issues, in contrast, refer to the correct operation of electrical equipment in the presence of unplanned electromagnetic disturbances.

Interference mitigation is achieved by addressing both emission and susceptibility issues, that is, quieting the sources of interference, making the coupling path between source and victim less efficient, and making the potential victim systems less vulnerable.

Radio frequency interference

Radio Frequency Interference (RFI) is a type of EMI in the radio frequency spectrum that causes an unwanted disturbance that affects an electrical circuit due to electromagnetic radiation emitted from an external source. The disturbance may interrupt, obstruct, or otherwise degrade or limit the effective performance of the circuit. The source may be any object, artificial or natural, that carries rapidly changing electrical currents, such as an electrical circuit, the sun, or the northern lights.

Susceptibility

Susceptibility is the capability of an electronic system to respond to unwanted electrical energy.

EMI/EMC compliance

This equipment complies with IEC 60601-1-2 EMC standard for medical devices.

The OmniTom Elite with PCD system is suitable to be used in an electromagnetic environment, as per the limits and recommendations described in the tables hereafter:

• Emission compliance level and limits (Table 11).

Immunity compliance level and recommendations to maintain equipment clinical utility (Table 12, Table 13, and Table 14).

Note: This system complies with the above-mentioned EMC standard when used with supplied cables. If different cable lengths are required, contact a qualified service representative for advice.

Acronym and abbreviation	Definition
AEC	Automatic Exposure Control
СТ	Computed Tomography
CTDI _{vol}	Volume Computed Tomography Dose Index
CTDI _w	Weighted average Computed Tomography Dose Index
DICOM	Digital Imaging Communication in Medicine
DLP	Dose Length Product (DLP)
DHCP	Dynamic Host Control Protocol
EMC	Electromagnetic Compatibility
ΕΜΙ	Electromagnetic Interference
FOV	Field Of View
HIS	Hospital Information System
HU	Hounsfield Unit
MAR	Metal Artifact Reduction
MIP	Maximum Intensity Projection
MPR	Multi-Planar Reformation, sometimes referred to as Multi-
	Planar Reconstruction
PACS	Picture, Archiving, and Communication System
PCD	Photon Counting Detectors
QA	Quality Assurance
RIS	Radiology Information System
RSO	Radiation Safety Officer
RFI	Radio Frequency Interference
SCP	Service Class Provider
SCU	Service Class User

Table 10: Acronyms and abbreviations



WARNING Medical, electrical equipment needs special precautions regarding EMC and needs to be installed and put into service according to EMC information provided in accompanying documents.



CAUTION Portable and mobile RF communications equipment can affect medical electrical equipment.

CAUTION Do not use or stack the equipment or system with other equipment and if adjacent or stacked use is necessary, the equipment or system should be observed to verify normal operation in the configuration in which it will be used.

Note: The EMC tables and other guidelines included in this user manual provide information to the user essential in determining the suitability of the equipment or system for the electromagnetic environment of use, and in managing the electromagnetic environment of use; to permit the equipment or system to

perform its intended use without disturbing other equipment and systems or non-medical electrical equipment.

Note: The EMISSIONS characteristics of this equipment make it suitable for use in industrial areas and hospitals (CISPR 11 class A). If it is used in a residential environment (for which CISPR 11 class B is normally required) this equipment might not offer adequate protection to radio-frequency communication services. The user might need to take mitigation measures, such as relocating or reorienting the equipment.

Table 11: Emission declaration for OmniTom Elite with PCD system

OmniTom Elite with PCD system is intended for use in the electromagnetic environment specified below. The user of the OmniTom Elite with PCD system should assure it is used in such an environment.

Emissions test	Compliance	Electromagnetic environment guide
RF emissions CISPR 11	Group 1	The OmniTom Elite with PCD system uses RF energy only for internal function. Therefore, the RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.
RF emissions CISPR 11 Harmonic	Class A	The OmniTom Elite with PCD system is predominantly intended for use in non- domestic environments, and not directly connected to the public mains network. The
emissions, IEC 61000-3-2 Voltage	Class A	OmniTom Elite with PCD system is predominantly intended for use (for example, in hospitals) with an appropriate power supply and the recommended
fluctuations/ flicker emissions, IEC 61000-3-3	Complies	shielding for mobile use.

Table 12: EMC immunity declaration for the OmniTom Elite with PCD system

OmniTom Elite with PCD system is intended for use in the electromagnetic environment specified below. The user of the OmniTom Elite with PCD system should assure it is used in such an environment.

lmmunity test	IEC 60601- 1-2 test level	Compliance level	Electromagnetic environment guidance
Electrostatic Discharge (ESD) IEC 61000-4-2	± 6 KV contact ± 8 KV air	± 6 KV contact ± 8 KV air	Floors should be wood, concrete, or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30%.
Electrical fast transient/burst IEC 61000-4-4	±2 KV for power supply lines ±1KV for input/ output lines	±2 KV for power supply lines ±1KV for input/output lines	Mains power quality should be that of a typical commercial or hospital environment.
Surge IEC 61000-4-5	± 1 KV line- line ± 2KV line- ground	± 1 KV line- line ± 2KV line- ground	Mains power quality should be that of a typical commercial or hospital environment.
Voltage dips, short interruptions, and voltage variations on power supply input lines IEC 61000-4-11	 >95% dip for 0.5 cycle 60% dip for 5 cycles 30% dip for 25 cycles >95% dip for 5 seconds 	 >95% dip for 0.5 cycle 60% dip for 5 cycles 30% dip for 25 cycles >95% dip for 5 seconds 	Mains power quality should be that of a typical commercial or hospital environment. If the user of the OmniTom Elite with PCD system requires continued operation during power interruptions, it is recommended that the OmniTom Elite with PCD system be powered from its internal batteries.
Power frequency (50/60Hz) magnetic field IEC 61000-4-8	3 A/m	3 A/m	Power-frequency magnetic- fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.

Note: The wireless receiver operates within the following bands. 2.412 to 2.462 GHz (11 channels) 5.180 to 5.240 GHz (4 channels) 5.260 to 5.320 GHz (4 channels)* 5.500 to 5.700 GHz (8 channels, excluding 5.600 to 5.640 GHz)* 5.745 to 5.825 GHz (5 channels) The preferred frequency band is 5.189 to 5.240 GHz at 40MHz bandwidth. The wireless transmitter operates within the following frequency bands and power. 802.11b: Typ. 26±1.5 dBm @ 1 Mbps, Typ. 26±1.5 dBm @ 2 Mbps Typ. 26±1.5 dBm @ 5.5 Mbps, Typ. 25±1.5 dBm @ 11 Mbps 802.11g: Typ. 23±1.5 dBm @ 6 to 24 Mbps, Typ. 22±1.5 dBm @ 36 Mbps Typ. 20±1.5 dBm @ 48 Mbps, Typ. 19±1.5 dBm @ 54 Mbps 802.11n (2.4 GHz): Typ. 23±1.5 dBm @ MCS0/8 20 MHz, Typ. 18±1.5 dBm @ MCS7/15 20 MHz Typ. 23±1.5 dBm @ MCS0/8 40 MHz, Typ. 17±1.5 dBm @ MCS7/15 40 MHz 802.11a: Typ. 23±1.5 dBm @ 6 to 24 Mbps, Typ. 21±1.5 dBm @ 36 Mbps Typ. 20±1.5 dBm @ 48 Mbps,Typ. 18±1.5 dBm @ 54 Mbps 802.11n (5 GHz): Typ. 23±1.5 dBm @ MCS0/8 20 MHz, Typ. 18±1.5 dBm @ MCS7/15 20 MHz Typ. 23±1.5 dBm @ MCS0/8 40 MHz, Typ. 18±1.5 dBm @ MCS7/15 40 MHz The device includes 4 dBi gain antennas.

Countermeasures against EMC related issues

Generally, it is difficult to solve issues related to EMC. It may take a variable amount of time and cost to identify issues causing interference.

General countermeasures to minimize EMI are as follows:

- Electromagnetic interference may be alleviated by positioning other equipment farther away from the source of the EMI.
- MI may be mitigated by changing relative location (installation angle) between system and other equipment.

- EEMI may be eased by changing wiring locations of power/signal cables of other equipment.
- EMI may be reduced by altering the power-supply path of other equipment.
- Electromagnetic environment specified (Table 12 and Table 13).

Table 13: EMC immunity declaration

OmniTom Elite with PCD system is intended for use in the electromagnetic environment specified below. The user of the OmniTom Elite with PCD system should assure it is used in such an environment.

Immunity	IEC 60601-1-	Compliance	Electromagnetic environment
test	2 test level	level	guidance
Conducted RF IEC 61000-4-6	3 Vrms 150kHz to 80MHz	V1 = 3 Vrms	WARNING: Portable and mobile RF communications equipment should be used no closer to any part of the OmniTom Elite with PCD system, including cables, than the recommended separation distance calculated from the equation appropriate for the frequency of the transmitter.
Radiated RF IEC 61000-4-3 (alternative method: IEC 61000-4-21)	3 Vrms 80MHz to 2.5GHz	E1 = 3 V/m	$d = [\frac{3.5}{\nu_1}]\sqrt{P}$ $d = [\frac{3.5}{E_1}]\sqrt{P}$ 80 MHz to 800 MHz $d = [\frac{7}{E_1}]\sqrt{P}$ 800 MHz to 2.5 GHz Where P is the maximum power rating in watts and d is the recommended separation distance in meters. Field strengths from fixed transmitters, as determined by an electromagnetic site survey, should be less. than the compliance levels (V1 and E1). Interference may occur in the vicinity of equipment marked with the following icon: $((\cdot))$

Table 14: Recommended separation distances

Recommended separation distances between portable and mobile RF				
	CD system is intended	for use in an electron		
Ommon Eine with P	CD system is intended	for use in an electron		
environment in which	1 radiated KF disturband	ces are controlled. In	e user of the	
Omni iom Elite with P	'CD system can neip pre	ent electromagnetic	interference by	
maintaining a minimu	im distance between p	ortable and mobile Ki	- communications	
equipment (transmitt	ters) and the Omnilom	Elite with PCD system	as recommended	
below, according to the	he maximum output po	wer of the transmitte	rs.	
Rated maximum	150kHz to	80MHz to	800MHz to	
output Power	80MHz	800MHz	2.5GHz	
(P) if	Separation	Separation	Separation	
transmitter	distance	distance	distance	
Watts (W)	meters ¹	meters ¹	meters ¹	
0.01	0.12	0.12	0.23	
0.1	0.38	0.38	0.73	
1	1.2	1.2	2.3	
10	3.8	3.8	7.3	
100	12	12	23	
For transmitters rated	d at a maximum output	power not listed abo	ve, the separation	
distance is estimated	using the equation in t	he corresponding colu	umn, where P is the	
maximum output pov	maximum output power rating of the transmitters in Watts (W) according to the			
transmitter manufact	urer.			
Note: At 80MHz and	d 800MHz senaration d	listance for higher fre	quency range	
applies				
NI			· · · · ·	
Note: These guide	lines may not apply in a	ill situations. Electron	nagnetic	

propagation is affected by absorption and reflection from structures, objects, and people.

¹ Separation distance according to frequency of transmitter, measured in meters (m).

Table 15: Test specifications for ENCLOSURE PORT IMMUNITY to RF wireless communications equipment

Test Frequency (MHz)	Band ª)	Service ^{a)}	Modulation	Max Power (W)	Distance (m)	Immunity Test Level (V/m)
385	380- 390	Tetra 400	Pulse Modulation ^{b)} 18Hz	1.8	0.3	27
450	430- 470	GMRS 460, FRS 460	FM ^{c)} ±5kHz deviation 1 kHz sine	2	0.3	9

710			Pulse			
745	704-	LTE Band	Modulation			2
780	787	13,17	b)	0.2	0.3	9
/80			217Hz			
810		GSM				
870		800/900				
930	800- 960	1ETRA 800, iDEN 820, CMDA 850, LTE Band 5	Pulse Modulation ^{b)} 18Hz	2	0.3	28
1720		GSM				
1845		1800; CMDA				
1970	1700- 1990	1900; GSM 1900; DECT; LTE Band 1, 3, 4, 25; UMTS	Pulse Modulation ^{b)} 217Hz	2	0.3	28
2450	2400- 2570	Bluetooth, WLAN, 802.11 b/g/n, RFID 2450, LTE Band 7	Pulse Modulation ^{b)} 217Hz	2	0.3	28
5240			Pulse			
5500	5100-	802.11	Modulation	0.2	0.3	9
5785	5800	a/n	^{⊳)} 217Hz	0.2	0.0	

Use recommendations

This product complies with IEC 60601-1-2 standard for medical devices and with radio frequency emission requirements per CISPR11 Group 1 Class A standard limits. The OmniTom Elite with PCD system is predominantly intended for use in hospitals.

Do not use devices that intentionally transmit RF signals (cellular phones, transceivers, or radio-controlled products) in the vicinity of this equipment as it may cause performance outside the published specifications. Keep the power to these types of devices turned off when near this equipment.

Adhering to the distance separation (recommended in Table 14) between 150kHz and 2.5GHz, will reduce disturbances recorded at the image level, but may not eliminate all disturbances. When installed and operated as specified herein, the system will maintain its essential performance by continuing to safely acquire controlled, radiological, x-ray exposures in a mobile radiography environment. For example, a 1W mobile phone (800MHz to 2.5GHz carrier frequency) must be 2.3 meters apart from the OmniTom Elite with PCD system (in order to avoid image interference risks).

The use of accessories, transducers, and cables, other than those specified, may result in degraded, electromagnetic compatibility of the OmniTom Elite with PCD system.

The medical staff in charge of this equipment is required to instruct technicians, patients, and other people who may be around this equipment to comply fully with the above equipment requirements.

Installation recommendations

This system complies with the above-mentioned EMC standard when used with supplied cables.

In order to minimize interference risks, the following requirements apply.

Cable shielding and grounding

All interconnect cables to peripheral devices must be shielded and properly grounded. Use of cables not properly shielded and grounded may result in the equipment causing radio-frequency interference.

Adjacent components and equipment

OmniTom Elite with PCD system should not be used adjacent to or stacked with other equipment; if adjacent or stacked use is necessary, the OmniTom Elite with PCD system should be tested and verified to make sure normal operation in the configuration in which it is used. Consult NeuroLogica and Facility Technical Support staff regarding device/system configurations.

Static magnetic field limits

In order to avoid interference on the OmniTom Elite with PCD system, static-field limits from the surrounding environment are specified. Static field is specified as less than 1 Gauss around the unit.

Electrostatic discharge environment and recommendations

- In order to reduce electrostatic-discharge interference, install a charge-dissipative floor-material to avoid electrostatic charge-buildup.
- The relative humidity must be at least 30 percent.
- The dissipative material must be connected to the system ground-reference.

Facility IT-NETWORK

The OmniTom Elite with PCD system utilizes the IT-NETWORK for the customer (as applicable) to communicate with the modality worklist and for supplemental, imagestorage space. As part of the installation, the applicable IT-NETWORK is reviewed to create the appropriate setup for the system. Setup is done to ensure no potential concerns arise with the system.



CAUTION It is possible that the IT-NETWORK connection from the system could result in previously unidentified issue(s) to the respective population. Should this occur, please contact Customer Service right away to identify, analyze, evaluate, and resolve the issue(s).



CAUTION It is possible that any changes to the IT-NETWORK made by the facility could introduce a new issue where Customer Service needs to be contacted to address the concern, right away.

Changes to the IT-NETWORK include – but are not limited to the following:

- Changes in network configuration
- Connection to additional items
- Disconnection to items
- Updating equipment
- Upgrading equipment.

Hazard Information

Please review this material before using the system and observe basic, common-sense safety rules when operating this scanner.

General safety considerations and statements

Review the following before using the system and observe basic, common-sense safety rules when operating the scanner:

- Become familiar with the functional hardware to help recognize serious problems.
- Do not use scanner if it appears damaged or fails.
- Wait for qualified personnel to correct any problem.

WARNING	The health software is installed on a medical device and is required for its operation. In order to securely remove the software from use, the system must be decommissioned.
WARNING	Modification of this equipment is not allowed.
CAUTION	All non-medical electrical equipment will comply with relevant IEC and ISO safety standards.
CAUTION	Federal law restricts the use of this device without a prescription by a physician.
CAUTION	Always store and/or use unit in a well-ventilated area. Keep air pollution to a minimum. Keep floor clean at all times.
CAUTION	Do not touch parts of non-medical electrical equipment in patient environment and patient simultaneously.
CAUTION	For disposal of any material emanating from the system; follow local regulations.
CAUTION	This system was designed for use by individuals trained in CT system operation. The user should be familiar with this user manual before scanning patients.
CAUTION	It is the user's responsibility to make sure that after installation or subsequent modification, the system will be in compliance with the requirements of collateral standard IEC 60601-1.
WARNING	Installation of this product is performed in accordance with Installation Manual (1-NL5100-059). All installation processes and personnel qualifications are outlined in that document.
WARNING	Proper disposal of batteries is required to ensure compliance with environmental safety guidelines. Contact authorized NeuroLogica representative for instructions.
WARNING	Observe safety-exposure factors and operating procedures to protect patient from physical harm during contact with this x-ray scanner.
WARNING	Observe safety requirements to prevent excessive dose exposure to patient and/or operator.
CAUTION	Improper system usage could endanger patients and/or users and void the warranty if not operated correctly.
CAUTION	Should the Tablet encounter a computer related virus, contact Technical Support for assistance with removing the virus from the equipment.
CAUTION	Radiation dose exposure to patients should not exceed 1Gy CTDI.

CAUTION	For proper disposal of material at equipment's end-of-useful life, as a service to the user, NeuroLogica can dispose of the device. Please contact NeuroLogica's dealer or customer service at 1-888-564- 8561.
CAUTION	Equipment that uses only basic insulation to protect against electric shock should not be used in this system.
CAUTION	If issues occur while obtaining patient dose report, contact Technical Support for assistance.

Laser safety



The lasers themselves are mounted internally to the disk assembly, which spins (as prescribed by the control panel) within the system's bore. Therefore, the laser's output light will always be aimed at, and rotating within the bore itself.

Laser parameters

Wavelength = 650nm Output Power = 1mW

Output Po	
WARNING	Complies with 21 CFR 1040.10 and 1040.11 except for conformance with IEC 60825-1 Ed. 3., as described in Laser Notice No. 56, dated May 8, 2019.
WARNING	Viewing the laser output with certain optical instruments (for example, eye loupes, magnifiers, and microscopes) within a distance of 100mm may pose an eye hazard.
CAUTION	Instruct the patient to close his/her eyes before activating (turning ON) the alignment light.
CAUTION	Closely monitor infants and infirm patients to prevent them from accidentally staring into the beam.
CAUTION	Class 2 laser radiation when open. <i>Do not</i> stare into the beam or view directly with optical instruments.
CAUTION	Use of controls or adjustments or performance of procedures other than those specified herein may result in hazardous radiation exposure.



CAUTION The warning label (below, for "laser in use") is located on the front of the scanner cover and also inside the scanner to identify the presence of a laser.

Scanner mobility safety

CAUTION Due to the mobility of the system, an external interlock is not available; however, a prescribed scan can, at any time, be terminated from the scanner or the Tablet. When the user activates the scan, a 10-second, countdown-clock, scan delay (adjustable to 99 seconds) triggers. This countdown allows the user time to perform needed tasks before the scan begins.

- **WARNING** To prevent involuntary movement, do not position scanner on an incline while in **Transport** mode.
- **WARNING** Contact Technical Support for assistance when movement is required on an incline.
- **WARNING** Do not move the system right or left if transport on an incline becomes necessary. Always keep the system in a straight motion.
- WARNING This system shall not be transported on an incline greater than 5°.

Note: Be sure there are no obstacles in front of the scanner when it is being transported.

- **CAUTION** Check to ensure proper clearance is provided to allow removal of patient from scanner in case of a power failure. This is accomplished by moving patient's support (after unlocking wheel-locks) away from scanner.
- **CAUTION** To prevent patient entrapment or entanglement with accompanying equipment, slowly move scanner away from patient using the LCD (touch screen) while observing patient.
- **CAUTION** Do not station or operate the system on an uneven floor. The flatness requirement is 3mm over a distance of 300mm.
- **CAUTION** Prior to transporting the scanner, verify that power cord is unplugged from wall to avoid damage to cord and outlet.

Floor level (even)

For proper operation, the system must be operated on an even, level, hard surface.

Carpeting

Do not use the system on a carpeted floor. Poor image quality could result due to unevenness of the floor.

Electrical safety

WARNING	The system's external AC power cord should be checked prior to use to verify there are no exposed wires or damaged insulation/prongs. Damaged prongs could result in sparking and fire. In case of such damage, contact Customer Service immediately.
CAUTION	Check to ensure the AC outlet is working properly before plugging in the system's AC power cord.
WARNING	To prevent electrical shock, do not connect items that are not specified as part of the system.
WARNING	To prevent electrical shock, do not remove the covers from the equipment. The covers protect the user and the patient from moving parts and electrical shock. Hazardous voltages are present within this equipment. The covers provide protection from radiation exposure given off from the x-ray tube. The covers also protect the equipment.
WARNING	An electrical shock hazard: no user should replace parts. Refer to qualified service personnel for any service.
WARNING	Always electrically isolate this equipment from the main electrical supply before cleaning and disinfecting it. This is necessary to prevent short-circuiting or possible electrical shock.
WARNING	Never position the mobile system in a manner that prohibits access to unplugging it or prohibits pressing the EMERGENCY STOP button.
WARNING	To minimize shock hazard, the system chassis must be connected to an electrical ground. The system is grounded through the ground conductor of the supplied, three-conductor power cord. The power cord must be plugged into a three-conductor electrical outlet receptacle. Do not alter the ground connection.
WARNING	Avoid all contact with any electrical conductor as follows:
	 Allow only people who know the proper procedures and use the proper tools to install, adjust, repair, or modify the equipment. Only use this equipment in rooms or areas that comply with all applicable laws (or regulations having the force of law) concerning electrical safety for this type of equipment.

• Always electrically isolate this equipment from the main electrical supply before cleaning and disinfecting it.

	 The detachable cord is the disconnecting device, which is used to remove mains power from the wall socket. The system is internally powered.
WARNING	For Class 1 equipment using an alternate internal source: a warning to use the alternate source if the integrity of the protective earth conductor is in doubt.
WARNING	Do not position the system so that it is difficult to access the AC power cord.
CAUTION	Protect the system power cord against mechanical damage.
	Where the integrity of external, protective conductor in the installation or its arrangement is in doubt, equipment is operated from its internal electrical power source.
	Parts of non-medical electrical equipment in the patient environment that, after removal of covers, connectors, without the use of a tool, may be contacted by the operator during routine maintenance and calibration, will operate at a voltage not exceeding 25VAC or 60VDC or peak value supplied from a source that is separated from the supply mains in accordance with one of the methods described in IEC 60601-1.
CAUTION	To help prevent tripping hazards, use care in the arranging of any cords (for example, AC cord, Ethernet cable, etc.) when connecting to the system.
CAUTION	To prevent damaging electrical outlet cords, check to ensure they have been removed and properly stored before transporting the scanner.
CAUTION	All systems within the patient environment provide the same level of safety as medical equipment complying with IEC 60601-1.
WARNING	The OmniTom Elite with PCD CT scanner contains high-voltage circuits for generating x-rays. Only trained and qualified personnel should be permitted access to the internal parts of this equipment.

Mechanical safety

WARNING In case of unwanted movement or motion, press the EMERGENCY STOP (E-STOP) button.

WARNING Physically assist all patients on and off the bed and into position on the scan board. Adjust the bed to the specified height for patient loading and unloading.

WARNING	When positioning the scanning platform, be careful when moving the patient support to avoid having it hit the scanner covers.
WARNING	Position any lines (IVs, respirator hoses, electrical leads, etc.) attached to the patient so the lines cannot catch on the scanner during scanner travel.
CAUTION	Prevent pinching or crushing of the patient's extremities. Keep patient's hands on the side of his/her body. Watch the patient and equipment carefully at all times during scanner movement.
CAUTION	To prevent pinching or crushing of the operator's feet/toes, be sure extremities are not positioned under the scanner when it is being lowered from Transport mode to Scan mode.

Radiation safety



Figure 4: Dangerous to patient and operator label

WARNING	Improperly used x-ray equipment may result in unwanted radiation exposure. Read and understand the instructions in this user manual before attempting to operate this equipment.
CAUTION	Use technique factors prescribed by the radiologist or diagnostician. Use a dose that produces the best diagnostic results with the least x-ray exposure.
CAUTION	All persons authorized to use the equipment must understand the dangers from excessive x-ray exposure. NeuroLogica recommends use of protective materials and devices.
WARNING	Everyone having anything to do with x-ray must take adequate steps to insure protection against injury.
CAUTION	The use of this device requires its users to receive proper training in accordance with local and national laws.
CAUTION	Never perform calibration with patients in the scanner or while personnel are present in the vicinity of the scanner.
CAUTION	Amber indicator lights (on the top of the scanner) illuminate during x-ray exposure.
CAUTION	Ensure that there is no potential for detrimental interaction of the system's irradiation with a patient's active implantable medical devices and/or body-worn, active, medical devices.



CAUTION For any questions on load factors or calculating the applicable dose, Contact NeuroLogica for service. See "Contact information" on page 23.

X-rays can only be produced during the following conditions:

- The scanner is in the down (Scan mode) position.
- The **Start** button is activated.
- The **Tablet** is connected.

Fire and explosion safety

DANGER This equipment is not suitable for use in presence of a flammable anesthetic mixture with air or with oxygen or nitrous oxide.

DANGER This equipment is not suitable for use in the presence of oxygen or an oxygen-enriched atmosphere.

- Fire regulation for the type of medical area being used should be fully applied, observed, and enforced. Fire extinguishers should be provided for both electrical and non-electrical fires.
- All operators of the OmniTom Elite with PCD scanner should be fully aware of and trained in the use of fire extinguishers and the firefighting equipment, and in local fire procedures.

WARNING Only use extinguishers on electrical or chemical fires that are specifically labeled for those purposes. Using water or other liquids on an electrical fire can lead to fatal or other serious injury.

If it is safe to do so, attempt to disconnect the equipment from electrical and other supplies before attempting to fight a fire. This will reduce the risk of electrical shocks.

EMERGENCY STOP button

CAUTION	Check the EMERGENCY-STOP (E-STOP) button at least one time a month to ensure proper function.
CAUTION	Every user should take a few minutes to locate the E-STOP before scanning the first patient.
CAUTION	In case of emergencies, stop scanner movement immediately by pressing the E-STOP red push-button located on the scanner, next to the LCDs (touch screens).



Battery system safety and information

The **System battery capacity** status icon shows an indication of battery capacity, which is identical to the indicator on the scanner. The user should always check the indicator on the scanner to verify the batteries' status.

- **CAUTION** The system unit contains batteries and will always be charging when plugged into AC mains.
- **CAUTION** In case of battery leakage, do not handle the batteries themselves nor continue to operate the system. Contact NeuroLogica for service. See "Contact information" on page 23.
- **WARNING** Do not immerse the battery in water or seawater and keep the battery in a cool dry surrounding when it stands by.
- WARNING Do not use or leave the battery near a heat source as fire or heater
- WARNING Do not reverse the position and negative terminals
- **WARNING** Do not discard the battery in fire or a heater.
- **WARNING** Do not short-circuit the battery by directly connecting the positive and negative terminals with metal objects
 - **WARNING** Do not transport or store the battery together with metal objects such as hairpins, necklaces, etc.
 - **WARNING** Do not strike, trample, or throw the battery
- **WARNING** Do not use or leave the battery at high temperature (for example, at strong direct sunlight or in a vehicle in extremely hot weather).

WARNING If the battery gives off an odor, generates heat, becomes discolored or deformed, or in any way appears abnormal during use, recharging or storage, immediately contact NeuroLogica for service. See "Contact information" on page 23.

System battery capacity

The range is 0 to 100. Battery voltage and current are used to calculate the system's battery capacity based on charging (plugged in) state.

Run time operation

During normal run-time operation (all components, up and running), the battery capacity is calculated one time per second.

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State changes

After each periodic update, battery capacity is checked to make sure it does not fall below certain thresholds, as follows:

- Low voltage alarm state
- When the scanners battery capacity goes below 25%. It will remain in this state until the battery capacity has gone back up to 27% or higher.

Predictive scanning

Before each scan, battery usage for that scan is predicted based on the selected load factors (for example, kV, mA, time), and is compared against the available battery charge. In the case that there is not enough charge to complete a scan, a pop-up appears on the Tablet screen. The user can cancel the scan at that time or continue the scan, with the understanding that the scan may abort due to a low-power fault.

Under voltage protection

When the **system battery voltage** drops below the low-voltage cutout-level while unplugged, a system power-down sequence is initiated.

Recovering the system

Make sure the system is plugged into an operational wall outlet.

Note: The system will not boot up fully until the system reaches 10% charge. This can take up to 30 minutes. Full charge can take up to ninety (90) minutes.

If the system does not resume normal operation as the batteries recharge, there is likely a fault in the battery system; contact NeuroLogica for service.

Tablet

To check the charging status of the tablet, select the Scanner Status Icon in the top, right corner to see the capacity of the battery, ranging from 0 to 100%. The Scanner Status screen also displays a lightning bolt when the tablet is plugged in and charging. This screen is only active if the tablet is connected to an operational scanner. The user should always check the screen to verify the status of the batteries.



Figure 5: Tablet battery capacity icon

Note: The operator panel provides a way to verify that the scanner is plugged in and charging. The blue light if solid means the batteries are fully charged. If the light is off, then the batteries are not charging. If the blue charge light is blinking the system is charging normally. If the system is not charging and is plugged into an

outlet, please check if the wall power is active, to avoid permanent damage to the batteries.

The battery system is designed to be replaced by authorized and trained NeuroLogica service personnel, *only*.



CAUTION The system can only be charged from a correctly rated wall outlet. A rating information plate is located on the product-marking plate (lower backside panel or lower left side panel).

CAUTION The system should be plugged in at all times when not in **Transport** mode, being transported or in scanning use, to help maintain battery life and proper system operation. Failure to do so could result in permanent battery damage, which will require a service technician to repair.

- **CAUTION** The system may not complete a scan when below 25% battery capacity while unplugged.
- **CAUTION** If the system is unplugged and battery capacity drops to 0%, permanent battery damage can occur.
- **CAUTION** The system can be charged only from a correctly rated wall outlet.

CAUTION The power cord selection must not be less than 220v/16A, made of 2.08mm (diameter) copper wire in accordance with local power supply cable standards.

Note: Medical grade power cords should be used at all times.

Battery component for the base power distribution assembly

- **Note:** All battery safety warning and charging methods are similar to the direction respectively noted for the battery system in the above section: Battery system safety and information of this document.
- **WARNING** Proper disposal of batteries is required to ensure compliance with environmental safety guidelines. Contact authorized NeuroLogica representative for instructions.

Hazard Information

Maintenance and service

WARNING	Equipment maintenance of non-medical electrical equipment should not be performed in the patient environment.
WARNING	Maintenance checks and all service must be performed by NeuroLogica trained technicians. Service personnel use Service manual (1-NL5100-062) to effectively perform needed service and preventive maintenance and inspection on the system. See "Contact information" for NeuroLogica's contact information.
WARNING	The only calibration performed by the user on this system is called daily calibration and is described in detail later in this user manual. All other calibration needs that arise must be performed by trained technicians at NeuroLogica Corp. See "Contact information" for NeuroLogica's contact information.
CAUTION	Service personnel must complete training at NeuroLogica Corp. for the system and its accessories prior to conducting any service activities.
	Users are not to perform service or maintenance on the system at any time. This includes battery maintenance.
Note: N	euroLogica recommends that a six-month preventive maintenance be onducted by NeuroLogica's service personnel/trained facility bioengineer.
Ν	euroLogica recommends a semi-annual service contract.
ln (1	nstructions for replacing serviceable parts are identified in the Service Manual I-NL5100-062).

Cleaning the system

Keep the equipment clean. Remove body fluids to prevent health risk and damage to internal parts.

Note: NeuroLogica recommends a solution of ≥99% pure Isopropyl Alcohol (IPA) to sufficiently clean the equipment.

When the system is between uses, NeuroLogica recommends keeping it clean as described below. This will help remove body fluids to prevent health risk and damage to internal parts.



WARNING Do not use flammable or potentially explosive disinfecting sprays, since resultant vapor could ignite, causing personal injury and/or damage to the equipment.

WARNING In order to prevent short-circuiting or possible electrical shock, do not spray cleaning agents or spill liquid cleaning agents directly onto the machine.

WARNING Always electrically isolate this equipment from the main electrical supply before cleaning and disinfecting it to prevent short-circuiting or possible electrical shock.

CAUTION The unit surfaces may be cleaned with a soft cloth and the recommended solution or a similar mild non-abrasive cleaning solution. General purpose liquid disinfectant may also be used as necessary. Apply the cleaning solution to the cloth, not directly to the unit.

Cybersecurity

WARNING Upon detection of a cybersecurity threat to the system or **Tablet**, do the following:

- Immediately contact Technical Support.
- Discontinue use of system (enabling the EMERGENCY-STOP if needed).
- Remove any Ethernet and/or wireless connection that has been made with the facilities' IT-network.

Continued use of the system can occur after Technical Support has assessed the situation and provided the "go-ahead" to do so.

Note: NeuroLogica Corp. recommends the customer facility utilize an IT-network that provides sufficient means of cybersecurity control to help maintain the requirements of HIPAA.

Contraindication(s)

There are no contraindications associated with CT x-ray scanning.

Personnel privileges and terminology

Qualified operator

The operator (for example, technologist, radiologist, and other professionals that are authorized to handle x-ray equipment) as determined by the healthcare facility and assigned by a user with administrative privileges – who by their education, certification, experience, and training, are sufficiently qualified to competently perform clinical scans

on the particular model of CT system which they are to use. See "Understanding the types of users" on page 91 for a description of the types of users.

Operator of record

The operator of record is an operator or health care professional currently logged onto the CT system with a unique username and password identifier.

Scanning privileges

Scanning privileges are granted to a qualified operator, assigned by a user with administrative privileges, to conduct clinical scans on the particular model CT system which they are to use. This privilege level allows use of all clinical protocols to properly scan the patient.

Protocol privileges

Protocol privileges are granted to a qualified healthcare professional (for example, radiologist, technologist, physicist), as determined by the healthcare facility and assigned to users with administrative privileges, who by their education, certification, experience, and training, is sufficiently qualified to competently save clinical protocols (either new or modified) on the particular model CT system they are working on. A healthcare professional with protocol privileges does not necessarily have to have scanning privileges on the particular CT system.

Administrator privileges

Administrator privileges are granted to qualified healthcare professionals (for example, radiologist, technologist, physicist, department administrator, etc.) as determined by the healthcare facility who by their education, certification, experience, and training, is sufficiently qualified to competently assign, maintain, and oversee the assignments of personnel to scanning privileges and/or protocol privileges on the particular CT system which they administer. In addition, qualified healthcare professionals are authorized and qualified to pull system logs associated with this standard for Quality Assurance review. Healthcare professionals with administrative privileges do not necessarily have to have scanning privileges or protocol privileges on the particular CT system.

Clinical operation

CT system operation that involves scanning live humans and/or creating or editing protocols intended for use on live humans.

Clinical scanning

CT system operation that involves scanning of live humans.

Clinical protocol

A protocol on the system intended for use on live humans.

Kernel

The kernel is defined as the reconstruction algorithm or mathematical equation used for convolution of the attenuation profiles and reconstruction of the CT images. The choice of the kernel determines the noise level and the contrast resolution of the reconstructed images.

Chapter 2 System Overview

To understand general aspects of the system, this chapter gives a brief, initial overview of the parts of the system. For example, it provides basic skills for powering on and off the scanner, familiarizing yourself with the LCD (touch screen) when the scanner is on, as well as how to locate and use **E-STOP**.

After you are familiar with basic parts of the system, you can learn how to use the scanner and **Tablet**.

The OmniTom Elite with PCD CT system lets you scan patients in a room or ward, an **Intensive Care Unit (ICU)**, an **Emergency Room (ER)**, a medical satellite facility, an **Operating Room (OR)**, and a private office or clinic.

OmniTom Elite with PCD system

The OmniTom Elite with PCD is a mobile, battery-operated CT scanner and software system with **Axial**, **Helical** and **Dynamic** capabilities. It has the following specifications:

- Standard Resolution (SR) scanning with 16 slices at 0.707mm slice thickness
- High Resolution (HR) scanning with 26 slices at 0.424mm slice thickness
- Ultra-high Resolution (UHR) scanning with 80 slices at 0.141mm slice thickness

Total scan coverage is 500mm in SR and HR and 67mm in UHR.

The OmniTom Elite with PCD core system consists of the scanner, scan board, bed adapter, QA phantom, interface cables, lead shielding curtains, and the OmniTom Elite with PCD Tablet. Consider the following:

- The Scanner and Tablet communicate using a wireless connection.
- The OmniTom Elite with PCD Tablet is a computer with custom software that allows the user to employ pre-defined system protocols or devise unique protocols for performing that user's patient studies. The viewing portion of the OmniTom Elite with PCD Tablet allows the user to view images in more detail and includes tools to help facilitate diagnosis by a physician.

Note: Although the tablet does have a high-resolution display, it is not certified as a diagnostic viewing station.

- The scanner uses a minimum slice-thickness: 0.625mm and a maximum coverage: 500mm. The maximum scout length is 500mm.
- Scan boards are headboards that support the patient's head and neck while the patient undergoes a scan or study.

Note: *It is advised* to power up the OmniTom Elite with PCD system hardware first, to allow time for the scanner to warm up, then power up the tablet once the scanner power-up has completed.

• Specially designed platforms are also available for scanning neo-nates and small pediatric patients who cannot be scanned in their own beds.

The LCD touch screen

The **Liquid Crystal Display** (LCD) is a touch screen that is located on both sides of the scanner. The touch screen displays patient information, machine-positioning buttons for a scan, and scanner-status information.

The touch screen contains three tabs that show different information, actions, options, and buttons which include: the **Acquisition**, **Positioning**, and **Transport** tabs. To make a tab active, and see what actions you can perform from the tab's panel, press the tab. The tabs are described later in this user manual. See "Overview of the scanner's LCD touch screen" on page 69.



Figure 6: Acquisition, positioning, and transport tabs on the LCD touch screen

The shielding curtains

The shielding curtains decrease radiation to those around the patient.

To learn more about the US and European findings regarding the scatter radiation measurements with open curtains, no curtains, or closed curtains, see "Scatter radiation" on page 163. To learn how to use the shielding curtains, see "Using the curtains for shielding" on page 88.

The tablet

The **Tablet** is an accompanying part of each scanner; it is the control unit that operates all respective functions of the system.

The **Tablet** is connected wirelessly to the scanner. All basic information related to the Tablet (for example, operating distance, warnings and cautions, connectivity, functionality, etc.) appears in Chapter 4 Basic Tablet Operations.



Figure 7: OmniTom Elite with PCD tablet

Table 16: Tablet display dimensions

Display	
Screen	12.3" PixelSenseTM Display
Resolution	2736 X 1824 (267 PPI)
Aspect Ratio	3:2
Touch	10-point multi-touch

Note: Although the tablet does have a high-resolution display, it is not certified as a diagnostic viewing station.



CAUTION Do not connect or use equipment beyond what is specified by NeuroLogica Corp.; this practice may lead to a reduced level of system safety.

The recommended distances, provided in Figure 8, relate *only* to distances specified by IEC 60601-1 and *do not* relate to specific distances required for ionizing radiation and/or stray radiation protection for operators and bystanders.



Figure 8: Tablet safe distance location (two views)

The silhouette scan board and universal transfer board

A scan board is supplied with the system. The silhouette scan board is always used with an adapter; the universal transfer board should always be used with mattress stiffeners.



Figure 9: Silhouette scan board and universal transfer board

Parts that potentially come into contact with the patient

While you use the system, be aware that the patient may come into contact with the following parts:

- The OmniTom Elite with PCD system, especially the painted, external surfaces of the system's scanner cover.
- Silhouette scan board
- Bed Adapter
- Patient Safety Straps
- Universal Transfer Board
- Pediatric and Neonate Scan Platforms.

Chapter 3 Basic Scanner Operations

Basic scanner skills include powering on and off the scanner, learning how to use and navigate the LCD touch screen, and how to use **E-STOP**.

Scanning basics you should know before you scan a patient include how your system should be set up, how to position the scanner and the patient before the scan, and how to perform a scan using the **Tablet**.

Powering on and off the OmniTom Elite with PCD system

The OmniTom Elite with PCD can be powered on and off with the power button on either side of the scanner. Press and release the power button to turn the scanner on. After pressing the power button allow time for the scanner to power up. To power off the OmniTom Elite with PCD press and hold the power button for approximately 5-10 seconds until the light ring around the power button blinks. The LCD will display a shutdown message which must be confirmed by tapping **OK**.

Note: The system will perform a Drive Bar Calibration upon power up. Additional information regarding the Drive Bar Calibration is discussed on page **73**.



CAUTION Do not hang, drape, lean, or otherwise have any objects in contact with the **Drive Bar** while the scanner is powering up. Doing so can cause an erroneous calibration of the driving subsystem producing erratic driving behavior.

When powering on and off the system, consider the following:

- Make sure the scanner is properly plugged in, whenever possible; be sure the outlet(s) provide the required power.
- Plugging the electrical cord into the wall charges the batteries; the batteries are the power source that allows the scanner to operate.
- When plugging in the scanner, make sure the cable lays flat on the floor to ensure the safety of hospital personnel. In addition, make sure that the floor behind the scanner is free of any obstructions or debris that could interfere with the Translate system during scanning.

Note: NeuroLogica recommends the scanner be restarted weekly.

When powering off the scanner, wait at least 30 seconds before turning the scanner back on.

Overview of the scanner's LCD touch screen

The LCD touch screen appears on both sides of the scanner. The touch screen lets you set up and activate a scan.

LCD tabs and icons

The LCD screen contains three tabs: **Acquisition**, **Positioning**, and **Transport**. To make a tab active, press the tab with your finger.



Figure 10: Acquisition tab



Figure 11: Positioning tab



Figure 12: Transport tab

Each tab contains buttons, icons, text boxes, and status indicators to perform actions.

lcon	Description
	Scanner Relative Position – Displayed on Acquisition and Positioning tabs.
	Laser – Displayed on Acquisition and Positioning tabs.
Ref.	Zero Reference position – Displayed on Acquisition and Positioning tabs.
	Raise scanner – Displayed only on Transport tab.
	All Movement – When selected will be highlighted in orange as seen in bottom picture. Scanner will move in all directions. – Displayed only on Transport tab.

Table 17. Icons	on acquisition	nositioning	and trans	nort tabe and	actions
Table 17: Icons	on acquisition,	positioning,	and transp	port tabs and	actions

lcon	Description
	Lateral Lock – When selected will be highlighted in orange as seen in bottom picture. Scanner will only move or 'strafe' in the lateral direction. – Displayed only on Transport tab.
	Forward Lock – When selected will be highlighted in orange as seen in bottom picture. Scanner will only move forward or backward. – Displayed only on Transport tab.
	Alignment Camera – When selected, activates a camera on the back of the scanner which identifies a QR code located on the bed adapter and automatically aligns the scanner to the patient's head on the scan board. – Displayed only on Transport tab.
	Lower Scanner – Displayed only on Positioning tab.
	Translate toward patient – Displayed only on Positioning tab.
	Translate away from patient – Displayed only on Positioning tab.
	Lock LCD Screen – Displayed on Acquisition, Positioning and Transport tabs.
SCAN	Scan – Only displayed after Patient Registration, Protocol Selection and pressing 'Begin' to initiate a scan.
CANCEL	Cancel – Only displayed after Patient Registration, Protocol Selection and pressing 'Begin' to initiate a scan.
83%	Air freshness level – Identifies the air calibration status. – Displayed on Acquisition, Positioning and Transport tabs.
\bigcirc	System State Orb – Identifies the system's current state. – Displayed on Acquisition, Positioning and Transport tabs.

lcon	Description
	Tube Heat Capacity indicates the remaining tube-capacity percentage available. – Displayed on Acquisition, Positioning and Transport tabs.
77%	Battery Capacity indicates the remaining battery percentage available. – Displayed on Acquisition, Positioning and Transport tabs.
93% 76%	Battery Capacity showing 80V percentage (left) and 400V percentage (right). – Displayed on Acquisition, Positioning and Transport tabs. *This icon will only be seen when the difference between the two percentages is greater than 5 percent
	Hospital Network Connectivity – Displayed on Acquisition, Positioning and Transport tabs.
	Collision Sensors Enabled – Displayed only on Transport tab.
	Collision Sensors Disabled – Displayed only on Transport tab.
	Mute Audible Alerts – Disables the audible alerts from the collision sensors. – Displayed only on Transport tab.
	Disable Collision Sensors – Disables all Collision Sensors, as shown in bottom picture. When this mode is activated the system speed is reduced to approximately 35% of full speed in the forward direction. – Displayed only on Transport tab.
	Radiation Icon Identifies X-ray as on or off. – Displayed on Acquisition, Positioning and Transport tabs.
	Emergency Stop (E-Stop) – Displayed on Acquisition, Positioning and Transport tabs.
lcon	Description
------	--
	Remote Support– When enabled, as seen in bottom picture, allows a NeuroLogica technical support representative to troubleshoot the system remotely. – Displayed on Acquisition, Positioning and Transport tabs.
	Engineering Screen – Displayed on Acquisition, Positioning and Transport tabs.

Patient information

The patient information that appears on the touch screen is there to help you confirm that the correct patient information was selected and will appear on the scan you perform. To learn more about how to enter patient information, see "Registering the patient" on page 170.

Drive bar calibration for transport

Upon power up, the system will perform a required **drive bar** calibration. The OmniTom Elite with PCD **LCD** displays the following screen during the start-up sequence:



Figure 13: Initial screen when drive bar calibration begins

If the **Drive Bar Calibration** fails on the first attempt, the **LCD** will display the following screen to remind the user not to touch the drive bar during calibration:



Figure 14: Retry calibration screen

If the **Drive Bar Calibration** fails again, the **LCD** will display the following screen and Customer Service should be contacted:



Figure 15: Drive bar calibration failed

`

CAUTION Do not hang, drape, lean, or otherwise have any objects in contact with the **Drive Bar** while the scanner is powering up. Doing so can cause an erroneous calibration of the driving subsystem producing erratic driving behavior.

Moving the scanner for transport

To move the scanner, the scanner must be powered on and in Transport mode. The **DOWN** icons are located on both sides of the scanner on the LCD screen on the Positioning tab. The **UP** icon is located on the LCD screen on the side with the **drive bar** on the Transport tab. To set the scanner in **Transport** mode, go to the side of the scanner with the drive bar and follow the procedure below.

CAUTION Before transporting the scanner, verify that the Ethernet cable (if wired to PACS) is unplugged from the scanner to avoid damage to cable and receptacle. Verify that the power cable is unplugged from the wall to avoid damage to the cord and outlet.

CAUTION When transporting the scanner, it is recommended that two trained staff members be in control of the system to avoid any potential collisions.

1. Select the Transport tab on the LCD screen.

83% 70%
((III-

Figure 16: Active transport tab on LCD

- 2. Press and hold **UP arrow** to raise the scanner for transport until the **scanner reaches** the transport state. You will know you are in Transport Mode when:
 - The **Mood Ring** on the front of the scanner alternates flashing light blue and dark blue.
 - The System Status Orb is light blue.
 - The **Up** arrow is greyed out.
 - The LCD opposite the Drive Bar side displays "Transport Mode".
 - The Entering Drive View pop-up box will appear.



Figure 17: Transport mode

Note: The **drive bar** is located on the right side of the scanner and can transport in lateral, diagonal, and 360-degree movements.

The **drive bar** is equipped with an **enable bar**. The **enable bar** is a darker material located on front of **drive bar**. While holding the **drive bar**, squeeze the **enable bar** to activate the system to move (transport). If you let go of the **enable bar**, the scanner stops.

- 3. Once you are in **Transport Mode**, use the Drive Bar to maneuver the scanner to the patient's location. The following points explain how to hold the **drive bar** to move it forward, backward, lateral (left and right), diagonal:
 - Push forward with both hands on the **drive bar** and use equal pressure on the **enable bar** to move the scanner forward. You can also use the **Forward Lock** icon on the LCD screen to force the system to only move in the forward direction.
 - Pull back with both hands on the **drive bar** and use equal pressure on the **enable bar** to move the scanner in reverse.
 - To perform a left lateral movement, select the Lateral Lock icon on the LCD screen, squeeze the **enable bar** and tilt the drive bar to the left.
 - To perform a right lateral movement, select the Lateral Lock icon on the LCD screen, squeeze the **enable bar** and tilt the drive bar to the right.
 - Squeeze the **enable bar** and apply pressure to either the right or left side of the **drive bar** to travel diagonally in that direction.













Lateral Right

Figure 18: Drive bar motions

Note: A three-point driving technique is required to turn in smaller spaces (for example, narrow corridors).

C/

CAUTION If a loss of control is encountered while moving the system, release the **enable bar** to stop **ALL** movement.

Using collision avoidance sensors

The scanner has **Collision avoidance sensors** that will alarm when in close proximity of an object when in **Transport** mode.

The **collision avoidance sensors** warn the user visually and audibly of upcoming obstacles, allowing the user to avoid incidents.

To ensure the Collision avoidance sensors are enabled:

- Put the Scanner in **Transport** mode.
- Select the Transport Tab on the LCD.
- The collision avoidance sensors will be illuminated when enabled.



Figure 19: Enabled collision avoidance sensors

When coming in close proximity with a large object, an audible alarm will sound, and the **collision avoidance sensors** will visually change from green to red on the LCD.

The scanner will also decrease the speed of transport and gradually stop the scanner from continuing to go forward if an obstacle is in the way.

To disable the collision avoidance sensors:

• Select the **Disable Collision Sensors** Icon, located on the **Transport** Tab on the **LCD**.



Figure 20: Disable collision sensor icon

You can also mute the audible alarm without disabling the Collision Avoidance Sensors.

To mute the Audible Alerts:

• Select the Mute Audible Alerts Icon, located on the Transport tab on the LCD.



Figure 21: Mute audible alerts icon

Transporting with drive view

The scanner has an optional Drive View camera mounted on the front of the scanner which connects to the tablet to allow you to see what is in front of the scanner while driving.





Figure 22: Optional drive view camera

To enable **Drive View** the scanner must be in Transport mode.

1. Select System.



Figure 23: System icon

2. Select System Settings.



Figure 24: System settings icon

3. In System Settings select Drive View.



Figure 25: System settings

Once enabled the tablet will display a forward-looking view as demonstrated below:



Figure 26: Drive view

You can use the **CLOSE** option on the top right corner of the screen to disable the Drive View on the tablet.

Drive View can also automatically be enabled when the scanner is raised from Scan Mode into **Transport** mode. When the scanner is fully raised, the **Entering Drive View** pop-up box will appear on the tablet. Select **Enter** to allow the camera to connect to the tablet.



Figure 27: Entering drive view pop-up



WARNING When transporting, use Drive View and the Collision Sensors as guides to avoid hitting objects.

WARNING When using Drive View the camera will not show the sides of the scanner, only what is directly in front of the system. Care should be taken to avoid collisions with objects on the sides of the scanner when going through doorways or tight spaces.

Transporting with mood ring lights

The scanner has a **Mood Ring Light** on the front of the gantry that can be disabled during **Transport** and **Scan** Modes by the operator.



Figure 28: Front of OmniTom Elite with PCD with mood ring light enabled

To disable the **Mood Ring Light,** tap the **System Status Orb** located on the LCD in the upper, right corner.

To enable, tap the **System Status Orb** again.



Figure 29: Mood ring light enable/disable feature

Positioning the scanner before a scan

1. Select the **Positioning** tab on either side of the scanner on the LCD.



Figure 30: Active positioning tab on LCD

- 2. Press and hold the **DOWN** arrow until it lowers the scanner to **Scan Mode**. You will know you are in **Scan Mode** when:
 - The System Status Orb changes to the blue IDLE state.
 - The Mood Ring on the front of the scanner is also in the blue IDLE state.
 - The **DOWN** arrow greys out when the scanner is completely lowered into **Scan Mode**.
- 3. Use the positioning **LEFT** and **RIGHT** arrows to align the patient and bed with the scanner, ensuring the patient is in the center of the **Field Of View (FOV)**.
- 4. Align the patient bed with scanner opening to prevent a collision with patient, patient support and/or any life supporting devices prior to scanning.

Note: Be sure the floor is clear of debris or anything that can cause interference with the scanner's translate wheels.

CAUTION Ensure there is adequate clearance for the scanner to move the prescribed distance during the scout and/or scan.

5. Make sure the patient's bed is locked.



- **CAUTION** Make sure to lock the patient bed or scan table to prevent it from moving during the scan.
- 6. Adjust the bed height so that the patient is centered within the bore.

Positioning the patient

WARNING	Prior to scanning, properly position the patient to ensure that
	extremities, hair, life support equipment, and any other in proximity
	of the scanner have sufficient clearance to prevent patient injury
	with the scanner itself and/or when used with accessories/options.

- **WARNING** Ensure the patient support is properly positioned (height and alignment) to prevent injury during scanning.
- **WARNING** Make sure the foot pedal brake on the patient support/bed is engaged to prevent it from moving during the scan.
- WARNING Never raise or lower the scanner when a patient is positioned in the system's bore. *Always* move the scanner away from the patient support *before* raising or lowering the system itself.



CAUTION The following-instructions for patient positioning should be performed in accordance with NeuroLogica Corp.'s clinical training.

Note: To determine where personnel should stand during a scan, consult with the hospital physicist. NeuroLogica recommends a distance of 8-10 feet.

1. Attach bed adapter to the bed.



Figure 31: Bed adapter without posts insertion



Figure 32: Bed adapter with T-square

3. Tighten the Bed Adapter to the scan board by turning the T-square handle until two (2) clicks are heard.

2. Attach the silhouette scan board to the bed adapter by inserting the scan board into



Figure 33: Attaching the scan board with bed adapter to patient's bed

4. Position the patient until the patient's head rests on the pad of the scan board as shown below.



Figure 34: Positioning the patient on the bed

5. Position the patient in front of the scanner opening. The patient's shoulders should rest flat against the face of the device.



• Make sure the patient is centered in the scanner bore.

Figure 35: Ensuring placement of patient's neck directly under laser light

6. Lock the gurney or bed wheels before proceeding with scan.

Using alignment camera

The OmniTom Elite with PCD scanner includes an automatic Alignment Camera that can be used to position the scanner so that it is centered on the bed adapter and scan board.

To enable the Alignment Camera:

- 1. Attach the Bed Adapter and Scan Board to the patient's bed.
- 2. Drive the scanner near the patient's bed so the alignment camera on the front of the scanner can 'see' the Alignment markers attached to the silhouette scan board.



Figure 36: OmniTom Elite with PCD alignment camera



Figure 37: QR code alignment indicators





Figure 38: Adult scan board

3. While the scanner is in Transport mode, and near the patient's bed, select the **Align** icon on the LCD.



Figure 39: Align icon

4. Once selected the camera will look for the Alignment markers on the scan board.

SCANNER ALIGNMENT Looking for markers
STOP

Figure 40: Looking for alignment markers

5. If the alignment markers are found, the following screen will appear.



Figure 41: Alignment found

6. If the markers are not found there may be something blocking the camera view or the scanner is not within the proper envelope, and the following screen will appear. If the markers are not found, check for proper placement of the scanner in the alignment envelope and obstructions blocking the camera view and select RETRY.

SCANNER ALIGNMENT
Markers not found. Retry?

Figure 42: Alignment not found



7. When the Alignment markers are found select **Move** which will show the following:

Figure 43: Alignment moving

8. When the scanner is in the proper location the LCD will indicate Alignment is complete.

SCANNER	ALIGNMENT
Move C	omplete!
STOP	DONE

Figure 44: Alignment complete

9. You can now move the patient onto the scan board and into position for your scan.

WARNING	Always align scanner to scan board before positioning the patient on the scan board.
WARNING	The Alignment camera can only be used on the adult silthoutte scan board. It is not currently available with the neo-nate or pediatric platrform.
WARNING	Please note that the collision sensors are not active when using the Alignment camera.
WARNING	If the scanner moves unexpectedly while aligining, press the E-stop to halt all motion.

Positioning the scanner using the laser light

The scanner provides a laser light to guide you to properly position the patient. The lasers indicate the center of the scan or Transverse plane, the mid-Sagittal plane, and the vertical height or Coronal plane. The laser light is centered on the actual x-ray beam at all times. For multi-slice protocols, this means that the laser light will indicate the middle position of all simultaneous scans being acquired. The accuracy of the position of the laser plane, with respect to the scan plane, is +/- 2mm.

To activate the laser light on the touch screen

- 1. Press the **Laser** button on the LCD screen.
- 2. The laser will automatically shut off, 30 seconds after pressing the **Laser** button. See the precautions regarding the laser in "Laser safety" on page 51.

If the patient is conscious, request the patient remain still with eyes closed throughout the entire scan.

If the patient is unconscious, secure the patient.

Follow the appropriate facility guidelines when scanning unconscious patients if the patient's eyes remain open.

Using the curtains for shielding

Using shielding curtains is recommended when performing equipment calibrations and patient scans, to ensure maximum efficiency and patient safety.

- 1. Pull up the back curtain ensuring that the curtains lie flat against the back of the scanner.
- 2. Check to ensure that the patient is properly positioned and comfortable.
- 3. Position the front curtains as close to the patient as possible to minimize the space between them.
- 4. Before scanning the patient, check to ensure that nothing interferes with the patient's life support or other external medical devices.

Note: For calibrations, the curtains should be completely closed against the front of the scanner.

The (shielding) front curtains minimum thickness is 0.5mm, the rear curtain is 0.25mm.

Operating the E-STOP button

- 1. Press the **E-STOP** button to perform the following:
 - Stop the system (if it loses control).
 - Stop all system motion and x-ray.

- Remove power to the gantry drives and x-ray system.
- If the OmniTom Elite with PCD system starts to move unexpectedly.
- 2. Make sure to resolve the situation.

Note: When E-STOP is activated, the moving gantry may overrun by less than 10mm.

Restoring the system from E-STOP

If you have pressed the **E-STOP** button to stop the system, follow these steps to restore the system using **E-STOP**.

- 1. Make sure any hazard is removed.
- 2. Twist the E-STOP button clockwise until the button pops out to restore the system.

Rebooting the system

If your scanner needs to be rebooted press the power button for 5 seconds or until the button begins to flash, then release the button. The scanner will display a prompt asking if you would like to power down. Press 'OK' to initiate the Power down sequence.



Figure 45: Power down pop-up

To restart the scanner, press the power button once. Allow a few minutes for the scanner to restart.

You will know your scanner is fully powered on when the **Mood Ring** and **System Status Orbs** change from grey (Powerup) to Idle Blue (**Scan Mode**) or alternates flashing blue (**Transport Mode**), depending on which mode the scanner was left in before rebooting.

Remote support feature

The Remote Support feature allows a NeuroLogica technical support representative to troubleshoot the system remotely, perform system updates, and transfer files from the

system for use in diagnosing issues. When enabled, Remote Support allows NeuroLogica personnel to take remote control of the system while you observe.

Enabling remote support

1. Select the Enable Remote Support icon on the LCD.



Figure 46: Enable remote support

2. Select OK on the Enable Remote Support Prompt.

Enable remote support?

Figure 47: Enable remote support prompt

3. When the remote support session is completed, select the **Disable Remote Support** icon on the LCD.



Figure 48: Disable remote support

4. Confirm that you want to end the Remote Support Session on the **Disable Remote Support Prompt.**

Disable remote support?	
CANCEL	

Figure 49: Disable remote support prompt

Chapter 4 Basic Tablet Operations

Basic Tablet information includes understanding the different kinds of user access, learning how to power on and off the Tablet, getting familiar with the Tablet screen, menus, and tabs.

Note: *It is advised* to power up the OmniTom Elite with PCD system hardware first, to allow time for the scanner to warm up, then power up the tablet once the scanner power-up has completed.

Understanding the types of users

There are three **User Levels** available on the Tablet: administrator, limited operator, and restricted operator. Usernames and passwords can be created for individual users, and specific User Levels can be assigned to each user. The following define the access levels for each User Level:

	Full access to the system and its configuration. Can create	
Administrator	protocols, user names and passwords, as well as access all	
	functions of the system.	
	Modified access to the system. Users with Limited access can	
Limited	modify protocols during scanning but cannot create and save	
	protocols; has no access to system configuration.	
	Users with Restricted access can scan with the system but are	
Restricted	unable to make any changes to protocol parameters while	
	scanning, they also have no access to system configuration.	

Powering on the tablet

- 1. After the scanner has completed its power-up sequence.
- Press the **Power on** button on the Tablet. The Tablet will boot up and the **Login** panel appears.



Figure 50: Login panel

Note: NeuroLogica recommends the tablet be restarted daily.

Logging in to the tablet

To gain access to the Tablet application, you must provide the system with credentials. These credentials consist of a **user identification** (**ID**) and **password**. Make sure you have a valid user ID and password before you log into the system.

Note: If you do not have a valid user ID and password, ask your supervisor or administrator for one.

- 1. Select your User ID from the dropdown under 'Select'
- 2. Click in the **Password** field and type your password in the field. Passwords are case sensitive.
- 3. Select the Tap to Login icon.



Figure 51: Tap to logIn prompt

If the user ID and/or password are invalid, a prompt appears denying access. Following the prompt, you are able to retry logging in with the correct credentials.



Figure 52: Access denied prompt

Note: You have a limited number of login attempts before the system locks the account. An administrator can unlock the account.

- 4. When the user ID and password are verified, you are logged into the system.
- 5. The Tablet software appears with the **Registration** tab active.



Figure 53: Registration tab

Locking the tablet with the LCD

Lock the tablet to guard your work while you are away from the OmniTom Elite with PCD. It is important to lock out unwanted users even if you are away for only a short period of time.

When you lock the scanner, it will automatically lock the tablet. Although both the tablet and scanner will remain on, no one can access the tablet without your username and password.

The **Lock button** located on the LCD indicates whether your system is in lock or unlocked mode.

In addition, you can:

- Log off the tablet
- Enable the privacy screen.

Note: The tablet will auto lock after 15 minutes of inactivity and will require the user to input their username and password to unlock.



Figure 54: Automatic lock out screen

Press the **Lock** button located at bottom of the LCD.

Posteric Information Protocol Information POSITIONING Retient: Information It4367 Retient: Init Petient: Di: 14367 Aritegh Young MrA: 20 Aritegh Young Aritegh Young Accession #: 51649668 DOSE Context CTD/vol: Dup: Coverage: 120	ACQUISITION		2.5	Ref.	
TRANSPORT Accession #: 5184966 Position: HFS COSE Common Type: AXIAL CTDMot D:P. Coverage: 120	POSITIONING	Patient Information Patient ID: 14387 Patient Name: Aeliairin Xana	Protocol Infor kV: mA:	rmation 120 20	83%
CTDNot Scen Type: AXIAL DLP: Coverage: 120	TRANSPORT	Accession #: 5164966 DOSE	Thickness: Position: Kernal:	5 HFS 1	77%
Scan Delay: 10 Start Pos: 10		CTDIvol: DLP: Scan Delay: 10	Scan Type: Coverage: Start Pos:	AXIAL 120 10	98%

Figure 55: System lock button

To unlock, input **3644** and press **enter** on the LCD keypad and both the scanner, and the tablet will unlock.



Figure 56: Login screen on LCD

Logging off the tablet

1. Select the System Icon.



Figure 57: System icon

2. Select the Logout Icon.



Figure 58: Logout icon

3. A prompt will display asking to **Confirm** or **Cancel** the logout.



Figure 59: Confirm logout prompt

4. Select **Confirm** to logout.

Powering off the tablet

1. Select the System Icon.



Figure 60: System icon

2. Select the **Shutdown Icon**.



Figure 61: Shutdown icon

3. A prompt will display asking to **Confirm** or **Cancel** the shutdown.



Figure 62: Confirm shutdown prompt

4. Select **Confirm** to shut down.

Enabling the privacy screen on the tablet

1. Select the System Icon.



Figure 63: System icon

2. Select the Privacy Screen Icon.



Figure 64: System screen

The Privacy Screen will appear.



Figure 65: Privacy screen

3. To leave the **Privacy Screen**, tap below the NeuroLogica Brain Logo, and then above the NeuroLogica Brain Logo.

Navigating around the tablet's main screen

The Tablet screen shows similar information, regardless of the screen elements change when you perform different actions. These constant screen elements are as follows:

Constant Screen Elements	Element	Element description
	Patient Registered Icon	Appears in the top left corner of the screen and when tapped displays the currently registered patient's information.
	Protocol Selected Icon	Appears in the bottom left corner of the screen and when tapped displays the current active protocol parameters.
	System Icon	Appears in the bottom right corner of the screen and when tapped displays scanner configurations and calibrations.

Table 18: Constant screen elements

Getting to know the scanner status screen

The **Scanner Status** screen appears once the **Scanner Status** Icon is selected. The status screen gives you a quick view of the current system status. The following identifies the **Scanner Status** screen:



Figure 66: Scanner status screen

Status bar icon	Status bar icon name	Status description
System Battery 100%	System battery capacity status	Indicates the remaining scanner battery percentage available. The capacity values are color coded as follows: Green 100% - 51% Yellow 50% - 25% Red 24% - 0%
Tablet Batteny 58%	Tablet battery capacity status	Indicates the remaining tablet battery percentage available. The capacity values are color coded as follows: Green 100% - 21% Yellow 20% - 11% Red 10% - 0% You will be prompted to plug the Tablet into an outlet to charge if the battery capacity is low; a scan cannot complete when the battery capacity is 10% (red) or lower. When the Tablet reaches the red capacity range it will shut down. A message appears informing the operator that the tablet will shut down due to a low battery. The lightning bolt icon signifies that the tablet is currently charging and goes away when unplugged
Daly Calbratic O%	Daily Calibration status	 Indicates the air freshness status; It is recommended that an air calibration be performed: Every six (6) hours. When the air freshness status falls below 50%. If the scanner is moved to an area with a dramatic change in humidity and or temperature. The calibration status values are color coded as follows: Green 100% - 51% Yellow 50% - 25% Orange 24% - 0%

Table 19: Status bar identification

Status bar icon	Status bar icon name	Status description
	System tube heat capacity status	Indicates the remaining tube-capacity percentage available. The capacity values are color coded as follows: Green 100% - 51% Yellow 50% - 15% Red 14% - 0%
	Radiation status	Identifies x-ray as on or off. The icon changes from a gray/black icon (when x-ray is off) to an animated (rotating) yellow/black icon when x-ray is on.
Patient Storage 63%	Image storage space status	Indicates the available disk space on the system for image storage. The available space values are color coded as follows: Green 100% - 51% Yellow 50% - 20% Red 19% - 0%
Hospital Network CONNECTED Hospital Network NOT CONNECTED	Wireless signal indicator	Indicates the scanner's connection to the Hospital Network.
Emergency Step OFF	System E- STOP status	Identifies when E-STOP is engaged. The icon will flash when E-STOP is pressed.
System State	System state	Identifies the system's current state. The orb changes color depending on the state the system is in. In addition, the mood ring located on the front of the scanner changes color to reflect the system's state. See Table 20, for a list of the different orb colors and system states they identify.
C Scencer Position	Scanner position	Identifies the system's current position relative to its zero reference.

System status orbs and mood ring lights

The system changes states as it performs different actions. The following table indicates what state the system is in and the colored orb that correlates to that state.

In addition, the mood ring light located on the front of the scanner also correlates to the state the system is in.



Figure 67: Mood ring light

Table 20: System state orbs and corresponding mood ring light colors

Orb	Color	State
	Dark gray	The system is powering up.
	Light gray	The system is powering down.
	Dark purple	The system is busy.
	Blue	The system is idle.
	Light Blue	The system is in transport mode.
	Green	The system is ready to perform a scan.
	White	The system is performing an air calibration.
	Yellow	The system is preparing.
	Amber	The system is scanning.
	Pink	The system is not ready.
	Red	The system is in fault.

Getting to know the system screen

The **System Screen** appears once the **System Icon** is selected. The **System Screen** contains options that can execute various commands from the Tablet and contains system configurations.



Figure 68: System screen

Table 21: System options	Table	m options	System
--------------------------	-------	-----------	--------

System bar icon	System bar icon name	Status description
	Daily Calibration	Executes Daily Calibration.
	Quality Assurance	Executes Quality Assurance Test.
	PACS Queue	Displays status of studies being archived to PACS.
	Protocol Manager	Allows user with Administrator rights to create, modify or delete protocols.

System bar icon	System bar icon name	Status description
$\langle \mathbf{i} \rangle$	About	Display's Software Version Information.
$\langle \bigcirc \rangle$	Shutdown	Shutdown Tablet
	Logout	Logs user off
<	Privacy Screen	Allows you to disable access to the tablet by covering it with the NeuroLogica Brain Logo.
	Audio	Turns sound on or off.
$\langle \not \sim \rangle$	System Settings	Contains System Configurations. See Chapter 5 System Settings for System Configuration options.

Getting to know the protocol screen

The **Protocol Screen** is available after a patient has been registered, the protocol has been selected and the **Protocol Selected Icon** is selected. The **Protocol Screen** contains the following information: protocol parameters, dose, and scan time.

- The **Protocol Screen** can be used to double check parameters before making an exposure.
- In case you need to make changes to your protocol before scanning, close the **Protocol Screen** by deselecting the **Protocol Selected Icon**. Then **Cancel** from the **Acquisition** Tab or **Cancel** from the LCD.

odv Parts:	Head	Step and Shoot:	False
can Type:	Scout	Scan Delay:	
cout Type:	Lateral	Enable AEC:	False
v: 0,40,40,40,40,40	100	Noise Level (1):	
	10	Min mA:	
lice Thickness:		Max mA:	
lice Spacing Placeholder:	-5	Window Width:	100
ernel:		Window Level:	-1000
otations:		Tube Angle:	
itch:	3	Orientation:	HFS
tart Position:	0	Max CTD:	1000
nd Position:	250	Max DLP:	2000
can Time:	0		
TDivol:	5.64		

Figure 69: Protocol screen

Getting to know the patient screen

The **Patient Screen** is available once a patient has been registered and after the **Patient Registered Icon** is selected. The **Patient Screen** displays patient demographics: ID, Accession, Study Date and Time, Date of Birth, and Sex.

• The **Patient Screen** is where **Finalize** and **Edit** are located.

	PAT	IENT	
ID:		DATE OF BIRTH:	05/04/1969
ACCESSION:	98214534	SEX:	
STUDY DATE:	01/03/2019		
STUDY TIME:	11:47:49		
NAME:	McMurphy, Randl		

• To finalize a study, select the **Finalize** button.

Figure 70: Patient screen



The tablet tabs

To perform a patient examination, you will use the following five tabs on the Tablet:

REGISTRATION ACQUISITION RECONSTRUCTION VIEWER BROWSER

Figure 71: Tablet tabs to perform a patient examination

The tabs include active tabs which will be blue if they are selected or white if they are active but not selected, and inactive tabs which will be gray. The active tabs are **Registration**, **Reconstruction**, and **Browser**. The **Acquisition** and **Viewer** tabs require additional steps to be performed before they become active. The following actions are available in each tab:

Pogistration	Allows you to register a patient either manually or from the
Registration	hospital's database sites.
Acquisition	Allows you to select a protocol and perform the examination.
Acquisition	This tab is inactive until a patient is registered.
Percentruction	Allows you to manipulate raw data in different parameters and
Reconstruction	settings after your scan is completed.
Viewer	Allows you to view patient images. This tab is inactive until a
viewer	study is loaded from the Browser.
Browcor	Allows you to view, manipulate, and archive scans already
browser	performed.

In the following chapters, you will learn how to perform necessary steps to conduct a patient examination and learn how to manipulate and store the data you acquire.

Tablet buttons

Table 22: Registration tab buttons and functions

Tablet button	Action
GUERY	Searches the HIS/RIS server for scheduled patients. The population of patients could take several minutes to appear, depending on the number of patient entries found from the HIS/RIS query.
SEARCH	Searches queried patient entries for specific information.
MANUAL	Allows you to manually enter a new patient but does not include Patient ID information.

Tablet button	Action
	Clears search information entered when using the Search function.
WORKLIST	Displays HIS/RIS query results.
STORED LIST	Allows you to select patient(s) from query results and move them into the Stored Results list. All Manually entered patient information will default to this list.
	Allows you to manually enter a new patient including the Patient ID information.
REGISTER	Registers the selected patient and then takes you to the Acquisition tab to select a protocol to be used for scanning.
FINALIZE	Found under the Patient Registered Icon and allows you to complete the examination. Finalizing the Exam completes all protocols, builds dose SR and images, and re-opens the Registration tab.

Table 23: Acquisition tab buttons and functions

Tablet button	Action
CONTINUE	After protocol selection, authorizes the scanner to move to the next step.
PAUSE	Pauses the current exposure within an Axial Scan. This is a toggle button with the Resume button.
RESUME	Resumes a paused series of scans within an Axial Scan. This is a toggle button with the Pause button.

Tablet button	Action
REPEAT	Repeats the last scan that was performed.
INITIATE EXPOSURE	Manually initiates x-ray exposure when using the optional Step & Shoot feature.
RESUME ALL	Disable's Initiate Exposure and scans remainder of planned axial scans when using the optional Step & Shoot feature.
MANUAL START	Used to stop a Dynamic CTA scan and move to the Helical CTA acquisition when using Bolus Tracking and the threshold in the ROI is not crossed.
ACCEPT	Used to accept protocol selection for exam.
BEGIN	Used to begin a scan or series of scans after accepting the protocol.
CANCEL	Cancels the current scan within a protocol.
PROTOCOL	Allows user to modify existing protocol selected in exam or choose a new protocol.
Chapter 5 System Settings

System settings overview

A user with Administrative privileges must set up the OmniTom Elite with PCD system configurations for other users (limited and restricted operators). System configuration, overall, is how the OmniTom Elite with PCD scanner is set up to meet site-specific needs. Most windows contain self-explanatory instructions and refer to elements that are known to the administrative user with radiological education and training. Additionally, brief instructions are provided to aid in completing those sections with more detail.

Configuration includes setting up user permissions (or access privileges) to manage other users, as well as servers, what is available through settings, presets, and other preferences for the use of the system at a site. Many system configurations are permissible to the administrator, *only*. While other configurations are permissible to users without administrative access.

Note: You must have administrative access privileges and be logged in as an administrator to set configurations for the site.

Incorrect changes to the system configuration may make the system inoperative.

The following table shows the **System Configuration** options that appear when you click the **System Icon** (from the bottom right corner of the screen), and then select one of the system options below. The table provides a brief description of these options.

lcon	Icon Name	Description
$\langle \bullet \rangle$	Manage Users	Allows the administrator to create and edit user accounts and permissions.
	Audit Trail Viewer	Allows the administrator to view and log all changes as well as actions in the system, which include logins, patient registrations, and series updates.
$\langle \mathbf{x} \rangle$	Dose Configuration	Allows the administrator to set up dose notifications, dose alerts, and configure dose limits for specific scans.
$\langle \not \! \! / \! / \! \! / \! \! / \! \! / \! \! / \! \! / \! \! / \! \! / \! \! / \! \! / \! \! / \! \! / \! / \! \! / \! / \! \! / \! \! / \! / \! / \! \! / \! / \! \! / \! / \! / \! \! / \! / \! / \! / \! / \! \! / \! / \! \! / \! / \! / \! \! / $	Privacy Screen	Allows the user to activate the privacy screen on the tablet to hide patient information.

Table 24: System configuration settings

lcon	Icon Name	Description
$\langle \bullet \rangle$	Recon Presets	Allows the administrator to define and customize reconstruction presets.
	Recon Groups	Allows the administrator to define and customize reconstruction groups.
	MPR Presets	Allows the administrator to define and customize MPR presets.
	PACS Configuration	Allows the administrator to select a different Picture, Archiving, and Communication System (PACS) server to Archive studies.
	Window Presets	Allows the administrator to set window width and window levels.
$\langle \mathbf{Q} \rangle$	SMPTE	Allows the user to view the SMPTE Medical Diagnostic Imaging Test Pattern.
	Drive View	Opens the drive camera view on the tablet when in transport mode.

Setting user accounts

The **Administrator**, *only*, can update another user's account, add a new user, delete a user, and lock or unlock a user's access.

NEW USER	Username Enter Username	Password (8+ characters) Le Se	evel lect	
	Username	Level			
	Admin			P	
	Test 2	Restricted		P (
	Test 3	Limited			
					ß

Figure 72: User accounts dialog box

1. For the Username field, enter the user ID name.

2.	For the User Level field	, enter one of the following user levels:
		Full access permission (rights) to the system

	Full access permission (rights) to the system and its
Administrator	configuration. Can create protocols, user names and
	passwords, and all functions of the system.
	Modified access permission (rights); can modify protocols
Limited operator	during system use but cannot create and save protocols;
	has no access to system configuration.
Destricted exercise	No access to creating or deleting protocols; has no access
Restricted operator	to system configuration.

3. For the Password field, enter the user's password.

Note: The password must contain 8 to 12 characters, and must include one number, a lower-case letter, a special character, and an uppercase letter.

- 4. Select the Save button. The user is added to the list.
- 5. Press the up and down scroll bar to reorder the user list, if desired.

Deleting a user

Note: The NeuroLogica administrator account cannot be deleted.

- 1. Select the user to delete from the list of users.
- 2. Press the Delete button. The following message will display:

USERS	Username	Level			
and and a second se Second second	Admin		(P		
	Are you sure you want to delete user	"cttech1"	YES	NO	

Figure 73: Delete pop up

- 3. Select **Yes** to delete and **No** to keep user account.
- 4. Select the **Close** button to exit.

Applying dose configuration

Note:	You must have administrative privileges to access this area in the application.
	Incorrect changes to the system configuration may make the system inoperative.

Note: You can check the **Audit Trail** to review the audit log that details what dose limit was removed, by whom, and the date and time it took place.

Dose Configuration consists of both Dose Notifications and Dose Alerts (defined as System Limits in OmniTom Elite with PCD System Settings).

Dose Notification	Notifies the user when the planned CTDI _{vol} and/or DLP value of a single series will exceed the value defined value set in each protocol.
Dose Alert (System Limit)	Notifies the user when the planned CTDI _{vol} and/or DLP value from the combination of all planned series will exceed the defined value set in System Settings . Dose Alerts represent a value which would be well above an institution's established CTDI/DLP range for the given examination and warrant a more stringent review and consideration before proceeding.

Setting dose notifications

Dose Notifications are set in Protocol Manager on a series-by-series basis. Every series in a protocol can have its own notification value. Use the following steps to edit **User** protocols to define the Dose Notification value for each series. Protocols that begin with "**NL**" are NeuroLogica reference protocols and cannot be edited within Protocol Manager using the below steps, see the **Editing Dose Notifications on NL Protocols** below.

1. Select System.



Figure 74: System icon

2. Select Protocol Manager.



Figure 75: Protocol manager icon

3. Select either Adult or Pediatric.



Figure 76: Adult or pediatric selection

4. Select the icon corresponding to the appropriate body part.



Figure 77: Body part selection

5. Select the protocol you want to add Dose Notifications to.



Figure 78: Protocol selection

6. Select **Edit** to open the protocol.



Figure 79: Edit icon

7. Select the appropriate series. In this case we are going to add a notification value to the Axial series.

Protocol Description		Sc	out			A>	kial	
Brain	7,00	(1000000)		(and corts)	Tites	Scen Time	[1100]	Man D.P.(mDY.arr)
		End Paeton		Max CDP (HOY (H)	Body Plants	Bart Pasters		
Anatomical Reference	Body Parts		Scan Time		Description			
Head				GTDHolings!	tioni Ke			
				OLP (mGY(cm)				
Study Description								
Brain								

Figure 80: Series selection

8. Select the dropdown menu in the Max DLP or Max CTDI_{vol} fields



Figure 81: Max field selection

- 9. Enter the desired Maximum CTDI or DLP value
- 10. Select the **Checkmark** to save the updated value(s)
- 11. Select Update



Figure 82: Update icon

12. Select Save



Figure 83: Save icon

13. Continue Editing protocols or select **Close** to exit protocol manager.



Figure 84: Close button

If when using a protocol that has a Dose Notification turned on, the CTDI or DLP of the planned scan exceeds the Notification value, a Dose Notification message will be displayed on the tablet.

DOSE NOTIFICATION
The Axial's CTD (65.03/50) and DLP (1495/1000) exceeds its own limits.
Tap to close

Figure 85: Dose notification message

When you 'Tap to close' the above screen the following message box will appear, asking you to enter a Diagnostic Reason for exceeding the set Dose Notification value.

DOSE NOTIFICATION	
1 scan(s) within this protocol exceed their respective CTD and DLP limits. Please enter your reason for exceeding the limit(s) to continue.	
Enter reason	
CANCEL ENTER	

Figure 86: Diagnostic reason dialog box

You can either enter a reason and select Enter or select Cancel and modify the protocol parameters to reduce the CTDI or DLP below the Dose Notification Limits.

Editing dose notifications on NL protocols

The NeuroLogica (NL) Reference Protocols installed on the scanner come with default Dose Notification values set at 1000 mGy CTDIvol and 2000 mGy cm DLP values. To modify those values, you must use the Dose Configuration feature found in the System Settings.

1. Select System.



2. Select System Settings.



Figure 88: System setting icon

3. Select Dose Configuration.



Figure 89: Dose configuration icon

4. The **Dose Configuration** dialog box appears.

System Limits	ADULT	1000	2000	SET CANCE
Brotrond		CTD)/d		
NL Adult Axial Head	Scout	1000	2000	
NL Adult Axial Head	Axial	1000	2000	
NL Adult Stroke	Scout	1000	2000	SET CANCE
NL Adult Stroke	Axial	1000	2000	BET CANCE
NL Adult Head Trauma	Scout	1000	2000	
NL Adult Head Trauma	Axial	1000	2000	SET CANCE

Figure 90: Dose configuration dialog box

- 5. Select the NL Protocol Series you want to modify.
- 6. Enter the **CTDIvol (mGy)** value in the text box using the dropdown menu and select the Checkmark to save the value.
- 7. Enter the **DLP (mGy.cm)** value in the text box using the dropdown menu and select the Checkmark to save the value.

Protocol	Scan	CTDVal DLP
NL Upper Extremity	Scout	500 BET CANCEL
NL Upper Extremity	Axial	
NL Lower Extremity	Axial	
NL Lower Extremity	Scout	
NL Brain - Axial	Scout	X 500 SET CANCEL

Figure 91: Entering mGy and mGy.cm values

- 8. Select **Set** to save the new values.
- 9. Repeat steps 5 through 8 for any other protocol series you wish to modify.
- 10. When finished select **Close** to exit the Dose Configuration feature.

Setting dose alerts (system limits)

- 1. Select System.
- 2. Select System Settings.
- 3. Select Dose Configuration.
- 4. The **Dose Configuration** dialog box appears.

Note: The default Dose Alerts (System Limits) which are set at 1000mGy CTDI and 2000 mGy cm DLP are designed to prevent the patient from receiving any possible deterministic effects due to excess dose. However, the system allows these values to be modified by the user. Any modifications to the Dose Alerts (Systems Limits) should be made by qualified medical personnel.

- 5. To define the Dose Alert (System Limits):
 - Enter the **CTDIvol (mGy)** value in the text box using the dropdown menu and select the Checkmark to save the value.
 - Enter the **DLP (mGy.cm)** value in the text box using the dropdown menu and select the Checkmark to save the value.

System Limits	ADULT PEDIATRIC	CTDiVol	DLP 2000	SET CANCEL
		1 2 3 4 5 6 7 8 9 ○ X ✓ X		

Figure 92: Define dose alert

- 6. Select **Set** to save the new values.
- 7. When finished select **Close** to exit the Dose Configuration feature.

If a Dose Alert is triggered by a series of planned scans who's combined CTDI or DLP exceeds the Alert value, a Dose Alert message will be displayed on the tablet.



Figure 93: Dose alert

When you 'Tap to close' the above screen the following message box will appear, asking you to enter a password and a Diagnostic Reason for exceeding the set Dose Alert value.

The protocol's CTD (84/75) the system's limits. Please en the limit(s), <i>admin</i> usernam	ALERT and DLP (1938/1000) exceed nter your reason for exceeding e, and password to continue.	
Admin Enter reason	Enter text	
	TER	

Figure 94: Dose alert dialog box

You can either enter a reason and select Enter or select Cancel and modify the protocol parameters to reduce the CTDI or DLP below the Dose Alert Limits.

Managing DICOM servers

Digital Imaging Communication in Medicine (DICOM) is the definition of the acronym **DICOM**. **DICOM** servers are used to export images from the scanner. The **PACS Configuration** option lets the administrator access all possible storage servers created in the system by the **field service engineer**.

See **DICOM** standards on the **NEMA.org website** for a full description of settings (and actions) that are available.

DICOM servers are initially set up by the **Field Service Engineer** and the appropriate IT person at the hospital.

Note: You must have administrative privileges to access this area in the application.

- 1. Select the **System** icon.
- 2. Select the PACS Configuration. The PACS Configuration dialog box appears.



Figure 95: PACS configuration dialog box

- 3. Select the new default **PACS Server** from the dropdown.
- 4. Select Close.

Note: Any changes you make to the default server will be reset when the tablet is restarted.

Setting up the audit trail viewer

The administrator sets up the Audit Trail Viewer to build an audit trail, which monitors and/or reports changes that are made – from all users who logged in. Changes may include modifying presets or any other activities. These users include operator, administrator, or service-related users that make changes to the system.

Note: You must have administrative privileges to access this area in the application.

- 1. Select the System Icon. The System screen appears.
- 2. Select the Audit Trail Viewer icon. The Audit Trail Viewer dialog box appears.

	AUDIT TRA	AIL VIEN	NER		
D-D-D-D-D-D-D-D-P	rom To	Audit Type	User ID		
Se	elect Select	Select	Select	CLEAR	
Description	Туре	User ID	Date	Time	
User Admin has logged in.	UserLogin	Admin	06/05/2018	09:18:46	
User Admin has logged in.	UserLogin	Admin	06/05/2018	09:19:52	
User Admin has logged in.	UserLogin	Admin	06/05/2018	09:20:25	
User Admin has logged in.	UserLogin	Admin	06/05/2018	09:23:22	
User Admin has logged in.	UserLogin	Admin	06/05/2018	09:23:35	
User Admin has logged in.	UserLogin	Admin	06/05/2018	09:30:54	
User Admin has logged in.	UserLogin	Admin	06/05/2018	16:17:35	
User Test has logged in.	UserLogin	Test	06/05/2018	16:19:15	
Patient has been registered: ID 735	65624. PatientRegistr	ration Test	06/05/2018	16:39:23	
Performing the following protocol: T	ype: Scout kV: 12 PatientExamin	nation Test	06/05/2018	16:40:45	

Figure 96: Audit trail viewer dialog box

- 3. From the Audit Trail Viewer dialog box, select a date or date range with the calendar to show when changes were made.
 - To select a single date, press on the date on the calendar to find audits for that date. See the calendar in the next figure.

◄		Au	gust 20	18		►
Sun	Mon	Tue	Wed	Thu	Fri	Sat
29	30	31	1	2	3	4
5	6	7	8	9	10	11
12	13	14	15	16	17	18
19	20	21	22	23	24	25
26	27	28	29	30	31	1
2	3	4	5	6	7	8
	ACCE	PT		CA	NCEL	



• To select a span of time, select the **From** box and select a start date (on the calendar) and then select the **To** box and select the end date on the calendar. This approach lets you select a range of audits done in a specified span of time. See the text boxes to the right of the calendar; the top text box is where the start date will show when you select a start date on the calendar; the bottom text box is where the end date will show when you select an end date on the calendar.

F	rom	То	Audit Type	User ID
Se	elect	Select	Select	Select

Figure 98: Audit trail viewer options

4. Select the **Audit Type** dropdown to select the kind of audit you are searching for.

- 5. From the **User ID** dropdown, select the type of user to track.
- 6. Select the **View** button to see the result of audits that met your criteria. The results will appear.

Setting recon presets

Recon Presets allow the administrator to define and customize reconstruction presets. Recon Presets are listed in the preset dropdown menu in the Reconstruction tab.

The administrator can delete existing recon presets, update an existing recon preset, or create and save new recon presets.

- 1. Select the **System** icon. The System Screen appears.
- 2. Select System Settings icon. The System Settings screen appears.
- 3. Select the Recon Presets icon. The Recon Presets dialog box appears.

	reset	Rec auto	on presets can omatically popul	be used in ate the reco	post recon	struction to settings with a		
Name		pre- Scan Type	defined configu Thickness & Soncing	ration. Width		Multi-Energ		
2.8 Soft Tissu	e	Helical	2.8 x 2.8	135	35	None		
0.8 Soft Tissu	e HR	Axial HR	0.8 × 0.8	135	35	None	()	
0.8 Soft Tissu	e HR	Helical HR	0.8 x 0.8	135	35	None	()	
2.8 Soft Tissu keV	ie 60	Axial	2.8 x 2.8	135	35	VMI	()	
2.8 Soft Tissu keV	ie 70	Axial	2.8 x 2.8	135	35	VMI	• ×	
2.8 Soft Tissu keV	ie 80	Axial	2.8 x 2.8	135		VMI	@ X	

Figure 99: Recon presets dialog box

4. Create a new Preset by selecting New Preset.

Preset Name	Slice Thickness & Specing	Metal Artifact
Enter name	Select	Select
Window Width	Window Level	Noise Reduction
		Select
Scan Type	Multi-Energy	Kernel
Select	Select	Select

Figure 100: New preset dialog box

- 5. Enter the Preset Name, Slice Spacing & Thickness, Metal Artifact, Window Width, Window Level, Noise Reduction, Scan Type, Multi Energy and Kernel.
- 6. Select **Save** to update new preset.

Setting recon preset groups

Recon Preset Groups allow the administrator to define and customize reconstruction groups. Recon Groups can be added to a protocol to automatically start reconstructions from within the Acquisition tab.

The administrator can delete existing recon groups, update an existing recon group, or create and save a new recon group.

- 1. Select the **System** icon. The System Screen appears.
- 2. Select System Settings icon. The System Settings screen appears
- 3. Select the Recon Groups icon. The Recon Preset Groups dialog box appears.

New Group	Organize your recon	presets into groups.		
Group Name		Presets		
lodine Map & 70 keV	Axial HR			×
0.707 Bone and 0.707 \$	ST Mixed	2	1	×
1.414 Bone and 1.414 S	ST Mixed	4		×
2.828 Bone and 0.707 \$	ST Mixed	2	1	×
5.656 Bone and 0.707 \$	ST Mixed			×
0.424 Bone and 0.424 S	ST Mixed HR	2		×
0.848 Bone and 0.848 \$	T Mixed HR	4	1	×

Figure 101: Recon preset groups dialog box

4. Create a new group by selecting New Group.

Group N	lame Scen		Add Recon Preset	
Enter te	ext	Select	Select	
Pre	eet Name			

Figure 102: New recon preset group dialog box

- 5. Enter the Group Name, Scan Type, and Recon Preset(s).
- 6. Select **Save** to update new preset.

Setting MPR presets

MPR Presets allow the administrator to define and customize MPR Presets.

The administrator can delete existing recon groups, update an existing recon group, or create and save a new MPR Preset.

- 1. Select the **System** icon. The System Screen appears.
- 2. Select System Settings icon. The System Settings screen appears

3. Select the MPR Presets icon. The MPR Presets dialog box appears.

Name Biab Thickmeas Biab Spacing 0.625 x 0.625 0.625 0.625 1.26 x 1.25 1.25 1.25		
0.625 × 0.625 0.625 0.625 1.25 1.25 1.25 1.25	Siab Thickness Siab Spacing	
1.25 x 1.25 1.25	.625 0.625 0.625	×
	25 1.25 1.25	×
2.5 x 2.5 2.5		×
5x5 5 5		×
10 x 10 10 10		×

Figure 103: MPR presets dialog box

4. Create a new preset by selecting New Preset.

MPR PRESETS	
New Preset	
Name (Optional) Slab Thickness Enter name 0.6132813 Slab Spacing 0.3066407	

Figure 104: New MPR preset dialog box

- 5. Enter the Name, Slab Thickness, and Slab Spacing.
- 6. Select Save to update new preset.

Setting windowing presets

Windowing presets allow you to quickly set the Window Level and Window Width for your images.

The administrator can delete existing windowing presets, update an existing windowing preset, or create and save a new windowing preset.

- 1. Select the **System** Icon. The System Screen appears.
- 2. Select System Settings icon. The System Settings screen appears
- 3. Select the Window Presets icon. The Windows Preset dialog box appears.

NEW PRESET	Name Enter name	Window Lev Enter	el Window V	Midth
PRESETS	Name	Window Level	Window Width	
	Abdomen			
	Angio	300	600	
	Bone	300	1500	
	Brain			
	Chest		400	
	Lungs	-400	1500	

Figure 105: Window preset options

- 4. Create a **New Preset** by entering the name, window level, and window width and select **Add.**
- 5. To edit a **Preset**, select the **Preset icon**.



Figure 106: Window preset option

6. Make desired changes and select **Set** to update **Preset**.

Selecting image orientation

NeuroLogica describes patient orientation as if the viewer were looking towards the front of the gantry (where the Mood Ring is). In other words, if the patient is lying face up with their head in the gantry, the image orientation displays the patient's Right side on the Left side of the Viewer. If the patient's feet are going into the gantry, the image orientation displays the patient's Left side on the Left side of the Viewer.

The MPR Image Orientation will always display in the preferred anatomical viewing orientation, where the patient's Right side is on the Left side of the viewer.



Figure 107: Image orientation

Note: Image Orientation is not a customizable configuration for OmniTom Elite with PCD.

A user can modify patient orientation, but the view will always be as if the viewer were at the patient's feet for the anatomical position.

HFS	Head First-Supine
HFP	Head First-Prone
HFDL	Head First-Decubitus Left
HFDR	Head First-Decubitus Right
FFS	Feet First-Supine
FFP	Feet First-Prone
FFDL	Feet First-Decubitus Left
FFDR	Feet First-Decubitus Right

Patient Orientation abbreviation list:

You can modify patient orientation by selecting either arrow as shown in Figure 108.



Figure 108: Patient orientation toggle

Note: Always verify that your patient is in the correct orientation, prior to scanning.

ORIENTATION

Please verify that your patient's orientation matches the image above. If they do not match, use the arrows to cycle through available orientations.

Figure 109: Orientation verification dialog

Chapter 6 Protocol Manager

Note: You must have administrative privileges and be logged in as an administrator to access this area in the application.

Incorrect changes to the system configuration may make the system inoperative.

Protocol Manager lets you set how the limited and restricted operator uses protocols.

Creating a new protocol

- 1. Select System Icon.
- 2. Select Protocol Manager. The Protocol Manager dialog appears.



Figure 110: Protocol manager

3. Select Adult.

Note: Protocol parameters are customized to meet your requirements in conjunction with local and nationally recognized published guidelines. These protocols *must be* approved by your facility physicist *before* the system's acceptance.

WARNING Any modification to an existing protocol, or any new protocol created, should be reviewed, and approved by a radiologist and/or residing medical physicist. Failing to do so could cause a patient to receive an excessive and/or unnecessary dose of ionizing radiation.

Note: The **New** button and others (depending on what was done previously) are active *after* you select a colored orb corresponding to the body part and then select the protocol that shows in the **Protocol** list. Existing protocols appear in the **Protocol** list box, as shown, in the figures below.

4. Select the **New** button to create a new protocol.

Resolution	PROTTOOOL NL Brain - Axaal UHH NL Head Trauma NL Head Trauma HR O X NL Brain CT Angiography O X NL Brain CT Perfusion O X NL Sinus/Facial/Orbit NL Sinus/Facial/Orbit Sinus/Facial/O	
	EDIT	

Figure 111: Protocol Manager New Protocol

NEW PROTOCOL				
Protocol Name				
Enter text				
Protocol Description				
Enter text				
Anatomical Reference				
Select				
Default Study				
Description				
Enter text				
		A A A A A A A A A A A A A A A A A A A		~

5. The **New Protocol** dialog box appears.

Figure 112: New protocol dialog box

The **Protocol Information** and **Protocol's Series** areas are *empty*. The **Patient Position** settings are identical whether it is for an adult, pediatric, or emergency patient.

Protocol Name	The name of the protocol as it will be displayed in the protocol manager. Factory Protocols begin with the letters NL.
Protocol Description	The description of the protocol you assign; for example, Axial head or Helical head.
Anatomical Reference	References what part of the anatomy will be scanned; for example, head or chest.
Default Study Description	The DICOM image tag; if entered, this description will also appear in PACS as a Study Description DICOM tag (00081010).

For **Protocol Information**, enter your information in the following text boxes:

- 6. After adding all required **Protocol Information:** Protocol Name, Protocol Description, Anatomical Reference, and Default Study Description, select **Add.**
- 7. Under **Protocol Series**, select the **New** button. The **New Series** dialog box appears.



Figure 113: New series dialog box

- 8. For **Type**, select one of the following:
 - Axial
 - Helical
 - Reference
 - Scout
- 9. For **Scout Type**, select one of the following:
 - AP
 - PA
 - Lateral
 - Scout Type is not available for Axial, Helical, and Reference scan modes.
- 10. For **Body Part**, select the appropriate Body Part from the dropdown menu.
- 11. For **Description**, enter the defined study description.
- 12. For **kV** (scan voltage), select one of the following:
 - 70 To set the scan kV to 70
 - 80 To set the scan kV to 80
 - 100 To set the scan kV to 100
 - 120 To set the scan kV to 120

The scan voltage obtainable from the x-ray tube ranges from 70 to 120 kV. At a nominal scan voltage of 120 kV, a maximum output power of 1 kW is maintained by the x-ray tube for at least 4 seconds if the current heat capacity of the tube does not exceed 80%.

13. For **mA** (scan current), select the appropriate selection (5.0 to 20 mA with an increment of 5 from the dropdown. X-ray tube power of any scan combination is computed as the product of the scan voltage and the scan current.

The scan time is determined by the x-ray tube's current heat capacity. Scan Power = Scan Voltage (kV) x Scan Current (mA) 14. For **Scan Time**, if applicable, the calculated number appears here, depending on other selections.

The scan time is automatically calculated based on the parameters selected. For example, the **Scan Time** for an axial 120mm scan with 2 seconds per scan is 24 seconds. The **Scan Time** for a helical scan with same coverage is 12 seconds (1 second scan).

- 15. For Coverage (mm), enter the total scan distance.
- 16. For **Number of images**, if applicable, the calculated number appears here. The number of images is calculated based on the slice thickness and length of the scan.
- 17. For **Window Width**, enter the range of CT numbers that are distributed over the viewable gray scale of the display device or film.
- 18. For Window Level, enter the CT number in the center of the viewable gray scale.
- 19. For Kernel, select the image reconstruction kernel from the following list of kernels:
 - Soft Tissue
 - Posterior Fossa
 - Sharp
 - Bone

Kernel allows *only* **Soft Tissue** for **Reference** scan mode and *only* Posterior Fossa for **Scout** scan mode.

20. Available **Slice Thickness/Spacing**, options vary depending on the resolution selected:

Table 25: Slice Thickness	and Spacing Options
---------------------------	---------------------

Standard Resolution	High Resolution	UltraHigh Resolution (UHR) (Axial Scan Mode Only)
0.7 x 0.7 1.4 x 1.4 2.8 x 2.8 5.6 x 5.6 11.3 x 11.3	0.4 x 0.4 0.8 x 0.8 5.5 x 5.5 11.0 x 11.0	0.1 x 0.1 0.2 x 0.2 1.4 x 1.4 2.8 x 2.8 5.6 x 5.6 11.2 x 11.2

- Slice Thickness/Spacing is not available for Scout scan modes.
- 21. For **Rotations**, select one of the following scan times:
 - 1 Second(s)
 - 2 Second(s)

Rotations is available for Axial and Reference scan mode, only.

22. For **Pitch**, there is only one selection for travel time (per scanner rotation):1; the scanner is moving at 10mm per second.

Pitch describes a scanner's travel and how fast the scanner is moving per rotation; **Pitch** is available for **Helical** scan mode, *only*.

23. For **Scan Delay**, select options and enter the delay time that will occur after pressing the **Start Scan** button and before the scan begins.

24. For **CTDIvol (mGy)**, if applicable, the calculated number appears here, depending on other selections.

CTDIvol (mGy), applies an unknown to **Scout** scan modes; it applies 12.6 for **Helical** and 50.4 for **Reference**.

CT Dose Index Volume (CTDIvol)_represents the dose for a specific scan protocol, which considers gaps and overlaps between the radiation-dose profiles from consecutive rotations of the x-ray source. The CT Dose Index Volume is noted as CTDIvol. The CTDIvol is calculated differently for both the **Axial** and the **Helical** modes:

- For Axial scan mode: CTDIvol = [(N x T)/I] x CTDIw
- For **Helical** scan mode: CTDIvol = 1/pitch x CTDIw

Dose Length Product (DLP) is the measure of ionizing radiation exposure during the entire acquisition of images. Therefore, DLP (mGy.cm) = CTDIvol (mGy) x irradiated length (cm).

25. Select the **Enable AEC** option, if applicable.

	Allows you to automatically adapt the tube current or mA
Enable AEC	according to the patient's body habitus in order to achieve
	the specified image quality at the lowest possible dose.

For detailed information on this feature, refer to Scanning with special features on page 190.

26. Select the **Noise Reduction** option, if applicable for Axial scan mode only.

	Allows you to select Low, Medium or High levels of Noise
Noise Reduction	Reduction to decrease noise in Axially scanned images
	only.

For detailed information on this feature, refer to Scanning with special features on page 190.

Adding a New Protocol using Copy from an Existing Protocol

- 1. Select System Icon.
- 2. Select **Protocol Manager**. The Protocol Manager dialog appears.

ADUT PEDATRC	PROTOCOL	
Resolution 🛛 👔	NL Head Trauma 👔 💿 🗙	
	NL Head Trauma HR 👔 💿 🗙	
	NL Brain CT Angiography 🧊 💿 🗙	
🕞 (📂) 🐼 -	NL Brain CT Perfusion	
	NL Sinus/Facial/Orbit	
rest. Aco	NL Sinus/Facial/Orbit HR 👔 💿 🗙	anga NL study C7
TEST, PCD	axial water 👔 💿 🗙	24 NL Study CT
- Masec assa	helical water	
water balance a	12/4	19091 NE Study CT
	EDIT COPY	

Figure 114: Protocol manager

- 3. Select Adult.
- **Note:** Protocol parameters are customized to meet your requirements in conjunction with local and nationally recognized published guidelines. These protocols *must be* approved by your facility physicist *before* the system's acceptance.
- WARNING Any modification to an existing protocol, or any new protocol created, should be reviewed, and approved by a radiologist and/or residing medical physicist.
 Failing to do so could cause a patient to receive an excessive and/or unnecessary dose of ionizing radiation.
 - Note: The New button and others (depending on what was done previously) are active *after* you select a colored orb corresponding to the body part and then select the protocol that shows in the **Protocol** list. Existing protocols appear in the **Protocol** list box, as shown, in the figures below.
 - 4. Select the protocol you want to Copy to create a new protocol and tap the Copy button.



Figure 115: Copy Protocol

Axial Brain High Res			
Anatomical Reference			
Head			
Study Description			
Brain - Axial High Re			
Resolution			
High			
· · · · · · · · · · · · · · · · · · ·		=~	_

Figure 116: Copied Protocol

5. The **Protocol Information** and **Resolution**, which is used for protocol filtering, is copied from the previous protocol, modify that information as required and select **Save.**

Anatomical Reference Head Study Description Brain - Axial High Re Resolution High	Protocol Description Axial Brain High Res			
Study Description Brain - Axial High Re Resolution High	Anatomical Reference Head			
Resolution High	Study Description Brain - Axial High Re			
	Resolution			
	High			

Figure 117: Save Modified Protocol

5. The copied protocol series will appear.

rotocol Description	Scout	Axial HR	New
Axial Brain High	Ten Tel	Tan Barline Internation	
natomical Reference Head	An and a second	Recher Southers MacCharge	-
Study Description			
Brain - Axial High			
Resolution			
High			
	s,	AVE	

Figure 118: Copied Protocol

6. Modify and add an additional series required and select Save.

	Scout	Axial HR	New
Protocol Description			
Axial Brain High	New Contraction of the Contracti	Tan Bar tree (n), (manual free free free free free free free fre	
Anatomical Reference	Marchan Barten Barten Marchange	Destroy Mathematical Mathematical	
Head	Description	International In	
	an and a second an		
Study Description			
Brain - Axial High			
Resolution			
High			
		<u> </u>	
	, s	AVE	

Figure 119: Save Copied Protocol

7. Adjust the protocol order if needed.



Figure 120: Adjust protocol order

Editing an existing protocol

The **Edit** button is used in **Protocol Manager** to edit an existing protocol. Protocols that begin with "**NL**" are NeuroLogica reference protocols and cannot be edited or deleted.

- 1. Select System Icon.
- 2. Select Protocol Manager. The Protocol Manager dialog appears.
- 3. Select Adult.

Note: Protocol parameters are customized to meet your requirements in conjunction with local and nationally recognized published guidelines. These protocols *must be* approved by your facility physicist *before* the system's acceptance.

4. Select the icon corresponding to the appropriate body part.

lcon	Icon Name
	Brain
	Spine
	Lung
	Upper Extremity
	Abdomen
	Lower Extremity

Table 26: Anatomical orbs

- 5. Select the protocol you wish to edit in the **Protocol** list.
- 6. Select the **Edit** button.

ADULT PEDIATRC	PROTOCOL	
	NL Head Trauma 👔 💿 🗙	
	NL Brain CT Angiography (1) (2) (2) (2) (2) (2) (2) (2) (2) (2) (2	
	NL Sinus/Facial/Orbit	HL Study CT
	axial water	NL Study CT
iveator, actai	helical water	

- Figure 121: Edit button illuminated
- 7. The **Edit Protocol** dialog box appears.

Bertrees Descarton	Axial	New	
axial water			
	100 Per		
Anetornosi Reference			
Head			
	1000		
	A REAL PROPERTY AND	- International statements of	المح
	-BAV		

Figure 122: Edit protocol dialog box

8. Make your changes.

See "Creating a new protocol" on page 128 to learn how the fields and options perform to make informed choices on what to change.

- 9. Select the **Update** button to save your changes to the existing protocol.
- 10. Select the Save button to finish saving to the Protocol Manager.
 Select the Cancel button to return to the previous dialog box.
- 11. Select the **Close** button to exit.

Deleting a protocol

- 1. Select System Icon.
- 2. Select Protocol Manager. The Protocol Manager dialog box appears.
- 3. Select Adult.

Note: Protocols that begin with "**NL**" are NeuroLogica reference protocols and cannot be edited or deleted.

- 4. Select the icon corresponding to the appropriate body part.
- 5. Select the protocol from list to be deleted.



Figure 123: Protocol manager with a protocol selected for deleting

6. Select the **Delete** button.



Figure 124: Delete protocol icon

7. The Delete Confirmation pop-up appears.

?		PROTCOL	E.
3	Are you sure you t proto AB	want to delete this col?	у -
	NL Parament	11-X	
	Cancel:	Confirm	

Figure 125: Delete protocol confirmation pop-up

Perform one of the following in the **Delete Protocol Confirmation** pop-up:

- Select the Confirm button to delete the selected protocol.
- Select the Cancel button to return to the Protocol Manager dialog box.
- 8. The **Delete Protocol Confirmation** dialog box disappears, and the **Protocol Manager** dialog box appears.

9. Select the **Close** button to exit.

Hiding protocols

- 1. Select System Icon.
- 2. Select **Protocol Manager**. The Protocol Manager dialog box appears.
- 3. Select the icon corresponding to the appropriate body part.
- 4. Select the **Eye** icon next to the protocol you want to hide. A line through the **Eye** icon indicates that the protocol is hidden, as seen in Figure 126.

/	PROTOCOL	Scheduled Date
	Axial Head	
	NL Adult Axial Head	(
	NL Adult Head Trauma	(
	NL Adult Helical Head	(
	NL Adult Stroke	1 Ø ×
	NL Adult Head CT Angiography	()
	NL Adult Head CT Perfus	ion 🛊 💋 🗙
	NL Temporal Bones	

Figure 126: Protocol manager hide protocol function

Moving protocols

- 1. Select System Icon.
- 2. Select Protocol Manager. The Protocol Manager dialog box appears.
- 3. Select the icon corresponding to the appropriate body part.
- 4. Select the Arrow icon next to the protocol you want to move. Grab the arrow and arrange the protocols in the order you want them listed.



Figure 127: Protocol manager move protocol function

Chapter 7 Daily Calibration and Quality Assurance

This chapter will cover the daily steps needed to get the scanner ready for imaging. Part of that is to learn how to perform a daily air calibration and how to run the **Quality Assurance** (QA) tool that verifies the system is at its optimum performance. In addition, we will discuss some of the radiation safety issues such as the CTDI and the scatter from the scanner. The chapter will teach you how to predict the CTDI or patient exposure.

Daily calibration

Keep in mind that *before* using the OmniTom Elite with PCD, you *must* perform a **Daily Calibration** to ensure that the system is at its optimum performance.

- **Note:** The OmniTom with PCD includes a two-sided phantom that is used for both the Daily Calibration and the QA test. Ensure you are using the correct side of the phantom for each procedure.
- **Note:** If a Daily Calibration has not been performed for three days, a Tube Seasoning prompt will appear when the user logs in.
- **Note**: You must perform a Daily Calibration for the resolution mode you plan to use for scanning patients. NeuroLogica recommends this be done once per day.

Performing a daily calibration

1. To provide optimum image quality, a **Daily Calibration** is required to be performed once per day, prior to normal scanning. The water phantom/QA phantom below is supplied with the OmniTom Elite PCD scanner.



Figure 128: PCD Water/QA Phantom

Note: The phantom is intentionally offset in the 'koozie', this offset accounts for patient anatomy that may be offset in the scanner bore.

2. Insert the water phantom into the bore prior to the Daily Calibration.



Figure 129: Water phantom inserted in the scanner bore

- 3. On the LCD screen, press the Laser On button.
- 4. Use the laser indicators to align the phantom to ensure the beam travels through the water portion of the phantom.



Figure 130: Set the laser in the middle of water portion of the phantom

5. On the tablet, select the **System Icon.**

6. Select Daily Calibration.



Figure 131: Daily Calibration Screen

7. The Daily Calibration screen allows you to select the desired Resolution mode and/or mA values to calibrate **Daily Calibration** times for the OmniTom Elite with PCD are dependent on the resolution selected and detailed in the table below:

Table 27: Calibration time per mA

STANDARD	Four (4) minutes per mA value selected
HIGH	Ten (10) minutes per mA value selected
ULTRA-HIGH	Twenty (20) minutes per mA value selected

Note: All axial and helical protocols shipped with the OmniTom PCD system are built using 120kVp and 20mA so most customers at this time will only calibrate 5mA (scout) and 20mA for each resolution mode they will scan with.

8. You can use the **'Show Calibration Percentages'** check box to confirm the status of each mA setting.

5 mA:	5mA: 0%	5 mA: 0%		
10 mA: 0%	10 mA:	10 mA: 0%		
15 mA: 0%	15 mA: 0%	15 mA: 0%		
20 mA:	20 mA:	20 mA: 0%		
SH	OW CALIBRATIC			
	RCENTAGES			
ESTIMATED	CALIBRAT	TION	Insert the water ph Press 'Begin' when ready	antom into the bore. and stand at a safe distance.

Figure 132: Show Calibration Percentages

- 9. If specific Resolution modes and/or mA values will NOT be used for normal scanning, you can clear the check box next to the Resolution and/or mA value to disable the calibration for that selection, which will shorten the calibration time.
- 10. Press **Begin** to start **Daily Calibration**. The progress of the **Daily Calibration** will appear on the tablet.



Figure 133: Daily Calibration Progress Bar

11. When the calibration is completed the **Daily Calibration Success** dialog box will appear.


Figure 134: PCD Daily Calibration success

12. Press the **Close** button. The **Daily Cal icon** will change to green and show that it is at 100%.



Figure 135: PCD daily calibration status

Tube seasoning

If the system detects that a Daily Calibration has not been performed in at least three days, a **Tube Seasoning** prompt will appear when the user powers up the system and logs in. Press the **Continue** button to perform the Tube Seasoning. The Tube Seasoning will take approximately 15 minutes.



Figure 136: Tube seasoning prompt

The **Tube Seasoning** Dialog Box will appear. Press **Begin** after checking that there is nothing in the bore.



Figure 137: Tube seasoning dialog box

The QA phantom overview

Performing a quality assurance test

To ensure the system is at its optimum, factory-specifications level, the tablet provides Quality Assurance tools to verify the system's state and to perform image-quality verification.

Before you begin this section, be sure to run a fresh **Daily Calibration** on the system using the tablet.

The QA protocol is shipped with the system and appears when you click **Quality Assurance** from the System Panel (on the bottom, right corner of the screen on the tablet). You cannot customize or modify the QA protocol.

Before beginning the **Quality Assurance** test, make sure the water phantom/QA phantom is available and ready to position in the bore.

Note: You must perform a Quality Assurance scan for the resolution mode you plan to use for scanning patients.

The Quality Assurance test should be conducted per the local (hospital) requirements, typically it should be done daily or prior to any scheduled use of the scanner.

- **Note:** The OmniTom with PCD includes a two-sided phantom that is used for both the Daily Calibration and the QA test. Ensure you are using the correct side of the phantom for each procedure.
- 1. Place the **QA phantom** in the bore.



Figure 138: OmniTom with PCD QA Phantom

2. On the LCD touch screen, press the Laser On button.

- **Note:** The phantom label should face the front of the scanner and be positioned at the bottom. The red insert should be on the operator's bottom right when facing the scanner. The position of the phantom will greatly affect the QA results.
- 3. Align the phantom by lining the QA phantom's etching (line(s)) with the laser light.



Figure 139: Phantom's etchings appear on top and sides

Note: The laser will automatically shut off 10 seconds after pressing the Laser On button.

4. On the tablet, select the **System Icon**, then select **Quality Assurance**. The **Quality Assurance** Dialog Box will appear.



Figure 140: PCD Quality Assurance dialog box

- 5. The Quality Assurance screen allows you to select the desired Resolution modes you want to perform. By default, all Resolutions are selected. If specific Resolution modes will NOT be used for normal scanning, you can clear the check box next to the Resolution to disable that Quality Assurance scan.
- 6. Press the Begin button. The scan delay (10 seconds) will appear on the LCD.
- 7. Press the **Start** button on the LCD.
- 8. Wait for the Quality Assurance results to appear.

- 9. Review the results.
- 10. If multiple Resolutions have been selected, the **Quality Assurance Continue** screen will appear. Click the **Continue** button to perform the next resolutions QA scan.



Figure 141: PCD Quality Assurance Continue

11. When all QA scans have been completed, click the **Close** button on the QA Results pop-up when finished reviewing.





Note: Items in orange are failed results; reposition your phantom to perform another scan. Often positional issues cause the failure. If you try multiple times and failures persist, call your service representative. Review the results.

The QA image and QA results screen are available for viewing from the Browser and will show as two separate files on the Browser screen.

Ensuring good image quality

In order to produce consistent image quality over the system's lifetime, it is strongly advised that you:

- Establish and maintain a regular Quality Assurance (QA) program. NeuroLogica typically recommends a weekly QA test. For infrequent use NeuroLogica recommends running the QA prior to scanning.
- **Keep** a record of the QA results. QA images are typically stored.
- **Review** the record periodically to check the consistency of the test results. Contact Customer Services if the QA performance is degrading over time.

OmniTom Elite with PCD dose information (21 CFR 1020.33 c)

Dose is measured using a standard 16cm CTDI phantom. Both Surface and Center values of CTDI₁₀₀ are obtained for each scan kV. The CTDI₁₀₀ basis values at the maximum scan technique, 120kV and 20mA, are provided in Table 28.

Table 28: OmniTom Elite with PCD CTDI100 values

CTDI100 Center (C)	CTDI100 Surface (S)	CTDIw (W)
14.08	17.97	16.66

Scan voltages other then 120kV are not yet supported for imaging with the OmniTom Elite with PCD system, but preliminary dose measurements are available. The scan voltage adjustment factors for scans other than 120kV are provided in Table 29.

kV	CTDI100 Center (C)	$CTDI_{100}$ Surface (S)	CTDIw (W)
120	1.0	1.0	1.0
100	0.64	0.68	0.67
80	0.37	0.42	0.40
70	0.25	0.30	0.28

Table 29: Scan voltage weighting factors for dose calculations

 $CTDI_{100}$ values for a particular kV are obtained by multiplying a $CTDI_{100}$ basis value from Table 28 with an adjustment factor from Table 29.

Error! Reference source not found. provides the Center, Surface and Weighted $CTDI_{100}$ values normalized to 1 mAs for all scan voltages. $CTDI_{100}$ units are mGy/1mAs.

Table 30: Normalized CTDI₁₀₀ as a function of kV

Normalized CTDI (mGy/1mAs)						
kV	120	100	80	70		

Note: If you notice degradation in image quality or a change in QA values; schedule a site visit and let a Field Service Representative or Imaging Physicist run more detailed tests.

Center	0.704	0.454	0.257	0.173
Surface	0.899	0.613	0.375	0.266
Weighted	0.833	0.560	0.335	0.235

CTDI₁₀₀ measurements

The 16cm CTDI phantom is used for dose measurements. The phantom should be positioned within 1cm of the scanner isocenter. Centering can be verified with imaging.

The exposure is measured at the center of the phantom as well as the four surface positions at 12, 3, 6 and 9 o'clock. It is recommended that four surface exposure measurements be averaged and used in the $CTDI_{100}$ calculations. Measurements from two opposite surface locations will also be sufficient, e.g., the 12 and 6 o'clock positions or the 3 and 9 o'clock positions. Using the 12 o'clock position only might overestimate the patient dose due to the configuration of the x-ray radiation.

For each position of the dose probe, CTDI₁₀₀ is computed using the following equation:

$$CTDI_{100} = \frac{E \times 0.0087 \left(\frac{mGy}{mR}\right) \times CL(mm)}{N \times T(mm)}$$

The detector collimation N x T = 11.15mm. The length CL = 100mm. Exposure E is measured in mR. The units of $CTDI_{100}$ are mGy.

Weighted CTDI is computed using the Surface and Center measurements:

$$CTDI_w = \left(\frac{2}{3} \times CTDI_{surface} + \frac{1}{3} \times CTDI_{center}\right)$$

Table 31 shows the measurement for the 100kV scan voltage using 30mAs. The exposures are measured three times to confirm repeatability.

Position	m1	m2	m3	Average	CTDI ₁₀₀	CTDIw
Center	174.4	174.3	174.8	174.5	13.6	-
12 O'clock	270.2	269.7	270.5	270.1333	21.1	18.6
9 O'clock	211	212	211.5	211.5	16.5	15.5
6 O'clock	206.6	206.5	206.3	206.4667	16.1	15.3
3 O'clock	253.9	253.7	253.6	253.7333	19.8	17.7
Average Surface	-	-	-	235.4583	18.4	16.8

Table 31: The detailed dose measurements and results for 100kV and 30mAs

Note regarding detector collimation

 $CTDI_{100}$ and dose efficiency measurements depend on the exposed length of the detector sensor, called detector collimation. This is the number of active detector rows multiplied by the row length at isocenter, N x T. The three supported resolution modes, Standard Resolution (SR), High Resolution (HR), and Ultra-high Resolution (UHR) are specified in Table 32.

Table 32: Detector collimation

Parameter	SR	HR	UHR
N rows	16	26	80
row length, T, mm	0.706	0.424	0.141
collimation, N x T, mm	11.3	11.0	11.3

The slight deviation of N x T for HR mode is due to the summation of detector sub-pixels into an integer number of rows. To simplify the dose calculations, we use a value of N x T = 11.5mm as representative of all resolution modes. The error associated with this assumption is less than 1.5 percent.

Calculation of CTDI100

 $CTDI_{vol}$ may be computed using Table 30 and the prescribed scan parameters. For axial scan mode, the $CTDI_{vol}$ can be calculated using the following equation:

$$CTDI_{vol}(kV, mA, S) = \left(\frac{mA \times S}{NSI}\right) CTDI_w(kV)$$

Where **mA** is the x-ray tube current in mA, **S** is the scan rotation time in seconds, and **CTDI**_w(**kV**) is the weighted value taken from Table 30 for the specific scan kV. **NSI** is the normalized scan increment, which is always equal to 1 for an axial scan.

Recall that $CTDI_w$ in Table 30 has been normalized to an exposure of 1mA.

For helical scan mode, $CTDI_{vol}$ is also a function of helical pitch:

$$CTDI_{vol}(kV, mA, S, Pitch) = \left(\frac{mA \times S}{Pitch}\right) CTDI_{w}(kV)$$

CTDI_{vol} is measured in units of mGy.

The Dose-Length Product (DLP) is defined as the product of $\mathsf{CTDI}_{\mathsf{vol}}$ and the scan length in centimeters.

$$DLP = CTDI_{vol} \times \frac{Scan \, length \, in \, mm}{10.0 \, mm/cm} \quad mGy \cdot cm$$

Sample calculation

For example, consider an axial scan of 120kV, 20mA and 1 second scan length of 170mm. NSI=1 because the scan increment is the same as the detector collimation. The CTDI_{vol} is:

$$CTDI_{vol}(120 \ kV, 20mA, 1s) = \left(\frac{20 \times 1}{1}\right) \times 0.833 \ mGy = 16.7 \ mGy$$

The DLP is:

$$DLP = 16.7 \times \frac{170}{10.0} = 283.2 \, mGy \cdot cm$$

A helical scan of pitch 1.0 with the same kV and mA would have the same $\mathsf{CTDI}_{\mathsf{vol}}$ and DLP as the axial example.

The scout dose

During a scout scan, the scanner moves at 30mm/sec, and the detector collimation is 11.3mm. The corresponding effective pitch is 2.65. The dose calculation is similar to that for helical scan mode:

$$CTDI_{vol}(kV, mA, S) = \left(\frac{mA \times S}{2.65}\right) CTDI_w(kV)$$

A scout scan using 120kV with 5mA will have the following $CTDI_{vol}$:

$$CTDI_{vol}(120kV, 5mA, 1s) = \left(\frac{5 \times 1}{2.65}\right) 0.833 = 1.57 \ mGy$$

Recommended dose verification procedure

The dose should be measured in clinical mode, using the tablet. The scan protocol should have no scout since that will affect the location of the initial scan. The protocol for each scan voltage is set as follows:

Axial
120 kV
20 mA
1 sec
SR
Posterior Fossa
0mm
11.3mm

The scan current and time may be adjusted according to physicist recommendation.

The calculated value of $\mathsf{CTDI}_{\mathsf{vol}}$ may be compared with the predicted value displayed on the tablet.

OmniTom Elite with PCD dose in air

For 2 second exposures of 20mA, the measured values of dose in air are shown in Table 33:

Table 33: kV vs. dose in air

kV	Dose in air (mGy)
120	42.4
100	30.8
80	19.6
70	14.8

QA measurements

The QA phantom provided by NeuroLogica is typically used to monitor the scanner on a daily basis. Detailed imaging performance measurements made using the ACR accreditation phantom are presented here.

CT Number Accuracy

Material	SR		HR		UHR			
	Axial	Helical	Axial	Helical	Axial	Helical	ACK LIMIUS	
Water	-1.7	-3.5	-1.7	-2.8	0.5	N/A	-7 to 7	
Polyethylene	-97.1	-98.9	-97.1	-95.6	-95.5	N/A	-107 to -87	
Bone	893.5	893.2	893.5	890.9	906.2	N/A	850 to 970	
Air	-983.7	-981.0	-978.5	-976.0	-977.0	N/A	-1005 to -970	
Acrylic	113.6	111.6	111.3	110.1	113.7	N/A	110 to 130	

Table 34: CT values of the ACR inserts for each resolution mode

Contrast-to-noise ratio and low contrast resolution

Module 2 of the ACR phantom is used to measure the low contrast resolution of the scanner. It consists of sets of four cylinders with diameters ranging from 2 mm to 6 mm, and a 25 mm insert. Each cylinder has a +0.6% (6 HU) difference from the background CT value. The first test involves the observer identifying the smallest diameter cylinders that can be discerned at a window center of 100HU and width of 100HU. The second part of the test involves calculating the contrast-to-noise ratio (CNR). Table 35 shows the low contrast results for all resolution modes. The results are within the ACR

requirement for CT accreditation. After 2012 ACR dropped the size of the smallest observable rods and kept the CNR as the only requirement for low contrast detectability.

	SR		н	२	UHR		
	Axial	Helical	Axial	Helical	Axial	Helical	
Slice thickness	5.66 mm 5.66mm		5.51 mm 5.51 mm		5.64 mm	N/A	
Diameter of smallest visible set of cylinders	6 mm 6 mm		6 mm	6 mm	6 mm	N/A	
Contrast-to-Noise Ratio	ntrast-to-Noise 1.5 0		1.13	0.46	1.2	N/A	

Table 35: Results of low contrast resolution test

Figure 143 shows the low contrast module images for all resolution modes in both axial and helical scan modes. The 6.0mm rods are visible on both images at 37.16mGy as required by ACR accreditation.



< Axial - SR >

< HR >

< UHR >





Uniformity

	SR			HR			UHR		
Location	CT Value	Δ CT	Noise	CT Value	Δ CT	Noise	CT Value	Δ CT	Noise
1	-3.46	0.36	6.59	-3.12	-0.25	7.13	-1.85	1.13	6.33
2	-2.76	1.06	6.24	-2.31	0.56	6.90	-1.86	1.12	6.42
3 (Center)	-3.82	0	7.01	-2.87	0	8.06	-2.98	0	10.80
4	-3.74	0.08	6.60	-4.07	-1.20	6.69	-1.74	1.24	6.13
5	-3.58	0.24	6.46	-2.88	-0.01	6.83	-1.66	1.32	6.36

Table 36: CT uniformity and noise level of ACR accreditation phantom for Axial scans

ROI	SR			HR		
Location	CT Value	ΔCT	Noise	CT Value	Δ CT	Noise
1	-4.56	0.33	7.24	-4.49	1.30	12.57
2	-4.03	0.86	8.03	-4.08	1.72	12.11
3 (Center)	-4.89	0	9.35	-5.79	0	14.62
4	-4.88	0.01	7.80	-6.26	-0.47	12.55
5	-4.42	0.47	7.73	-5.31	0.49	12.88

Both scan modes satisfy the ACR requirements of uniformity below 5.0HU.

Noise

The noise was measured within the center of the ACR module using an ROI of 100mm diameter. Table 38 and Table 39 list the CT values and the measured noise in 5mm slices for all resolution modes in both Axial and Helical scan modes.

Table	38:	Axial	image	noise	results
Tuble	50.	/ with	nnuge	10150	results

	SR		HR		UHR	
Slice Index	Mean	SD	Mean	SD	Mean	SD
Slice 1	-3.32	7.90	-2.72	8.81	-4.48	7.69
Slice 2	-3.42	7.96	-2.56	7.88	-4.16	8.35
Average	-3.37	7.43	-2.64	8.34	-4.32	8.02

	S	R	HR		
Slice Index	Mean	SD	Mean	SD	
Slice 1	-4.35	8.85	-4.52	14.21	
Slice 2	-4.90	8.78	-5.04	13.98	
Average	-4.62	8.82	-4.78	14.10	

Table 39: Helical image noise	results
-------------------------------	---------

MTF

Table 40 and Table 41 present the MTF curves and their cutoff frequencies for the Soft Tissue, Posterior Fossa, and Bone kernels for both Axial and Helical scan modes. They sufficiently match in their frequency responses and therefore prove equivalency between them.

Table 40: Cutoff frequencies of Axial scans for the kernels at different MTF responses in lp/cm

Resol.		SR			HR			UHR	
	MTF	MTF	MTF	MTF	MTF	MTF	MTF	MTF	MTF
Kernel	50%	20%	10%	50%	20%	10%	50%	20%	10%
Soft tissue	3.77	5.30	5.98	3.84	5.52	6.31	4.00	5.67	6.41
Post Fossa	4.56	6.05	6.69	4.84	6.86	7.75	5.19	7.02	7.81
Bone	6.97	7.59	11.43	11.56	13.09	14.00	13.34	15.00	16.09

Table 41: Cutoff frequencies of Helical scans for the kernels at different MTF responses in lp/cm

Resol.	SR			HR		
	MTF	MTF	MTF	MTF	MTF	MTF
Kernel	50%	20%	10%	50%	20%	10%
Soft tissue	3.21	4.69	5.40	3.73	5.35	6.09
Post Fossa	3.51	5.09	5.89	4.19	6.44	7.72
Bone	5.34	6.57	7.10	7.36	8.99	9.94

Slice Width Test

The CT linearity module of the ACR phantom contains two ramps which consist of wires that are visible in 0.5mm z-axis increments. The sets are slanted in the z-axis such that thinner slice thicknesses will show less wires in the reconstructed images compared to images of thicker slice thicknesses. The slice thickness is measured by counting the

number of visible wires for a given image reconstruction. Wire with less than 50% of their density should not be counted. Resolution of z-direction for each binning mode is set to be similar to Standard Resolution. It should be noted that this test is less accurate for smaller slice thicknesses because the wires are 0.5mm in length and the max resolution of the scanner in the z-direction is limited to 0.71mm. As Table 42 shows, each slice thickness measurement was within the final test specification of ±1.5mm of the nominal slice thickness. It should also be noted that in their latest update of the CT accreditation the ACR have dropped the slice sensitivity test from the CT accreditation requirements.

Nominal Slice	S	R	HR		UH	R
(SR/HR/UHR)	Axial	Helical	Axial	Helical	Axial	Helical
0.71/0.85/0.71	1.0mm	1.0mm	1.0mm	1.0mm	1.0mm	N/A
1.41/1.70/1.41	1.5mm	1.5mm	1.5mm	1.5mm	1.5mm	N/A
2.83/2.54/2.82	3.0mm	2.5mm	3.0mm	2.5mm	3.0mm	N/A
5.66/5.51/5.64	5.5mm	6mm	5.5mm	5.5mm	5.5mm	N/A
11.31/11.02/11.28	11.5mm	11.0mm	11.0mm	11.0mm	11.5mm	N/A

Table 42: The measured	slice	thickness
------------------------	-------	-----------

High Contrast Resolution

The high contrast resolution, or spatial resolution, describes the smallest distance between two high density objects. The spatial resolution depends mainly on the dimension of the detector. Module 4 of the ACR phantom is used to test high contrast resolution. It consists of eight sets of high-density wires arranged along the phantom periphery. Each set represents a different resolution (4, 5, 6, 7, 8, 9, 10, and 12 lp/cm counting counterclockwise from the 12 o'clock position).

Figure 144 shows the line-pairs patterns with FOV 308mm. Both scan modes using born kernel satisfy the ACR accreditation requirements of having at least 6 lp/cm pattern visible for the top images and the 8 lp/cm pattern for the bottom images. Figure 145 shows the line-pairs patterns with FOV 50mm. The 12 lp/cm pattern is visible in the UHR mode.



< Helical – SR >

< HR >

Figure 144: Sample images of the ACR phantom for Axial scan (top) and Helical scan (bottom) with 308mm FOV



Figure 145: Sample images of the ACR phantom for Axial scan with 50mm FOV

Noise Power Spectrum (NPS)

The NPS was measured for three typical axial and helical kernels that range from smooth to sharp. Noise images generated by taking the running difference of 10 consecutive scans of the 20cm water phantom were used to generate the noise power spectrum for each kernel. The normalized NPS of the three kernels is shown in Figure 146 which displays the spatial distribution of the noise over the entire image Field of View (FOV). It is observed that approximate frequency bandwidths for Soft Tissue, Post Fossa, and Bone kernels are 7, 8, and 12 lp/cm, respectively, in axial scans. In addition, the frequency bandwidth of the helical scan is 7, 11, and 12 lp/cm for Soft Tissue, Post Fossa, and Bone kernels. The bandwidths in the NPS plot match the expected frequency bandwidths which kernels are designed to have.



Figure 146: Normalized Noise Power Spectrum magnitude plot for the three reconstruction kernels for Axial and Helical scan modes (Top: Axial scan, Bottom: Helical scan)

Beam width

The ACR accreditation requires measurement of the Beam Width. The beam width is defined as the width of the irradiated region along the z-axis. The beam can be measured using a radio chromic film or strip that is properly positioned at the iso-center of the scanner. Figure 147 shows the measured beam width of the scanner. Due to the positional accuracy of the radio chromic film and the inaccuracy of the x-ray focal spot, the measured beam width of the scanner. 11.7 and 14.3 mm.



Figure 147: The measured beam width of the OmniTom Elite with PCD scanner

Geometric efficiency in the Z axis direction

The geometric efficiency is the ratio of the integral of the dose profile integrated over the detector width in Z divided by the total $CTDI_{100}$. The detector width in Z, or collimation, is 11.0mm for HR mode. The geometric efficiency is 75%.



Figure 148: OmniTom Elite with PCD dose profile

Slice sensitivity

The OmniTom Elite with PCD scanner provides three resolution modes. Row thickness and number are configured electronically. With reference to Table 32, the SR mode uses 16 detector rows with thickness 0.0706mm.



Figure 149 and Figure 150 show the bead in the ACR phantom and its profile in the Z-direction. Since the bead is effectively a high contrast impulse, its appearance in the series of thin images is a measure of the slice sensitivity profile in the Z-direction of the CT system.



Figure 149: Coronal reformation of the ACR scan using SR mode



Figure 150: Slice sensitivity profile of SR mode axial scan of the ACR bead phantom

Half-value layer

Table 43: Half-value layer

Scan voltage	70	80	100	120
Half value	3.8	4.5	5.7	6.8

Allowable variations

The following are allowable variations:

Dose	A \pm 5% variation in dose may occur due to variations between systems and measurement differences. The maximum variation is \pm 10%. The ACR required that the measured dose be within 20% of the measured dose.
High Contrast Resolution	The variation in values on the MTF curve may be $\pm 10\%$. These will occur mainly due to phantom placement errors, measurement inaccuracies and system variations.
Noise	The variation in standard deviation may be $\pm 10\%$ due to variations between systems.

Uniformity	The maximum difference between ROI means in an image is 4 HU.
Officiently	The maximum error in the CT number of water is ±3 HU.
Beam Width	The measured beam width should be between 11.7 and 14.3mm.

Scatter radiation

WARNING Exposure to secondary radiation can be harmful, and OmniTom Elite with PCD usage should only be done under the direct supervision of the facility's qualified Radiation Safety Officer (RSO) in compliance with site, local, provincial, and national regulations. Only this RSO can perform the calculations necessary to determine what additional safety precautions are necessary, such as shielding, personal protections, etc.

Note: The OmniTom Elite with PCD scanners are compatible with IRR1999 and EU Directive 96/29/EURATOM.

The OmniTom Elite with PCD is the latest addition to the NeuroLogica family of mobile scanners. The OmniTom Elite with PCD is a head scanner. The opening is 15cm bigger than its predecessor the CereTom to allow more diversified scan applications, for example, scanning the C-spine is now possible with the larger opening. The front cover of the scanner is self-shielded with 0.3mm lead. The scanner is also equipped with lead curtains in the front and the back. The back curtain will limit the scatter from the top of the scanned subject. The front curtains can be closed around the scanned patient to limit the side scatter.

The scatter is measured using the 16cm CTDI phantom. The scatter was measured as follows:

- 1. The phantom is placed in the beam with the laser positioned at the middle point of the phantom.
- 2. The back curtain is fully closed.
- 3. The front curtains are partially closed. The opening is limited to 50cm, to emulate the presence of a patient.
- 4. The scatter was measured using the RadCal 2026c dosimeter with the 1800cc chamber for low level radiation measurements.
- 5. The chamber is positioned at a height of 1.0 m from the floor.
- 6. Measurements were taken at 1.0 and 2.0 m from the fan beam location.
- 7. The data was collected between -90° and 90° at 45° increments.
- 8. The scan protocol used for measuring the scatter was:
- 9. The scan voltage: 120 kV
- 10. The scan current: 35 mA
- 11. The scan time: 2 seconds
- 12. Scan resolution mode is SR

Figure 151 shows the setup of the scatter measurement. The data is used to create the isodose curves. The data is in μ R. The data can be converted to mRem by multiplying by

0.877e-04 and to nSv by multiplying by 8.77. Table 44 shows the conversion rates between different radiation units.

Table 44: The energy	conversion rates
----------------------	------------------

	Multiply by
From μR to mGy	0.0000877
From μR to μSv	0.00877
From μR to mRem	0.000877
From mGy to μR	114025.0855
From mGy to mRem	100 mRem



Figure 151: The scatter measurement setup

Scatter exposure

Figure 152 shows the isodose curves generated from the above measurements. Figure 151 shows the measurement setup. The data is for a single axial scan position. In order to get the scatter for a longer scan, the data should be multiplied by the scan coverage. During a long scan, the scanner will be moving which might affect the measurement slightly.



Exposure isolines for 120 kVp, 70 mAs. Front curtain partially closed. Rear curtain fully closed. No patient absorption.



Figure 152: The isodose curves for 120 kV, 70 mAs using the CTDI 16cm phantom. The distances are in meters, and the scatter is measured in μ R.

Note: The scatter measurements were done without considering the scatter absorption through the patient or the patient bed. This can be simulated using water bottles stacked in the back of the scanned phantom. A body phantom can also be used to simulate the patient absorption. This should reduce the scatter in the front of the scanner. We have seen a typical reduction of up to four between the uses of a water or CTDI phantom and the body phantom. The body phantom that can be used is manufactured by the Kyoto Kagaku company. Figure 153 shows the body phantom.



Figure 153: The pediatric body phantom

Note: In compliance with IEC 60601-2-44, section 203.11, the previous figure shows the scatter radiation measured at the edge of the gantry in the tomographic plane is @20% of the scatter radiation measure at the same distance along the axis of rotation in the horizontal plane.

Additional scatter measurements

Additional scatter measurements along the perpendicular plane were also taken using the CTDI head phantom with the back curtain closed and the front curtains partially closed to simulate a real scan; however, it should be noted that the scatter measured with the CTDI phantom *only* is typically higher than the actual scatter radiation from a real patient. Due to the scanner height, the measurements were done at the positions noted in **Table 45** and

Table 46.

The scatter measurements were done using the RadCal survey meter and the 1800cc probe.

The scatter was measured using the following scan protocol: 120 kV, 45 mA, 10 seconds. The following table lists the measured scatter per second in the back of the scanner.

Table 15: Seatter in Vertical paraller to axis of rotation (pricings)				
	Distance from iso. (cm)			
	75cm	125cm	175cm	
50cm below iso	16.8519	7.1688	4.06725	
At iso	19.20525	8.61735	4.52835	
50cm above iso	10.94025	6.6729	3.9759	
75cm above iso	2.36205	4.6719	3.64095	

Table 45: Scatter in vertical parallel to axis of rotation (µRem/s)

	Distance from iso. (cm)			
	75cm	125cm	175cm	
At iso	0.428	0.155	0.075	
50cm above iso	0.375	0.149	0.069	
75cm above iso	0.274	0.140	0.070	

Table 46: Scatter	in vertical p	lane on pa	tient side	(mRem/	s)

Note: By design, this scanner's frame (about the bore) provides the primary shielding in the vertical plane perpendicular to the axis of rotation, therefore scatter radiation is negligible in this plane.

Dose linearity with tube voltage and current

The dose or radiation output of the tube is linear with the tube current. The exposures are measured at two different scans at isocenter with and without a CTDI phantom. The following table shows the exposure in the CTDI head phantom.

Table 47: Exposure at two different mAs

120 kV, 20 secs, head CTDI	Exposure at iso (mR)
30mA	4550
10mA	1467

Table 48: Exposure at two different mAs

120 kV, 4 secs, in air	Exposure at iso
30mA	1389
10mA	455

Two linearity factors are computed. The first is the ratio of the exposure at different scan currents; *F= Exposure (I1)/Exposure (I2)*. The linearity factor is then compared to the current ratio. The F linearity factor should be within 10% of the scan current ration.

$$L_1 = \left| \frac{E(l_1)}{E(l_2)} - \frac{l_1}{l_2} \right| \le 0.1 \frac{l_1}{l_2}$$

Table 49: Linearity calculations (mGy)

	CTDI head phantom	Air
E ₁ /E ₂	3.102	3.053
I ₁ /I ₂	3.0	3.0
L ₁	0.102	0.053
Linearity Test (< 0.1)	0.034	0.017

The second linearity factor is computed as described in the IEC 60601-2-44 standard.

$$L_2 = \left| \frac{E(l_1)}{l_1} - \frac{E(l_2)}{l_2} \right| < 0.2 \frac{\left| \frac{E(l_1)}{l_1} + \frac{E(l_2)}{l_2} \right|}{2}$$

Table 50: Linearity calculations in accordance with IEC

	CTDI head phantom	Air
E1/I1	151.67	46.3
E2/I2	146.70	45.5
(E1/I1+E2/I2)/2	149.19	45.9
L2	4.97	0.8
Linearity test (< 0.2)	0.033	0.017

The radiation output is not linear with respect to the scan voltage; however, it is approximately linear with respect to scan voltage to the power of 2.3, that is $E \propto kV^{2.3}$.

Note: Actual results on installed units can vary +20% due to machine and test tolerances.

Chapter 8 Registration

Registration is the first step in the patient scan process.

You can register a patient in the following ways:

- Manually register a patient for examination from the Registration tab.
- Perform a query to acquire already-entered patient data from the hospital (Hospital Information System (HIS)) or radiology system (Radiology Information System (RIS)).

It is assumed that the Tablet is connected to the site's **HIS/RIS** system. If you are not connected, you can always manually register the patient.

REGISTRATION	ACQUISITION	RECONSTRUCTION	VIEWER	BROWSER	1
				a a a a a a a a a a a a a a a a a a a	

Figure 154: Active registration tab

Navigating the Registration Screen

Make sure the Registration tab is selected; press it if necessary.

Notice the buttons at the top and bottom of the **Registration** dialog box. Many of these buttons are active *only* if you are already connected to the site's **HIS/RIS** or if you selected the **Query** button (to query for patients and the list of patients populated in the **Query Results** list). When a patient is selected, the buttons are active.

GUERY	SEARD+ MANJAL ADD					wi 🔪
WORKLIST						
Patient Name	Patient ID Date of Birth	Accession	Scheduled Date	Study Description	Stored List	Details
test, water			12/11/2024	NL Study CT	<	
test, gammex			12/18/2024	NL Study CT	 Image: A start of the start of	
test, test	1	121924 1009	12/19/2024	NL Study CT		
TEST, QA	1	1	11/13/2024	NL Study CT	 Image: A start of the start of	
TEST, QA		3	11/19/2024	NL Study CT	 Image: A second s	
TEST, QA			11/19/2024	NL Study CT	\checkmark	
TEST, PCD		2	12/18/2024	NL Study CT	\checkmark	
TEST, PCD			12/18/2024	NL Study CT	\checkmark	
water, axial			12/31/2024	NL Study CT	\checkmark	
watrr, helical			12/31/2024	NL Study CT	 Image: A start of the start of	

Figure 155: Registration dialog box

Registration buttons	Action	
GUERY	Searches the HIS/RIS server for scheduled patients. The population of patients could take several minutes to appear, depending on the number of patient entries the query retrieves after pressing the Query button.	
REGISTER	Registers the selected patient and then takes you to the Acquisition tab select a protocol to be used for scanning.	
	Shows selected patient details.	
SEARCH	Searches queried patient entries for specific information.	
	Selects patient(s) from query results and moves them into Stored List.	
	Removes patient(s) from the Stored List.	
	Manually enters a new patient including the Patient ID information, and when completed, takes you to the Acquisition tab to select a protocol to be used for scanning.	

Registering the patient

The following procedures show you how to enter patient information into the system before scanning the patient.

Querying patient information

1. If necessary, select the **Registration** tab on the main screen.

GUERY		TALAL ADD Dear	teal.	eel. Driar tert.	Select.		
WORLD	etometruet						
Potient Name	Patient ID	Date of Birth	Accession	Scheduled Cate	Boaty Description	Bearing.	-
Mark Harrill,	73565624	110000	78602600	510000			
Farmer, Ford	10110400	17/2000	78112708	112000			
Carle, Folw	20203051	11,0000	61306487	512000			
Lote: Stywatter	12001088	11/2500	91207248	512000		1	
Han, Sala	67710809	11,2000	47085860	1/1/2000			
Lein, Organa	39096817	11/2000	84370865	112000			
Berjaren, Kenda	102109417	11/2000	02402000	112000		-	
Michael, Linol	89124480	11/2500	13557968	512000		V	
Ross Calme	01726809	11,0000	06583192	512000		1	
Jesse, Riger	32907224	11,2000	45448577	512000			
						-	<u> </u>

Figure 156: Registration tab

2. Select the **Query** button at the top of the screen. The **Query Information** dialog box appears.



Figure 157: Query information dialog box

3. Select any of the named **Query Fields** you would like to use to query for patients by entering the value in the highlighted tabs.

A pop-up associated with the **Query Field** you are setting a value for appears. For example, if you press the **Scheduled Start Date** value row, the **Calendar** pop-up appears. Select the (new) date for the start. Another example would be to select the **Patient Name** value row. The **keyboard** pop-up appears so the user can type the patient's name to query.

The user can select any of the **Value** rows to fill in data to help query the patient you are searching. You can enter as much or as little information as needed. If no information is available, leave the value blank.

Pa	tient Name	Patient ID	Accession	Schedul	e Start Date		T
Enter to	ext	Enter text	Enter text	Se	lect		
				•	2016 -	2031	•
Birth	Accessio	on Schedu	Iled Date St	²⁰¹⁶ 2016	2017	2018	2019
				2020	2021	2022	2023
				2024	2025	2026	2027
				2028	2029	2030	2031
					/)	K

Figure 158: Query fields

- 4. When you are finished filling in query selections, perform one of the following:
 - Press the **Search** button to query based on the newly entered data to help narrow down your search.
 - Press the **Clear** button to remove any changes and return to the previous **Query Information** pop-up.
- 5. A list of patients matching your selected criteria variables populates in the Query Results list on the Registration tab.

Vicential and Reference and Re		
Parlent Name Patient ID Date of Birth Accession Scheduled Date Bruty Description		
		-
ark Hand, 7260634 110200 7700000 512200		
Sertex, Ford 70116480 1152500 78112788 1152500		
Carte, Failwer 20000031 5150000 85066487 5150000		
ake, Bryweker 12001088 1/12000 9/20/248 1/12000		
lan, Sale 67710809 1/1000 4706560 1/10000		
ekir Organa 30006817 1/1/2000 84370665 1/1/2000		
lengaren, Kunda 15076947 150000 00402588 5192000		
funael Linux 89124-991 11/2500 1368/369 512590		
	1	
Tues. Calue 01728889 1/12000 06551192 1/12000		

Figure 159: Registration query results table

- 6. Select a patient and select the **Register** button to register the patient for the exam.
- 7. The system enables and opens the **Acquisition** tab. To perform the acquisition steps, see "Performing A PCD Scan" on page 180.

Storing patients in the stored list

This list is helpful when multiple patients need to be scanned and connection to a worksite like **HIS/RIS** is unavailable at the exam location.

- 1. If necessary, go to the **Registration** tab to query the patients(s).
- 2. Perform steps 2 through 5 in "Querying patient information" on page 171.
- Select the Query button.
 Let the criteria you selected populate into the Query Results list area.
- 4. Select one or more patient entries from the Query Results list.

Select patients in the following ways:

- To select one patient, press the **Stored** icon located on the right side of the registration screen.
- To select more than one patient at a time, select the **Stored** icon for each patient you would like displayed in the stored list.
- 5. Select the **Stored** list.

The patients and subsequent patient information you selected appear in the Stored Results list on the Registration screen.

aun							
WORKLE	n Stranger						
Patient Rame	Patient D	Date of Birth	Accession	Scheduled Date	Study Centription	Beerler B	-
from the	A STREET	13,000	14002800	1/1/2000			-
			_			~	
				and the second se			

Figure 160: Registration stored results table

- 6. Select the patient you want to select from the **Stored Results** table.
- 7. Select the Register button to register the stored procedure in order to perform an examination.

The system enables and opens the Acquisition tab, which is discussed in more detail later on in this user manual.

Removing patients from the stored list

Patients can be manually deleted from the **Stored List** or if configured in Service, patients can automatically be removed from the stored list when the patient study is finalized. The pop up below will appear after finalizing a patient, if configured.



Figure 161: Stored list

Manually registering a patient using 'Manual Add'

You manually register a patient for examination when the **HIS/RIS** server is unavailable, the patient cannot be found, and/or was never entered into the system.

1. If necessary, go to the **Registration** tab.

2. Press the **Manual Add** button.

REGISTRATION ACCULISITION	RECONSTRUCTION	BROWSER
		0000000000000
GUERY SEARCH MANUAL ADD		

Figure 162: Manual Add button

3. The Add Patient Info dialog box appears.

First Name Last Name Date of Birth Accession: Patient	t ID Sex	
Enter text Select Enter text Enter text	Select	

Figure 163: Manually Add Pt Info Dialog Box

- 4. For **Patient Name**, press the **Patient Name** value. The **Keyboard** dialog box appears.
- 5. Enter patient name information in the fields provided and press the **Add** button to save your entries.

Notice that the value appears with the last name first, separated by ^, the first name next, separated by ^, the middle name (if you supplied that information), and any other information you entered.

- 6. For the **Patient Date of Birth**, perform the following:
 - Press and hold the **Patient Date of Birth** field.
 - Enter the patient's birth date in the **Patient date of birth** field using the **Calendar** pop up.
 - Select the Add button to save your work.
- 7. For the **Patient's Sex**, perform the following:
 - Press and hold the Patient's Sex field.
 - A drop-down list will display:
 - o Female
 - o Male
 - o Other
 - Select the **Patient's Sex** from the drop-down list and press the **Add** button to save your entries.

- Select the Register button to register your patient data. When you press the **Register** button, the system enables and opens the Acquisition tab.
- 8. After your patient is registered, view the **Patient Exam Details** to ensure your data is correct by selecting the **View Details** button.

Viewing patient information

This procedure lets you view (but not change) the patient's information.

- 1. If necessary, select the **Registration** tab on the main screen.
- 2. Select a patient from the Query Results list or the Stored Results list.
- 3. Press the **Details** button.
- 4. Review the patient's information. This pop-up presents static information that you cannot change.
- 5. Press the **Close** button to exit the **View Details Information** pop-up. The patients you selected in step 2 appear in the **Stored List.**

Chapter 9 Patient Scanning

Patient scanning overview

After registering the patient, the **Acquisition** tab is enabled and automatically opens. The **Acquisition** tab lets you check that the selected patient information is accurate before you perform the examination (scan). The **Acquisition** tab is also where you can set protocols for the scan before you scan the patient. A protocol lets you assess how you will capture the image you scan during the patient examination.



Figure 164: Active acquisition tab

After the protocol is selected, you can scan the patient by initiating the scan from the Tablet. See "Performing A PCD Scan" on page 180.

The following table provides information on the buttons on the **Acquisition** tab and what they are used for. Later you will learn how to set protocols for the scan.



CAUTION When conducting multiple or repeat scans, make sure the total exposure does not exceed maximum limit of 1Gy.

Table 52: Acquisition buttons

Acquisition buttons	Action
ACCEPT	Accepts the chosen protocol for the current study.

Acquisition buttons	Action
CONTINUE	Authorizes the scanner to move to the next step (if applicable).
BEGIN	Begins the countdown on the LCD to perform the study.
PAUSE	Allows the user to Pause the scan acquisition and then resume.
REPEAT	Allows the user to repeat a portion or all of the scan.
INITIATE EXPOSURE	Allows the user to start the exposure when using the Step and Shoot option.
RESUME ALL	Allows the protocol with Step and Shoot activated to resume the remainder of the acquisitions automatically.
MANUAL START	Allows the user to manually start the acquisition when doing CT Angiography protocols.
CANCEL	Allows the user to cancel the scout or scan.
PROTOCOL	Allows user to modify existing protocol selected in exam or choose a new protocol.

Identifying protocol types

Protocol types identify how to capture an image during a scan. The following are protocol types you can select from.

Axial

This protocol type lets you scan in **Transverse** plane. Data is acquired as the x-ray tube rotates around the patient.

Helical

This protocol lets you acquire data continuously as the x-ray tube rotates around the patient; the scanner translates over the patient in the Z axis.

Helical scan coverage

Helical scan coverage is typically truncated by 0.625cm to cover a half rotation at each end. "Figure 166: Scan coverage and imaged region for a true coverage of 60mm" shows the exposure using a radiographic film. The film shows exposure of the **Helical** scan-type coverage of 60mm. X-ray was on for 60mm. It took the scanner one extra second to completely stop after x-rays turned off. Figure 166 also shows exposure length to be 60mm.



Figure 165: Radiographic film of the 60mm scan coverage

The number of 0.625mm slices generated is 49, which covers about 60mm. The scanner parallel images start at the laser location and end where the scan ends. "Figure 165: Radiographic film of the 60mm scan coverage" shows the scan markers as described above. For a typical **Helical** scan with 23cm coverage, excess dose is 5.4% of scan dose.





Reference

This protocol lets you acquire a single 10mm slice to review anatomical position or place the **Region of Interest (ROI)** for **Bolus Tracking** scans. **Reference** scanning can only be used in conjunction with **Helical** scanning during a CTA protocol or perfusion.

Scout

This protocol lets you acquire data continuously as the x-ray tube remains stationary at a designated angle; the scanner translates over the patient in the Z axis. The resulting **2D** projection is used during examination planning.

Performing a PCD scan

You cannot complete this procedure without a registered patient.

Note: If the scan needs to be stopped, perform the following:

For an immediate or hard stop, press the **E-STOP** button. This stops x-ray, translate movement, and gantry rotation immediately.

For a controlled stop, press the Cancel button.

- 1. From the Tablet, go to the **Registration** tab to assign the patient to the scan in one of the following ways:
 - Query an already existing patient from the HIS/RIS.
 - Manually register the patient.

The **Acquisition** tab will be activated when the patient is registered.

2. From the **Acquisition** tab, select the **Adult** patient type. (Currently the PCD scanner does not support scanning of pediatric patients.)

Note: Protocol parameters are customized to meet your requirements in conjunction with local and nationally recognized published guidelines. These protocols *must be* approved by your facility physicist *before* the system's acceptance.

3. You can use the **Resolution** option to refine the available scanning protocols to select.
| ADUT PEDIATRIC | PROTOCOL | ORIENTATION | |
|----------------|-------------------------|-------------|-----|
| Resolution . | MULTI RES | | IFS |
| | NL Brain - Axial | | |
| | NL Brain - Axial UHR | | |
| | NL Brain - Axial HR | | |
| | NL Head Trauma | | |
| | NL Head Trauma HR | | |
| | NL Brain CT Angiography | | |
| | NL Brain CT Perfusion | NL Study CT | |
| | NL Sinus/Facial/Orbit | NEW | |
| | | | |

Figure 167: PCD Resolution options

4. Each Resolution provides different Slice Thickness and increment options as seen in the table below.

Standard Resolution (SR)	High Resolution (HR)	Ultra-High Resolutio (UHR) Axial scan mode onl
0.7 x 0.7 1.4 x 1.4 2.8 x 2.8 5.6 x 5.6 11.3 x 11.3	0.4 x 0.4 0.8 x 0.8 5.5 x 5.5 11.0 x 11.0	0.1 x 0.1 0.2 x 0.2 1.4 x 1.4 2.8 x 2.8 5.6 x 5.6 11.2 x 11.2

Table 53: PCD Slice Thickness Options

Note: Ultra-High Resolution scanning currently only supports a scan length of 67mm.

- 5. Select the desired protocol from the list.
 - The OmniTom Elite with PCD scanner allows multiple resolution scans to be planned and performed at the same time. The steps below include both a Standard and Ultra-High Resolution scan sequence.
- 6. Select the **Accept** button to review the selected protocol.



Figure 168: PCD accept icon

Note: The selected Resolution mode will be shown in the bottom right corner of the screens.

7. The Exam Planner dialog box appears.

	MITON RECONSTI MITOM, PCD		BROWSER	
Protocol Description AXIAL STD & UHR Anatomical Parls errors Head Study Description AXIAL STD & UHR Resolution Standard	Boout			
Tablet lé no	t cleared for diagnosis. Tablet is por	GN Dr use with pedialtric patients.	HR	

Figure 169: PCD Exam planner dialog box

The **Protocol Information** tabs displayed on the left and the **Protocol's Series** boxes displayed on the right show the series that are already created. The **Patient Position** is chosen before a protocol is selected.

Note: You can modify a protocol; however, changes you make from **Acquisition** will not be saved permanently. Permanent changes to protocols can only be made by the administrator in Protocol Manager.

Assuming you have the proper user privileges, you can modify protocol parameters such as, kV, mA, and coverage at the time of the scan, but the modifications will not be saved for future use.

- 8. To edit an existing protocol, perform the following:
 - Select an existing protocol series by pressing into the area within the box.
 - The selected **Protocol Series** will display.
 - Enter a description and make any other appropriate changes for your protocol.
 - Select the **Update** button in the **Protocol dialog** box.
 - Alternatively, select the Select Protocol button to return to the Exam Planner.

Protocol Description Adduction Protocol Description Adduction	ISTIN REC Partient card	INSTRUCTION Recent public Econ Trive 40 Counting 230 Number of Images Number of Images Standard	VEWER Mondow Wedth 135 Window Wedth 35 Kernel Soft Tissue Blem Trichness & Soft Tissue Soft Tissue Blem Trichness & Foctories & Pictures &	BROWSER Max CDP (mGY/GP1) 2000 Max CTD-vd (mgy) 1000 000X CTD-vd (mgy) 1000X CTD-vd (mgy)	
	toj clearod for diagnosa. Tabl	et is not far use with pedi	atric patients	HR	00

Figure 170: PCD protocol series dialog box

- 9. Set up the patient and scanner.
 - See "Positioning the patient" on page 82.
- 10. On the LCD Touch Screen on the scanner, press the Laser button to turn on the laser.
- 11. Move scanner and align patient as needed.
- 12. Select **Begin** on the Tablet from the **Exam Planner** dialog box.
- 13. The Ready. Press 'Scan' on scanner pop-up appears.



Figure 171: PCD Ready, press 'scan' on scanner

14. Press **Scan** from the LCD touch screen to acquire your scouts(s) and/or scan.

SCAN CANCEL	

Figure 172: PCD initiate scan on LCD

Note: During the scan, observe the following:

A yellow light on top of the scanner and an audible beep indicate x-rays are being emitted.

The scanner translates away from the patient.



15. After Scout or scouts are acquired, parameters can be set on the scout.

Figure 173: PCD setting scout parameters

- 16. To plan the first resolution, press the edge of the yellow box. A square will appear at the top right corner of the plan box, drag that square to modify the start and end location of this scan.
 - To plan the second resolution, press the edge of the green box. A square will appear at the top right corner of the plan box, drag that square to modify the start and end locations of this scan.

- You can also center the **Field of View** by selecting the circle on either plan box and dragging it to the desired location.
- 17. Select the **Continue** button to acquire the first scan or press the **Cancel** button to cancel the scan.
- 18. The Ready. Press 'SCAN' on scanner pop-up appears again.



Figure 174: PCD ready. Press 'SCAN' on scanner

- 19. Go to the scanner and press the **Scan** button to initiate the first scan.
- 20. To initiate the second scan, repeat steps 17 thru 19 above.
- 21. When the scans are complete the **Complete** dialog box will pop-up. Select **Yes** to finalize the patient and generate the **Dose Report.**
 - Selecting **No** will return you to the **Scan Complete** screen and would allow you to Repeat a scan or select a new protocol and continue scanning.
 - To Finalize the patient after pressing No on the Complete dialog box and generate the Dose Report, select the Patient Icon, then select Finalize.



Figure 175: PCD Complete dialog box

- Note: You must press the Finalize button before you can send the patient's data to PACS.
- **Note:** If you have additional reconstructions built into your protocol, you must finalize the scan before the additional reconstructions will be started.



Figure 176: Finalize before additional reconstructions start

Re-using Scouts with a New Protocol

1. Following the scan, select the **Protocol** button on the acquisition screen.



Figure 177: Scan Complete

2. Select the Start New button.



Figure 178: Start New

3. Select "Yes" to Re-use the Scout





4. Select new protocol and tap **Accept** button



Figure 180: Accept new protocol

5. Review Scout and scan parameters and select Begin.

Present Descriptor	tins.e	Padod	Name
Antonio Calendaria Antonio Calendaria Antonio Calendaria Antonio Calendaria			+
The Did			



6. Reset scout parameters, if needed, and select **Continue** to acquire the new scan.



Figure 182: Reused Scout

Repeating an Image

The repeat function can be used to repeat a scan if necessary. The entire scan can be repeated, or after reviewing the images, a new start position and coverage can be selected if only a portion of the scan needs to be repeated.

- 1. While the **Acquisition** tab remains active (before finalizing), click the **Repeat** button.
- 2. The **Repeat Protocol** pop-up appears.
- 3. Review the repeat scan coverage parameters.

You are o	poing to repeat the Axial. position or coverage	You can modify the Axial's before repeating.	start
	COVERAGE		
	Start Position	Coverage	
	45	181	
	CANICE	DEDEAT	
	LANLEL	HEPEAT	

Figure 183: Repeat scan pop up

Note: You can change the start position and coverage or use what appears.

- 4. Click the **Repeat** button, on the **Repeat Scan** pop-up.
 - The scanner will move to the start position.
- 5. Press the **SCAN** button on the scanner's control panel to begin the repeat scan.

Scanning with special features

The following features are available for use in protocols.

Using the step & shoot option

The **Step & Shoot** option in the protocol lets the user control the start of the scan acquisition. This is helpful in the case of an uncooperative or ill patient where motion is an issue.

- 1. If necessary, change the Scan Type to Axial.
- 2. Enable Step and Shoot by selecting the **Step & Shoot** button located in Options in the protocol.

OPTIONS	XAZiodow VAZidob
	Scan Delay
STEP & SHOOT	10
AUTOMATIC EXPOSU	RE CONTROL
	Noise Level
ENABLE AEC	0
MinmA	MaxmA
0	0
\checkmark	X

Figure 184: Step & Shoot dialog box

- 3. Select the check mark to save Step & Shoot setting.
- 4. Select **Update**.
- 5. Make sure your patient and scanner are positioned.
- 6. Press the **Begin** button to begin the scan.
- 7. The System 'Ready' to Scan message will appear.
- 8. Go to the scanner and press the Scan button on the LCD Touch Screen.
 - The first set of images are acquired at this position.
- 9. Press the **Initiate Exposure** button to initiate the next acquisition. To cancel the scan, select the **Cancel** button.
- 10. **Resume All** will turn off manual control of exposures and resume scanning automatically.

Performing a scan with Automatic Exposure Control (AEC)

Computed Tomography (CT) is responsible for the largest contribution to the collective effective dose to patients in radiology. The challenge to radiologists and medical

physicists is to establish adequate image quality with the lowest radiation exposure to the patient.

Tube current (mA) is one of the key technical scanning parameters for adjusting radiation dose in CT. To optimize radiation dose in CT, users can adjust mA either with manually selected values or with the application of Automatic Exposure Control (AEC). **AEC** refers to the automatic adaptation of mA on the basis of user specified image quality and attenuation characteristics of the scanned body region.

In OmniTom Elite with PCD, scout scans provide a graph of mA values based on object density and desired noise level. A single Axial or Helical scan in the protocol can utilize AEC, limiting the mA value of each slice to the minimum necessary to achieve the desired image quality. This ability to modulate the mA throughout the scan to achieve the desired noise level can reduce patient dose.

When using AEC, it is vitally important that the patient is well-centered in the gantry. AEC aims to deliver the specified image quality across a range of patient sizes. The use of AEC may change the planned $CTDI_{vol}$ and DLP values. It tends to increase $CTDI_{vol}$ for large patients and decrease it for small patients relative to a reference patient size stored in the scanner.

Note: Ensure patient is accurately centered in gantry. Do not use AEC when any type of metal is going to be scanned.
Do not use AEC with a small FOV; that is; tiny neonatal patients.
An automatic adjustment of the tube's current cannot occur when the tube potential is changed.
AEC can only be used with the Posterior Fossa kernel.

1. Enable AEC by selecting the ENABLE AEC button located in **Options** in the protocol.

OPTIONS	NA Ametowy VA Amtoh
	Scan Delay
STEP & SHOOT	10
AUTOMATIC EXPOSUR	RE CONTROL
End Position	Noise Level
ENABLE AEC	5
MinmA	Max mA
5	25
✓	X

Figure 185: Options dialog box with AEC enabled

- 2. Select the **Minimum mA** dropdown to set the minimum allowed mA value to be used for scanning.
- 3. Select the **Maximum mA** dropdown to set the maximum allowed mA value to be used for scanning. The available mA range is 5 to 20.
- 4. Select the **Noise Level** to set the standard deviation of the noise value for the acquired images. The available noise range is 1-200.

- 5. Select the check mark to save **AEC** settings.
- 6. Select Update.
- 7. Make sure the patient and scanner are positioned, and the patient is centered.
- 8. Select the **Begin** button to begin the scan.
- 9. Press Scan from the LCD touch screen.
- 10. After the scout(s) are acquired and the scan region is set, select the **Toggle AEC Graph Icon.**



Figure 186: Toggle AEC graph icon

11. The **AEC Graph** will be displayed on the scout. Review the mA modulation to ensure it meets your clinical needs.



Figure 187: AEC graph

- 12. To return to the scout parameter view, select the **Toggle AEC Graph Icon** again.
- 13. If the desired level is achieved according to your department policy, select the **Continue** button.

14. Press **Scan** from the LCD.

Note: While reviewing the scan you will see mA modulation as per your graphs.



Figure 188: mA modulation

Adding Noise Reduction to Axial Reconstructions

1. Enable **Noise Reduction** by selecting the **Options** button in the protocol.



Figure 189: Options

2. Select Noise Reduction

OPTIONS	
	Scan Delay
STEP& B-CCT	10
AUTOMATIC EXPOSU	RECONTROL
	Noise Level
ENABLEAEC	
MinmA	MaxmA
5	45
Noise Reduction Off	
✓	x

Figure 190: Noise Reduction

3. Select the desired level of **Noise Reduction**



Figure 191: Noise Reduction Levels

4. Select the **Check Mark** to save your selection.



Figure 192: Save

Note: Noise Reduction is only available for protocols using the Axial scan mode.

Performing a CT angiography scan with bolus tracking

CT angiography is a technique that uses contrast to visualize arterial and venous vessels throughout the body. This ranges from arteries serving the brain to those bringing blood to the lungs, kidneys, arms, and legs.

- 1. Perform steps 1 through 6 in "Performing a PCD scan" on page 180, selecting your approved CTA protocol.
- **Note:** Adult and pediatric protocol parameters are customized to meet your requirements in conjunction with local and nationally recognized published guidelines. These protocols *must be* approved by your facility physicist *before* the system's acceptance.
- 2. After selecting a CTA protocol, review Dynamic CTA options including **Delay, Bolus Mode, Contrast Threshold, Duration,** and **Injector Start** and make your selections.



Figure 193: Dynamic-CTA options

The following table explains the bolus tracking parameters and icons:

Table 54: Bolus tracking icons

Bolus Tracking Icons		Description
Duration	The amount of time allowed to monitor the bolus	
AutoStart	Begins the scan after the specified bolus scan time if no bolus is detected	
AutoStop	Stops contra	the scan after the specified bolus scan time if no ast is detected

TestBolus	A small amount of contrast is injected, and a timing graph is given after specified bolus scan time
Contrast Threshold	Hounsfield Unit detection at area being monitored - ROI
Manual	Manual Injector Start. The user is required to manually start both the scanner and the injector
Auto	Auto Injector Start. The scanner and injector will start when pressing start from injector

3. Perform scout(s) and set parameters.



Figure 194: Parameters on scout

4. The series boxes below show the steps in the CTA process.



Figure 195: Series boxes

5. To move the scout or reference parameter, select the appropriate series box to activate the scan region



Figure 196: Active scan regions on scout

- 6. Select **Continue** to acquire a Reference Scan.
- **Note:** A Reference scan is a single 10mm slice at a prescribed location. The Dynamic CTA is a monitoring scan and will be performed at the same location as the Reference scan.
- 7. Select the **Bolus ROI** tool.
- 8. Draw the ROI at the region you wish to monitor the bolus.



-

Table 55	Bolus	tracking	tool	table
Table JJ	. Doius	tracking	1001	lable

Bolus Tracking Tool	Description				
$\langle \overline{\mathbf{I}} \rangle$	Bolus ROI Tool				
$\overbrace{\longleftrightarrow}$	Moves Bolus ROI location				
	Resizes Bolus ROI				

- 9. Select **Continue**.
- 10. Load the injector and set the desired amount of contrast flow rate.
- 11. When the scanner is ready, press **Start** and manually start the injector at the same time, if you have chosen the **Manual Injector Start** option. If you have selected the **Auto Injector Start** option, press **Start** on the injector only.

Note: The Auto Injector Start is a purchasable option for the OmniTom Elite with PCD. Please contact your Customer Service Representative to purchase.

12. Scanner will trigger when bolus enters the reference point/Bolus ROI.

Note: During the Dynamic-CTA scan, if the Contrast Threshold is not reached, you can manually trigger the Helical CTA scan by clicking **Manual Start**.

13. Review completed scan.



Figure 198: Completed CTA

14. Select Finalize and perform reconstructions and MPRs as needed.

Performing test bolus

Test Bolus tests the timing of the contrast injected.

A small amount of contrast is injected, and a timing graph is given after the specified bolus scan time. When the contrast is detected, the system stops scanning and a report on the recommended delay time appears.

- 1. Select Test Bolus under the Bolus Mode options in the Dynamic CTA protocol.
- 2. Review Duration to determine length of **Test Bolus** monitoring time.
- 3. Select Update.
- 4. Select **Begin** to acquire the scout(s) and reference scan.
- 5. When the scanner is ready select **Start** on the scanner LCD.
- 6. Perform scout(s) and set parameters.



Figure 199: Parameters on scouts

- 7. Select Continue to acquire a Reference scan.
- **Note:** A Reference scan is a single 10mm slice at a prescribed location. The Dynamic CTA is a monitoring scan and will be performed at the same location as the Reference scan.
- 8. Select the **Bolus ROI** tool.
- 9. Draw the Bolus ROI.
- 10. Select **Continue** to begin Test Bolus.
- 11. Load the injector and set your desired amount of contrast (typically 10ml) and flow rate.
- 12. If you have chosen the **Manual Injector Start** option, once the scanner is ready, press **Start** on the scanner and injector at the same time. If you have selected the **Auto Injector Start** option, press Start on the injector only.
- **Note:** The Test Bolus graph will appear at the end of the prescribed duration time, showing the bolus tracking time. The recommended Scan Delay time for the CTA protocol will be displayed, as shown below in Blue.



Figure 200: Bolus timing graph

- 13. Select **Continue** to perform the scan with appropriate **Scan Delay**.
- 14. Review completed scan.
- 15. Select Finalize to complete.

Performing a CT perfusion scan

CT Perfusion (CTP) is a technique used to evaluate cerebral perfusion or the level of blood flow in the brain by monitoring the initial passing of iodinated contrast media through the vasculature of the brain.

- 1. Perform steps 1 through 6 in "Performing a PCD scan" on page 180, selecting your approved CTP protocol.
- 2. After selecting CTP protocol, review Dynamic CTP options including **Duration** and **Injector Start** and make your selections.



Figure 201: CT perfusion dynamic scan options



3. Perform scout and set dynamic CTP scan location.

Figure 202: Set dynamic CTP scan location

- 4. To move the Dynamic CTP location, move the yellow line using the circle on the reference line.
- 5. Select **Continue** to acquire the reference scan.
- Load the injector and set your desired flow and rate. When the scanner is ready, press Start and manually start the injector at the same time, if you have chosen the Manual Injector Start option. If you have selected the Auto Injector Start option, press Start on the injector only.
- 7. Review completed scan.



Figure 203: Review completed scan

8. Select Finalize.

Viewing images in the CTP viewer

Table 56: CTP Viewer Tools

CTP Viewer Tools	Description
Perfusion	Select to place the arterial and venous ROIs on the image
A	Hides the arterial ROI
$\langle \mathbf{v} \rangle$	Hides the venous ROI
Calculate	Select to calculate the CT Perfusion maps

Blending	Blends the original scan with the perfusion maps
$\langle \mathfrak{S} \rangle$	Reset
Capture	Captures a screenshot of the maps
Graph	Shows a graph of the arterial and venous blood flow
Layout	Changes the layout of the screen
	Allows the user to scroll through the images with their finger
$\langle 0 \rangle$	Brightness/Contrast
$\langle \overline{\mathbf{Q}} \rangle$	Zoom
	Pan
	Scroll Up through images
	Scroll down through images

Examining the scanned image with tools

You cannot work with image tools without a registered patient, the **Acquisition** tab enabled, and a scanned image showing. You can also use image tools from the **Viewer** tab – after you select a patient and open the associated images. In either case (from the **Acquisition** or the **Viewer** tab), the tools appear at the bottom of the screen and will illuminate when selected.

From the **Acquisition tab**, you can zoom, pan, modify window width, window level, and change layout; see the table below to understand the basics of what each tool looks like and how it performs.

Using tools on the acquisition tab

The following table describes some of the tools available to you when the **Acquisition** tab is active. For a comprehensive list, see Table 57.

Image tool	Tool name	Action
	Preset	Pre-defined Window Width and Window Level settings are located in a dropdown. Width and level presets can be saved or deleted by an administrator.
C)	Reset	Reverts all images back to their original mode.
	Page Down	Press and hold to page down through the images.
	Page Up	Press and hold to page up through the images.
	Toggle AP/PA	Displays scouts from Acquisition .

Table 57: Image tools

Image tool	Tool name	Action
	Toggle Lateral	Displays scouts from Acquisition .
$\langle \mathfrak{S} \rangle$	Toggle CT	Displays Axial or Helical scan from Acquisition.
	3 Panel Viewer	Displays Scout(s) and scan from Acquisition simultaneously.
	Toggle AEC Graph	Allows you to automatically adapt the tube current or tube potential according to the patient's body habitus in order to achieve the specified image quality at the lowest possible dose. AEC must be enabled in the protocol prior to the scan to use graph.
$\langle \overline{0} \rangle$	Window Width & Level	After selecting tool, press and drag in chosen direction to adjust image width and level.
	Pan	Allows you to move the image on the screen.
	Zoom	After selecting tool, press and move in upward direction to zoom in (enlarge) and downward to zoom out (shrink).
	Scroll	Allows the user to scroll through images slowly by dragging along the tablet surface.

Image tool	Tool name	Action
$\langle \mathbf{I} \rangle$	Bolus ROI	Allows the user to draw the ROI for use with bolus tracking.

Chapter 10 Browser

Browser overview

The **Browser** lets you view patient information and images associated with the patient information – after the patient's scan.



Figure 204: Active browser tab

The following table identifies the buttons found on the **Browser**.

Table 58: Command buttons

Button	Action
	Allows you to Edit Patient Details prior to sending to PACS.
	Selects the exam you want to archive.
×	Selects the exam you want to delete.
	Allows you to compare two series of images by tapping the compare arrow on each series you wish to compare.
ARCHIVE	Selects the archive destination for selected information.
DELETE	Deletes the selected exam information from the Browser tab.
PREVIEW	Allows the user to preview the selected exam information before being archived or deleted.

Navigating the browser

The Browser lets you perform tasks on existing series, for example archiving and previewing the series. This section will introduce you to the **Browser** and identify the symbols, areas, and buttons you can use.

The Browser can be broken down into the following sections:

- Studies Table
- Series Table

				cMurph	y, Randi	8 982	14534			N.
Studies										
Patient	Birth Date	Sex	Physician	Accession	Date	Time				
McMurphy Randle	19690504	NA	NA	98214534	2019/01/21	15:56:19.250	L ×	Scout	83	2019/01/ 3:56:19 I
Mickey, Mouse	NA	NA	NA	98214534	2019/01/21	15:25:39.439				
Gunderson Marge	19060509	NA	NA	08483495	2019/01/15	12:02:04.310		Image count: 1	PACS No	
Hudson William	19630713	NA	NA	53119816	2019/01/15	09:58:14.245				
Gunderson Marge	19060509	NA	NA	08483495	2019/01/15	00:00:00		Scout	83	2019/01/ 3:54:43 (
Doe, John	NA	NA	NA	78502800	2018/12/18	00:00:00				
Mickey, Mouse	NA	NA	NA	55555	0000/00/00	15:49:38.673	L X	Image count: 1	PACS NO	
						South Second		0000000000	0000000000	99999
Constant of the local division of the local	ARCHIVE			DELETE						2006

Figure 205: Browser studies and series tables

Selecting PCD Images in the Browser

Depending on the Material Decomposition reconstructions created from the scans, the patient browser will show multiple series of images, which can include those shown below.

Patient	Birth Date	Sav	Physician	Accession	Data	Time			
OMNITOM, PCD	NA	N/A	NA	TEST	2023/11/08	07:43 AM	1	×	
QUALITY	20231108	N/A	NA	QA	2023/11/08	07:41 AM		×	
TEST, PCD	NA	N/A	NA		2023/11/07	08:32 PM		×	Primary O
TEST, PCD	NA	N/A	NA		2023/11/07	07:37 PM	1	×	Irrage count: 10 PACS No 🔛 🛃
hel, tam	NA	N/A	NA	testhel	2023/11/07	11:54 AM	1	×	Axial 80% 000 2023
ax, tam	20170630	N/A	NA	12345	2023/11/07	11:09 AM	1	×	Iodine Map
tam, cta	NA	N/A	NA	testcta	2023/11/07	10:54 AM	1	×	Secondary
tam, cta	NA	N/A	NA	testcta	2023/11/07	10:48 AM	1	×	Irrage count: 30 PACS No
tam, cta	NA	N/A	NA	testcta	2023/11/07	10:45 AM	1	×	Axial 75 keV- 2023
tam, cta	NA	N/A	NA	testcta	2023/11/07	10:40 AM	1	×	
tam, cta	NA	N/A	NA	testcta	2023/11/07	10:37 AM	1	×	100000000000000000000000000000000000000

Figure 206: Material Decomposition results

Studies											
Patient	Birth Date	Sex	Physician	Accession	Date	Time			Axial Calcium	82	2023/06 03:18:52
PAULINE, PCD	20230621	N/A	NA	1234	2023/06/21	03:17 PM		×	TUU Map		
PCD, Pauline	19680520	N/A	NA	135577	2023/06/21	02:36 PM	2		Secondary		
CGUARD, HELICALCON	NA	N/A	NA	NA	2023/05/08	03:09 PM	1		Image count: 1	PAGS NO	
CGUARD, HELICALCON	NA	N/A	NA	NA	2023/05/08	02:57 PM	2		Axial Calcium	ൺര	2023/06
CGUARD, LC	NA	N/A	NA	NA	2023/05/08	12:22 PM	2		300 Map	89	
CGUARD, LC	NA	N/A	NA	NA	2023/05/08	12:17 PM	2		Secondary		
CGUARD, LC	NA	N/A	NA	NA	2023/05/08	12:08 PM	1		Image count: 1	PAGS No	
CGUARD, LC	NA	N/A	NA	NA	2023/05/08	11:56 AM		×	Axial Calcium	070	2023/06
									300+ Map	83	
									Secondary		
										annar Martha	44000

Figure 207: Calcium Maps

Compare Series

- 1. Select the Browser tab, if necessary.
- 2. Select the desired patient from the **Studies Table**.
- 3. In the **Series Table**, select the **Compare Arrow** for the first series to compare, then select the **Compare Arrow** for the second series.



Figure 208: Compare Series Arrow

4. The **Viewer** will display the two selected series side by side.



Figure 209: Compare Series Viewer

- 5. The active series is highlighted with a green border and can be changed by tapping the image.
- 6. Each series of images can be manipulated independently of the other using the image viewing tools.
- 7. The **Series Left** and **Series Right** buttons on the right side of the user interface can be used to select a different series from the study to display in that window.



Figure 210: Series Selection Buttons

8. When the **Series Left** or **Series Right** buttons are activated a dropdown menu appears in the selected series in the **Viewer** allowing selection of any additional series of images in that study.



Figure 211: Series Selection Dropdowns

Loading a series to viewer from the browser

- 1. Select the Browser tab, if necessary.
- 2. Select the desired patient from the **Studies Table**.
- 3. Select the desired series from the **Series Table**.
- 4. The selected series will autoload to the **Viewer** tab in **2D**.



Figure 212: 2D viewer

Deleting a series

- 1. If necessary, tap the **Browser** tab.
- 2. Select the study or the series to delete by pressing the **Delete Icon** located on the right side of the series or study.
- 3. The Delete **Preview** and **Clear** buttons will illuminate.



Figure 213: Delete pop up

- 4. Choose from the following:
 - Select **Preview** to review the study or series to be deleted.
 - Select Clear to deselect the study or series.
- 5. After selecting **Preview**, choose from the following:
 - Select **Delete** to delete study or series.
 - Select **Cancel** to keep the study or series.

Note: It is recommended to always archive to PACS prior to deleting from the browser.

Editing patient details

You can edit patient details in the browser prior to sending the data to PACS.

- 1. If necessary, select the Browser tab.
- 2. Select the patient in the Studies section
- 3. Select the Edit Patient Details icon. The Edit Details pop up will appear as shown below.

DIT PATIENT INF	D				
First Name	Last Name	Date of Birth	Accession:	Patient ID	
Susan	Johnson	Tue, 06/17/1952	13579	Enter text	CANCEL ENTER

Figure 214: Edit patient details

- 4. Edit patient details as needed.
- 5. Select Enter to save or Cancel, if needed.
- 6. All series data will be updated.

Archiving patient images

You can archive patients and studies (or series) to PACS.

Archiving to PACS

- 1. If necessary, select the Browser tab.
- 2. Select the patient study for **PACS** in the following way:
 - To select one patient and all associated series select the Archive icon from the Study Table.
 - To select specific series for a patient, select the **Archive** icon for each individual series from the **Series** Table.
 - The Archive **Preview** and **Clear** Buttons will illuminate.

ARCHI	VE
PREVIEW	CLEAR

Figure 215: Archive pop up

- 3. Choose from the following:
 - Select **Preview** to review the study or series to be archived.
 - Select Clear to deselect the study or series.
- 4. Select the PACS Server you wish to send to. You can have multiple servers configured.



Figure 216: Select the PACS server

- 5. After selecting the PACS Server, choose from the following:
 - Select **Archive** to archive study or series.
 - Select **Cancel** to keep study or series from being archived to PACS.
- 6. Click the **Archive** button to begin the archive process.
 - To verify the status of your image transfer, select System, then select PACS
 Queue. The PACS Queue dialog box will appear to show the status of your image transfer.
- 7. Watch the status of each series:
 - **Connecting** informs you that the series is in the process of archiving to its targeted location. Each series will move from the top portion of the pop-up to the bottom portion of the **PACS Queue** pop-up when it has been processed.
- 8. If the series is not successfully stored to its targeted destination, the "Store Failed" message appears in the **Failure** column. This means the series was not successfully archived. If the series was successfully archived, no message appears in the **Failure** column.



Figure 217: PACS queue dialog box

If the job is sent successfully, the series disappears from the queue.

Note: Any **Storing Failure** status appears as a pop-up to inform you why the failure occurred. If an archive job fails, it will be sent to the **Failures** list.

- 9. When the archiving is complete, click the **Close** button to exit the **PACS Queue**.
 - You can also click the **Close** button, and the archiving process will continue as you do other tasks.

Note: While the archive is in process, you can perform one of the following from the buttons in the PACS Queue dialog box. See the table below for a description of the buttons and their actions.

Table 59: Store/print queue buttons

Store and Print Queue button	Action
FAILURES	Shows the user the series that failed to archive. Select the Active button to return to the previous screen.
RETRY	Allows the user to Retry when an export fails.
ACTIVE	Shows the user the PACS Queue.
CLOSE	Closes the Store/Print Queue pop-up.
Chapter 11 Viewer

Viewer overview

The **Viewer** lets you see already scanned images from previous examinations. To view images, select the patient in **Browser** and then select the series to autoload to **Viewer**.

REGISTRATION	ACQUISITION	RECONSTRUCTION	VIEWER	BROWSER
anna an the second s			titititi	

Figure 218: Active viewer tab

The following table identifies the image tools from the **Viewing** tab that let you manipulate a scanned image. To see the following image tools, the **Viewing** tab must be enabled and open. Some image tools appear on specific viewing tabs, only. The view tabs are 2D, MPR, and 3D.

Table 60: Common tools

Image tools	Tool name	Action
PRESET	Preset	Changes Window Width/Level also includes presets of common WW/WL.
RESET	Reset	Reverts all images back to their original mode.
	Zoom	Magnifies the image.
$\langle \overline{0} \rangle$	Windowing	Adjusts window width and center of image. This icon remains active.
$\langle \mathbf{A} \rangle$	Pan	Adjusts image on X or Y axis.

INVERT	Invert	Inverts black to white and white to black.
$\langle - \rangle$	Line	Draws a line on the image and is used for measurement.
ROI	Region of Interest (ROI)	Defines a circular ROI and displays the ROI information.
RCRobal 10	ROI Radius	Allows you to type in a value for the placed ROI
GRID 1/1	Grid	Toggles layout from a 1 x 1 grid to 2 x 2 grid when selected.
	Page Up	Press and hold to page up through the images.
	Page Down	Press and down to page down through the images.
$\langle \mathbf{I} \rangle$	Scroll	Press and hold to scroll up or down through the images.
Annotation	Delete Annotation	Select to delete individual annotations.

Image tools	Tool name	Action
Enable Slab	Enable Slab	Enables slab lines on the anatomy.
SLAB	Slab	Allows you to change the Slab Thickness, Spacing and Rendering mode of the MPR series.
	Tilt	When selected a White 'steering' wheel allows you to correct a rotated image.
CAPTURE	Capture	Allows you to set WW/WL, enter a Series name and save the MPR series.

Table 61: MPR tools

Table 62: 3D tools

Image tools	Tool name	Action
Render Mode	Render Mode	Displays the image in Color, MIP, or Grayscale.
Reset orientation	Reset Orientation	Resets the image to the acquired orientation.
5	Rotate	Rotates the image.
Anterior	Anterior	Displays 3D image from the anterior view.
Posterior	Posterior	Displays 3D image from the posterior view.
Inferior	Inferior	Displays 3D image from the inferior view.

Superior	Superior	Displays 3D image from the superior view.
Left	Left	Displays 3D image to the left.
Right	Right	Displays 3D image to the right.

Setting window width and center

Note: Any modifications you make are not saved to the image.

- 1. Select a patient from **Browser**, select the series to view.
- 2. To open the image, click the **Viewer** button. The **Viewing** tab is enabled and open.
- 3. Click the **2D**, **3D**, **MPR**, or **CTP Viewer**.



Figure 219: 2D, 3D, MPR, and CTP viewing tab

To adjust the window width and/or level of the image, choose from the following options:

• To adjust with a preset, click the **Preset** button and select a preset: Abdomen, Angio, Bone, Brain, Chest, and Lungs.



Figure 220: Windowing preset list, text boxes, and the apply button

• To adjust the windowing width and level with the text boxes, enter the width and level values in the **Width** and **Level** text boxes, and click the **Apply** button.

Viewing images in 2D

2D lets you view scanned images in a **2-Dimensional (2D)** space. Standard **2D** mode is used when *only* one dataset is loaded. The default layout is a 1 x 1 grid.

Note: Any modifications you make are not saved to the image.

The **Viewer** tab is active immediately following acquisition. The current exam will autoload into the **Viewer** tab.

The 2D viewer will also open when you review a dataset from the Browser component.



Figure 221: 2D tools

- 1. Select a patient from **Browser**, select the series to view. The **Viewer** tab is enabled and opens to the **2D** tab.
- 2. Use any of the image tools to view the image differently (zoom, pan, invert, etc.) to manipulate your image.
- 3. Click the **Reset** button to reset images back to the original setting(s). You cannot undo this action.

Viewing or creating PCD keV images

1. Select a patient from **Browser**, select the series to view.

Studies										
Patient	Birth Date	Sex	Physician	Accession	Date	Time				
OMNITOM, PCD	NA	N/A	NA	TEST	2023/11/08	07:43 AM	🖉 🛓	Primary		
QUALITY ASSURANCE	20231108	N/A	NA	QA	2023/11/08	07:41 AM	2	Image count: 30	PACS NO	
TEST, PCD	NA	N/A	NA		2023/11/07	08:32 PM		Axial lodine	<u>972</u>	2023/11
TEST, PCD	NA	N/A	NA		2023/11/07	07:37 PM		Мар	8	
hel, tam	NA	N/A	NA	testhel	2023/11/07	11:54 AM	2	Secondary		
ax, tam	20170630	N/A	NA	12345	2023/11/07	11:09 AM	2	Image count: 1	PAGS NO	
tam, cta	NA	N/A	NA	testcta	2023/11/07	10:54 AM		Scout	\$ *2	2023/11/
tam, cta	NA	N/A	NA	testcta	2023/11/07	10:48 AM	1		8	
tam, cta	NA	N/A	NA	testcla	2023/11/07	10:45 AM		Primary		
tam, cta	NA	N/A	NA	testcta	2023/11/07	10:40 AM		Image count: 1	PACS NO	
	NA	N/A	NA	testcta	2023/11/07	10:37 AM		NOOCOOOC		

Figure 222: PCD Series

2. The Viewer tab is enabled and opens to the 2D tab.



Figure 223: PCD 2D Viewer

3. When loading Material Decomposition images, the 2D Viewer will include a keV and lodine button.



Figure 224: keV and Iodine buttons

- 4. Select the **keV** button.
- 5. This opens the keV dialog box in the 2D viewer.



Figure 225: PCD keV dialog box

6. Swiping up or down on the image changes the keV value displayed, keV values range from 40keV to 140keV in 1keV increments. The displayed keV value is shown in the keV field of keV dialog box.

- PECISITEATION
 ACCURTININ
 PECONSTRUCTION
 VENUE
 PROVISE

 Image: Construction
 Image: Const
- You can also press the keV value field to display a numbered keyboard and enter a value in the keV type in field to display a specific keV value.

Figure 226: keV type in

- The Save option allows a series of keV specific images, shown in the keV box, to be saved to the Patient Browser.
- The four-way arrow allows the keV dialog box to be moved anywhere on the screen.
- 7. To scroll through the images at the selected keV value, select any of the scroll buttons and swipe through the images on screen.



Figure 227: PCD keV Scroll buttons

Viewing or creating PCD lodine Overlay images

1. Select a patient from **Browser**, select the series to view.

Studies										
Patient	Birth Date	Sex	Physician	Accession	Date	Time				
OMNITOM, PCD	NA	N/A	NA	TEST	2023/11/08	07:43 AM 🖉		×	Primary 20	
QUALITY ASSURANCE	20231108	N/A	NA	QA	2023/11/08	07:41 AM 📝) 🛃	×	ingli com. 30	
TEST, PCD	NA	N/A	NA		2023/11/07	08:32 PM 💋) 💵	×	Axial lodine	2023/
TEST, PCD	NA	N/A	NA		2023/11/07	07:37 PM 🧪		×	Мар	
hel, tam	NA	N/A	NA	testhel	2023/11/07	11:54 AM 💋) 📘	×	Secondary	
ax, tam	20170630	N/A	NA	12345	2023/11/07	11:09 AM 🥖) 🛃	×	Image count: 1	PACS NO
tam, cta	NA	N/A	NA	testcta	2023/11/07	10:54 AM 🥖		×	Scout	2023/
tam, cta	NA	N/A	NA	testcta	2023/11/07	10:48 AM 📝		×		
tam, cta	NA	N/A	NA	testcta	2023/11/07	10:45 AM 🧪		×	Primary	
tam, cta	NA	N/A	NA	testcta	2023/11/07	10:40 AM 🧭		×	Image count: 1	PACS NO
tam, cta	NA	N/A	NA	testcta	2023/11/07	10:37 AM 🧪		×	N	

Figure 228: PCD Series

2. The Viewer tab is enabled and opens to the 2D tab.



Figure 229: PCD 2D Viewer

3. When loading Material Decomposition images, the 2D Viewer will include a keV and lodine button.



Figure 230: keV and Iodine buttons

- 4. Select the **lodine** button.
- 5. This opens the lodine dialog box on the 2D viewer.



Figure 231: PCD lodine dialog box

- 6. Swiping up or down on the image changes the lodine Overlay percentage displayed on the image, the lodine percentages range from 0% to 100% in 1% increments. The displayed lodine percentage value is shown in the lodine field of the lodine dialog box.
 - You can also press the lodine value field to display a numbered keyboard and enter a value in the lodine type in field to display a specific lodine percentage value.



Figure 232: Iodine percentage type in

- The Save option allows a series of Iodine Overlay percentage images, shown in the Iodine box, to be saved to the Patient Browser.
- The four-way arrow allows the keV button to be moved anywhere on the screen.
- 7. To scroll through the images at the selected Iodine Percentage value, select any of the scroll buttons and swipe through the images on screen.



Figure 233: PCD Iodine Scroll buttons

Viewing images in MPR

Multi-Planar Reformation (MPR) allows you to view the volume of images in Transverse (Axial), Coronal, and Sagittal planes.



Viewer layout is 2 x 2 as seen below.

Figure 234: MPR tools

- 1. Select a patient from **Browser**, select the series to view.
- 2. The series will autoload to the **Viewer**.
- 3. Select the MPR Icon.
- 4. Use any of the image tools to view the image differently (zoom, pan, invert, etc.) to manipulate your image.
- 5. The tilt or rotation tool can be used to modify the rotation of images.
- 6. Adjust the image angle by moving the circle clockwise or counterclockwise.
- 7. Click the Reset button to reset images back to the original setting(s). You cannot undo this action.

Understanding and using slab

Multi-Planar Reformation (MPR) allows images to be created from the original **Axial** plane into **Coronal**, **Sagittal**, or **Transverse** planes. **MPR** is fast, uses all the attenuation values in the dataset, and can easily be performed at the Tablet. **MPR**, however, provides only a **two-dimensional (2D)** display of the image data.

Sliding slabs are an additional technique used to create and save a series of MPR images in any plane. Through the reformation process, axial images are stacked creating a volume that can be dissected in different planes. The thickness and spacing of each dissection or slab can be varied to meet the needs of the viewer.

The reformations can be displayed in a minimum, maximum, or average projection.

Note: MPRs should be created from 0.625 or 1.25mm slices.

Creating the slab

- 1. Select a patient from **Browser**.
- 2. Select the desired series from the Series Table.
- 3. The selected series will autoload to the Viewer.
- 4. Select the MPR lcon.
- 5. Click the Sagittal, Coronal, or Transverse plane to create your slab.

To specify a stand	lard plane in	this vi ew , cl	lick the required image below.
	Coronal	Sagittal	Transverse

Figure 235: Coronal, sagittal, and transverse plane options

- 6. Select Enable Slab.
- 7. Set the **Cyan** lines to determine the beginning and end of the slab.



Figure 236: Set the cyan lines

8. Select Slab to set the thickness of the slices throughout the slab. Thickness and spacing can be entered manually or by selecting a pre-defined MPR Preset.



Figure 237: Slab option

9. Press the **Slab Rendering Options** dropdown to select the appropriate option.

The following options are available in MPR Mode:

Slab Thickness	The value that defines the thickness of the slab.
Slah Spacing	The value that defines the space between the start of one slab
Siab Spacing	and the next.
Slab Rendering Options	Where you define what pixel values are displayed in each slab:
	options include Average, Maximum Intensity, and Minimum
	Intensity.
Maximum Intensity	The highest pixel values for all slices within the slab displayed.
Minimum Intensity	The lowest pixel values for all slices within the slab is displayed.
Average	The pixel values of all slices within the slab are added up and the average value for each pixel is displayed.

Table 63: MPR Options

Vellow lines	Changes made to the slab thickness using the markers are
icitow inics	reflected in the value defined in the control panel.
	Define the slab FOV and dictate the range of a new series when
Cyan lines	generated. The cyan lines are adjustable by pressing and
	dragging on the lines themselves; both lines (at one time) are
	moved by clicking and dragging the central circle marker.
Crosshairs – Red,	Define the areas sections of the anatomy being viewed
Green, Blue	Define the cross sections of the anatomy being viewed.
Contune	Generates a new series with the name given in the Series
Capture	Description field, based on the selected MPR view pane.

10. Select the Tilt Tool to correct any rotation on the image



Figure 238: Using the tilt tool

11. Move the white circle on the image by placing your finger directly on the circle or just outside of it.

Note: The circle does not represent the Field of View.

- 12. The slab can be previewed in the bottom right viewport.
- 13. When you are ready to save, press the **Capture** button.



Figure 239: Capture dialog box

- 14. The **Capture** dialog box opens. Adjust Window **Width** and **Level** with the text box or use the windowing **Presets**.
- 15. Enter the series name and press **Apply** to keep settings.
- 16. To save the series press the **Capture** button. The new **MPR** image(s) appear in the **Browser** under the patient with the description (identifier) entered in the **Series Name** field.
- 17. To create additional MPRs, select the Reset button in MPR mode, select the MPR view you want to create and perform the steps above to create the new view.

Viewing images in 3D

In **3D** viewing, a **3-Dimensional** image is created by stacking all the images of a scan on top of one another to create a 3-D volume.



Note: Any modifications you make are not saved to the image.

Figure 240: 3D viewing

- 1. Select a patient from Browser and select the series to view.
- 2. To open the image, select the series and it will autoload to the Viewer.
- 3. Click the **3D** tab to dropdown the panel.
- 4. Use any of the image tools to view the image differently (zoom, pan, invert, rotate, etc.) to manipulate your image.

5. To rotate the image up to 360°, click (**Rotate**) and move the image by pressing on the screen to the rotation of choice. Use any of the image tools to view the image differently (zoom, pan, invert, rotate, etc.) to manipulate your image.

Viewing images in CTP

Computed Tomography Perfusion (CTP) is viewer software that enables your system to view dynamic scans to evaluate cerebral perfusion in the brain. The **CTP** viewer functionality is only possible if the optional perfusion package is set up and enabled on your system. A scan must have been performed with perfusion protocols.

- 1. Perform a Dynamic-CTP scan as outlined in Chapter 9 Patient Scanning.
- 2. Go to the Browser screen to load the Dynamic-CTP scan into Viewer.



Figure 241: Dynamic CTP viewer

- 3. Select CTP button on the left.
- 4. Select the **Scroll**, **Window Level**, **Zoom**, or **Move** viewer buttons at the bottom to set the viewer mode and display of the scanned image. Select and drag in the viewer to adjust the selected mode.
- 5. Select and hold the **Scroll Forward** or **Scroll Backward** buttons to scroll through the viewer images.
- 6. The CTP Viewer Tools are reviewed in the table below.

CTP Viewer	Description
Perfusion	Select to place the arterial and venous ROIs on the image.
A	Hides the arterial ROI.
$\langle \mathbf{v} \rangle$	Hides the venous ROI.
Calculate	Select to calculate the CT Perfusion maps.
Blending	Blends the original scan with the perfusion maps.
$\langle 5 \rangle$	Reset
Capture	Captures a screenshot of the maps.
Graph	Shows a graph of the arterial and venous blood flow.

Table 64: CTP Viewer Tools

7. Select **Perfusion ROI** button in the upper right.



Figure 242: Arterial and venous ROI's

- 8. Tap the scan image in the viewer to place the **arterial ROI**.
- 9. Tap the scan image in the viewer to place the venous ROI.
- 10. Select and drag the scale box on either ROI to adjust the size of the ROI.
- 11. Select and drag the move box on either ROI to adjust the position of the ROI.



Figure 243: Adjust position of ROI

- 12. Select the **A** (Arterial) and **V** (Venous) buttons to toggle the ROI's on and off or hide the respective ROI's if they overlap.
- 13. Select the **Calculate Map** button to calculate the perfusion map. After calculation completes a 4-port window displays 3 ports with the calculated maps and the original scan image.



Figure 244: Perfusion maps

- 14. Select the **Blending** button to activate blending mode. The mode displays the scan image overlayed on the calculated maps. Select the Window Level button and select and drag on the scan image to adjust the amount of blend.
- 15. Select the **Reset** button to set the viewer back to the uncalculated, starting state.
- 16. Select the **Capture** button to save a copy of the current scan image and perfusion maps to the browser.
- 17. Select the **Graph** button to display the Arterial Venous Flow values mapped over time.
- 18. Select the **Layout** button to view various layouts of the scan image and perfusion maps.

Chapter 12 Reconstruction

Reconstruction overview

The system stores multiple patient series of raw data to allow post reconstruction of images. **Reconstruction** allows reconstructing of the acquired data using different algorithms, slice thicknesses, or use of image enhancement algorithms, such as **Metal Artifact Reduction** and **Noise Reduction**.

Reconstruction Groups can be added to the protocol, as seen in Chapter 5 System Settings, or performed manually.

0.5x			
REGISTRATION ADDUSTION	RECONSTRUCTION	VIEWER	BROWSER

Figure 245: Active reconstruction tab

The tools available for **Reconstruction** are identified in the table below.

Table 65: Reconstruction tools

Image tools	Tool name	Action
LOAD ALL IMAGES	Load All Images	Loads image series into the viewer.
RESET	Reset	Reverts all images back to their original parameters.
$\langle 0 \rangle$	Windowing	Adjusts the width and level of selected image.
	Zoom	Magnifies the image.

Image tools	Tool name	Action
$\langle \Box \rangle$	FOV	Adjusts the Field of View (FOV) prior to reconstruction. *This icon will remain inactive until the user selects "Load All Images".
Delete	Delete FOV	Deletes the Field of View (FOV) *This icon will remain inactive until the user selects "Load All Images".
$\langle 1 \rangle$	Scroll	Press to activate – allowing you to scroll up or down through the images by swiping up or down on the image.
$\langle \mathbf{r} \rangle$	Page Down	Press or hold to page down through the image set.
	Page Up	Press or hold to page up through the image set.

Performing a Material Decomposition Image reconstruction

- Select a study in the **Studies** table. When you select a study, all of the scanned series for that study appear in the series table.
- 2. Select the series in the **series** table.



Figure 246: PCD Reconstruction studies and series tables

- 3. The settings table containing the available reconstruction parameter options will illuminate.
- 4. To view the study in the reconstruction viewing pane, select Load All Images.



Figure 247: PCD reconstruction viewing pane

- 5. From the Multi-Energy dropdown select the desired Material Decomposition result.
- 6. Select Add to List.



Figure 248: Material Decomposition Options

- 7. When you have selected all your desired results click **Begin** to generate the new dataset(s).
- Note: Prior to any Multi-Energy results being created you will see the system create the following Virtual Monoenergetic Image (VMI) datasets: 30-120keV
 40-120keV
 50-120keV
 These will not be saved in the Browser they are only used to create the Multi-Energy maps.
 Currently if Multi-Energy results are created, Noise Reduction and Metal Artifact

Reduction are not available.



Figure 249: PCD begin reconstruction

8. When reconstruction is complete, the image series appear in the **Browser.**

Chapter 13 Accessories and Options

Accessories and options overview

In this chapter, you will learn how to convert a bed, stretcher, or any type of adjustable surface into a scanning platform through the use of scan boards.

To request the catalog(s) to reference product descriptions/details and part numbers for the available accessories/options that are used with the respective scanner, see "Contact information" on page 23.

When using a fixed scanner, the table moves from one portion of anatomy to another while the gantry remains stationary. With the OmniTom Elite with PCD, an in-place scanning platform remains stationary while the gantry or scanner translates from one point to the other to cover the anatomy.

The universal transfer board is used for most beds or stretchers that are not compatible with the customer's bed adapter. It is placed under the patient and secured to the bed or stretcher with straps.

Note: The Universal Transfer Board may not be applicable for all beds.

A bed adapter is used as a secure mount to hold a silhouette scan board, which supports the patient's head. Different beds or scanning platforms require different adapters. NeuroLogica manufactures many different bed adapters to fit a wide variety of hospital beds, stretchers, and Operating Room (OR) tables. The majority of the bed and stretcher adapters are designed to hold a silhouette scan board.

WARNING

ING NeuroLogica Corp. recommends that the weight of the patient being positioned on the scan board does not exceed the bed manufacturer's safe, recommended, operating patient load. Realizing patient safety is of the utmost importance, it is recommended that safe judgment be exercised at all times when it comes to the clinical care of patients. There are a number of varying factors, such as the condition of the bed being used, unique patient anatomy, as well as the proper scan board and positioning of the patient, per NeuroLogica Corp.'s clinical training guidelines and product labeling. If any excessive wear or damage is noticed to any scan board, do not use it for a patient scan; contact a qualified service technician to assess, repair, and/or replace the device.

Using bed adapters

Bed adapters are manufactured for specific bed models. Prior to installing the OmniTom Elite with PCD at a facility, a precise survey is conducted to ascertain the type of bed used the most with the OmniTom Elite with PCD.

Adapters are classified as follows:

- Adapters that do not have posts and fit onto the frame of the bed by attaching to existing posts (below, left).
- Adapters that do have posts and fit onto the frame by inserting the posts into an existing hole in the frame (below, right).
- Adapters for OR tables, used in surgical cases.
- Adapters for a neonatal scan platform for children.



Figure 250: Bed adapter without posts (left) and bed adapter with posts (right)

Using bed adapter safety straps

The bed adapter comes with two straps that attach to the rings on the adapter and can be used to secure the patient to the silhouette scan board.



Figure 251: Safety Straps

1. Identify the rings on the bed adapter as highlighted on the Hill-Rom and Sytrker In Touch adapters below.



Figure 252: Hill rom bed adapter



Figure 253: Stryker In Touch bed adapter



2. Connect the hooks on the straps to the rings on the bed adapter as seen below.

Figure 254: Straps attached to bed adapter

3. Attach the silhouette scan board to the bed adapter.



Figure 255: Silhouette scan board attached to bed adapter

4. Position the patient on the Silhouette scan board and position the straps across the patients' shoulders and upper chest to provide support and immobilization.



Figure 256: Patient properly immobilized on scan board

Attaching bed adapter with or without posts

- 1. Check the bed adapter label and bed model to make sure the adapter is designed to fit the bed.
- 2. Remove the headboard from the bed.
- 3. Insert the bed adapter *without* posts onto the posts on the bed frame.





- 4. Make sure the adapter is flush against the bed frame.
- 5. Secure the adapter to the posts by adjusting the setscrew (see the yellow arrow in Figure 257).



Figure 258: Bed adapter T-square handle

6. Attach the silhouette scan board to the bed adapter by inserting the scan board into the adapter block.



Figure 259: Bed adapter without posts shown being inserted into scan board

Note: Ensure that the bed's IV pole does not obstruct or interfere with the Silhouette scan board. Ensure T-square handle is properly tightened: listen for two clicks.

- 7. Press down on the silhouette scan board while tightening the T-square handle until two (2) clicks are heard.
- 8. Add the cushion to the head holder for the patient's comfort.



Figure 260: Silhouette with cushion shown on head holder

9. The headboard is now ready to receive the patient.

Using the removable T-square handle with the bed adapter

- **Note:** If the desired scan starting point is limited by the handle touching the front cover of the scanner, the handle can be temporarily removed once the scan board is in place without disturbing its position.
- 1. Confirm that the scan board is set to the desired height and the T-Square handle is secured as outlined above and T-Square handle is tightened until you hear two clicks.
- 2. Pull the knob below the handle down, approximately 1/8 inch and while holding it down slide the handle straight out from the scan board clamps as shown below.



Figure 261: Steps showing removing T-square handle

- **Note:** Put the handle somewhere safe where it will not interfere with the scan or just let it hang down from the clamp.
- 3. To reattach the T-Square Handle, first ensure the wire rope is not wrapped around anything. Pull the locking knob below the handle location down and slide the T-Square Handle back into the clamp.
- 4. Keep hold of the handle and release the locking knob. You may need to rotate the handle a small amount, less than 1/4 turn, and you will feel the handle slide in further around 1/4", and the locking knob will snap into place as shown in the figures below.



Figure 262: Steps showing reattaching T-square handle

5. Confirm that the line on the handle shows that it is installed to the correct depth.



Figure 263: Correct install depth

Transfer boards for adult, pediatric, and neonate patients

Scan boards are customized for adult, pediatric, and neonate patients. Universal transfer boards and silhouette scan boards are used for adults; pediatric and neonate scan boards, and platforms are designed for smaller patients.

Scan board/platform	Weight
Neonate scan platform	7.5kg / 17lbs
Pediatric scan platform	24.9kg / 55lbs
Silhouette scan board	Weight limit of board is equal to the weight limit of the patient bed; weight limit on portion supporting patient head is 7.5kg / 17lbs
Universal transfer board	Weight limit of board is equal to the weight limit of the patient bed; weight limit on portion supporting patient head is 7.5kg / 17lbs

Table 66: 9	Scan boards	and their	weight-bea	aring re	estrictions
	Scan Souras	and then	weight bet	an ing i v	.50100115

See also "Parts that potentially come into contact with the patient" on page 67.



WARNING The weight limit for the superior portion of all scan boards (head) cannot exceed 7.5kg or 17lbs.

Using the universal transfer board and silhouette scan board

The universal transfer board and silhouette scan board are both carbon-fiber, radiolucent boards that are designed to work with any ICU bed or stretcher. The carbon-fiber board comes with a 0.5 in. (thick) headboard and 2 in. x 5 ft. straps to strap the board to the ICU bed or stretcher.

You can use the universal transfer board or silhouette scan board on most beds, tables, or stretchers. Because you can attach the universal transfer board to almost any type of surface, it is used anywhere throughout the hospital including the ICU, OR, and ER. The universal transfer board is placed on the mattress and secured with a strap or placed directly on a surgical table under the cushions. The patient lies on the board with the patient's head in the head holder. The OmniTom Elite with PCD is moved into position and the scan is performed.

The universal transfer board is always used with mattress stiffeners.

The mattress stiffeners provide a hard surface at the head of the bed to prevent the mattress from sagging when a scan is performed. There are usually four mattress stiffeners stored with the OmniTom Elite with PCD for easy transport.

Note: The universal transfer board and pediatric scan board are an optional accessory that does not come with the system.



Figure 264: Universal transfer board and stiffeners



Figure 265: Four types of mattress stiffeners

Note: Tipping of the board is a major concern. The universal transfer board **MUST** be securely fastened to the surface prior to placing the patient on the board.

- 1. Obey all warning labels when using the scan board.
- 2. Select the appropriate mattress stiffener for the mattress size and insert.



Figure 266: Mattress stiffener in place

- 3. The universal transfer board requires mattress stiffeners that provide a hard surface at the head of the bed to prevent the mattress from sagging with the weight of the patient when a scan is performed.
- 4. With the proper mattress stiffener properly inserted, apply the universal transfer board on top.
- 5. Position the board in accordance with the yellow, safety-warning stickers to avoid a tipping hazard. Do not extend the board beyond the mattress for proper placement.



Figure 267: Universal transfer board properly positioned on the bed on a mattress stiffener

- 6. When the board is properly positioned on the bed, secure it by using the safety strap.
- 7. The safety strap must be attached to the board, passed completely under the bed, and secured on the other side.



Figure 268: Universal transfer board with safety strap installed

- 8. When the universal transfer board is securely fastened to the bed, transfer the patient to the board, and secure the upper strap to the patient and the scan board.
- 9. When the patient is positioned and securely strapped in, position the scanner over the patient.
- 10. Initiate the scan.

Using the pediatric scan platform

Note: The Pediatric scan platform is used for larger children that cannot be supported by the infant/neonatal scan platform.



Figure 269: Pediatric scan platform

WARNING	The maximum weight limit for the Pediatric Scan Platform is 24.9kg / 55lbs.
WARNING	The pediatric scan platform is not a transportation device and should never be used to move a child from one location to another.
CAUTION	Always secure the child to the scan platform with the safety strap as described below to prevent motion and falls.
WARNING	<i>Never</i> leave the child unattended on the platform!

CAUTION Read and observe all warning labels.



CAUTION *Never* remove the scan board from the platform in order to use it separately.

Note: The pediatric scan platform is used to scan on adolescents too large to be scanned on the neonatal scan platform, but too small to be scanned on their regular hospital bed.

1. Place the child on the scan platform and secure with the safety strap.



Figure 270: Child placed on pediatric platform with safety strap (two views)

- 2. Position the scanner over the platform to ensure the patient is centered using the laser light as a guide.
- 3. After the scanner and patient are properly positioned, lock the platform in place by stepping down on the brake lever. Red to lock platform and green to unlock platform.



Figure 271: Platform brake


Using the infant and neonate scanning platform

Figure 272: Infant and neonatal scan platform

WARNING	The maximum weight limit for the neonatal scan platform is 7.5kg / 17lbs.
WARNING	The neonatal scan platform <i>is not</i> a transportation device and should <i>never</i> be used to move a child from one location to another.
WARNING	Always secure the child to the scan platform with the pediatric strap, as described in the following procedure, to prevent the child's motion and falls.
WARNING	<i>Never</i> leave the child unattended on the platform!
CAUTION	Read and observe all warning labels.

Positioning using the infant and neonate scanning platform





Figure 273: Pediatric strap

2. Drape the strap over the patient and secure one side around the patient with Velcro, then continue with other side.



Figure 274: Applying pediatric strap onto the platform (three views)

The neonate or infant patient can be positioned on the platform head-first for head scans, and feet-first or head-first for body scans.





Figure 275: Proper position of neonate/infant for head scans (left) and body scans (right)

3. The patient is now ready to be placed in the scanner.

Note: The platform remains stationary during the scan while the scanner moves during the scan.

- 4. Position the scanner over the platform to ensure the patient is centered using the laser light as a guide.
- 5. After the scanner and patient are properly positioned, lock the platform in place by stepping down on the plungers located on both sides.



Figure 276: Neonate platform foot brake location

Adjusting the infant and neonate platform handle

The Infant and Neonate Platform Handle is used for transporting and moving the imaging platform. The handle is adjustable and may be moved to accommodate for more space or user preference.



WARNING The neonatal scan platform *is not* a transportation device and should *never* be used to move a child from one location to another.

1. To adjust the handle, remove the two locking pins located on the insides of the handle by pressing down on the pin buttons and pulling back (see yellow boxes below).



Figure 277: Locking pins for the handle

2. Position the platform handle in one of the three set locations (see figure below).



Figure 278: Three positions for the platform handle

3. Once adjusted to the desired position, secure the handle in place by inserting both of the locking pins into the slots. When pins are at proper depth and correctly seated, they will lock into place.



WARNING Ensure both pins are locked in a secure position prior to moving!

Chapter 14 Using the OmniTom Elite with PCD in the Operating Room

The three methods used to convert the operating table into a scanning platform are as follows:

- An OR table adapter with the silhouette scan board
- Universal transfer board
- Doro[®] OmniTom Elite with PCD intra-operative cranial stabilization system

Each of the above-mentioned devices are covered in the following sections. It is important to note that all these devices are used pre, post, and intra-operatively.

Using the operating room table adapter

The Operating Room (OR) table adapter functions in the same way as a bed adapter.

1. Attach the OR table and then the silhouette scan board to the adapter.

The OR table adapter has two posts that fit into the adapter supplied by either the OR bed manufacturer or by Mayfield.



Figure 279: OR table adapter

2. Adjust the OR table adapter to the openings in the mount and attach it to the mount.



Figure 280: OR table adapter attached to mount

- 3. Attach the silhouette scan board to the OR table adapter.
- 4. Add the padding to the head holder and slide the patient onto the board.

You can perform this step at scan time or before the start of surgery. If you perform it prior to surgery, the surgeon may choose to perform the surgery on the head holder.

5. Drive the scanner into place to start the study.

Using the universal transfer board in the OR

The universal transfer board is a carbon-fiber, radiolucent board that is designed to work with most ICU beds or gurneys. The carbon-fiber board comes with a 0.5 in. (thick) headboard and 2 in. x 5 ft. straps to strap the board to the ICU bed or gurney. It also comes with 3 in., 4 in., 6 in., and 12 in. bed stiffeners.

In addition to using the OR adapter and silhouette board in the OR, the universal transfer board is often used in the OR. The universal transfer board is secured to the surgical table by placing the board under the table cushions (if used) and securing the safety strap to the table by passing the strap under it and securing it on the other side. The board is secured to the table prior to the patient's arrival. The patient is then placed on the table with the patient's head in the head holder. The surgery is performed with the patient on the universal transfer board. When a scan is needed, the OmniTom Elite with PCD is driven into place over the patient's head and the scan is performed. The universal transfer board is used for almost any non-invasive surgical procedure.

Appendix A Glossary

Α

Algorithm	Mathematical filter applied to raw data during CT image reconstruction to remove blurring artifact inherent to back-projection. Also referred to as a kernel
Annotation	User comments or text added to an image.
Anterior	Front of the patient's body
Application Entity (AE)	An end point of a DICOM information exchange, including the DICOM network or media interface software; that is, the software that sends or receives DICOM information objects or messages. A single device can have multiple AEs.
Attenuation	The reduction in intensity of a radiation beam as it passes through a substance.
Automatic Exposure	Software used to adjust or modulate the mA
Control (AEC)	throughout an acquisition to reduce patient radiation dose to a minimum.
Axial scan mode	Data acquisition while the scanner remains stationary. The scanner position may be incremented between exposures to collect data over a longer Z axis range.

В

Beam hardening	The phenomenon whereby low energy photons are absorbed as the x-ray beam passes through an object, resulting in an increase in the average photon energy of the beam.
Bolus tracking	Monitors flow of contrast media in vessel and triggers scan at optimal timing. This is a scanner feature to automatically initiate a prescribed Helical scan when a threshold level of contrast enhancement is reached at a specified region of interest.

С

Center X, Center Y	Reconstruction coordinates of an image.
Collimation	Restricts x-ray to only the selected anatomy,
	minimizing dose to patient and reducing scatter.
Computed Tomography	A test that uses x-rays to provide detailed pictures of
Angiography (CTA)	the heart and the blood vessels that go to the heart,
	lung, brain, kidneys, head, neck, legs, and arms. A CT

	angiogram can show narrowed or blocked areas of a blood vessel.
Computed Tomography Dose Index (CTDI)	An approximate measure of the radiation dose received in a single CT section or slice.
Computed Tomography Dose Index Volume (CTDI _{vol})	Represents the dose for a specific scan protocol, which takes into account gaps and overlaps between the radiation-dose profiles from consecutive rotations of the x-ray source. The CT dose index volume is noted as CTDI _{vol} . The CTDI _{vol} is calculated differently for both the Axial and the Helical mode: 1. For Axial scan mode: CTDIvol = [(N x T)/I] x CTDIw 2. For Helical scan mode: CTDI _{vol} = 1/pitch x CTDI _w
Computed Tomography Dose Index (CTDI _w) weighted average	The measure of ionizing radiation exposure per slice of data acquisition. CTDI represents the integrated dose along the Z axis from one axial CT scan (one rotation of the x-ray tube). The CT Dose Index is noted as CTDI _w .
Computed Tomography (CT) number	Relative value assigned to each pixel to quantify the attenuation occurring in each pixel in comparison with the attenuation of water. The calculated CT number for a given pixel is given in Hounsfield units (HU).
Computed Tomography Perfusion (CTP)	Evaluates cerebral perfusion or level of blood flow in the brain by monitoring the initial passing of iodinated contrast media through the vasculature of the brain.
Contrast media	Used to improve sensitivity and specificity of clinical diagnoses.
Contrast resolution	The ability of a CT system to detect an object with a small difference in linear attenuation coefficient from the surrounding tissue. Also referred to as low-contrast detectability or sensitivity.

D

Digital Imaging	Digital Imaging and Communications in Medicine, or
Communication in	DICOM, is a standard that helps people doing work in
Medicine (DICOM)	the field of radiology. The DICOM standard is designed
	to promote communication and integration between a
	variety of radiology imaging systems and equipment
	used in filmless radiology.
Digital tilt	The ability to correct the image post-acquisition and
	correct positional inaccuracies prior to sending to
	PACS.
Dose	Amount of ionizing radiation absorbed by patient per
	unit mass.
Dose Length Product (DLP)	The measurement of dose for an entire series of CT
	images. DLP is equal to the calculated dose per section

	multiplied by the length of a CT acquisition along the
	Z- axis.
Dynamic Host Control	A standardized network protocol used on Internet
Protocol (DHCP)	Protocol (IP) networks. The DHCP is controlled by a
	DHCP server that dynamically distributes network
	configuration parameters, such as IP addresses, for
	interfaces and services.
Dynamic scan mode	Data acquisition at multiple time points over the same
	anatomic location(s).

Electromagnetic	The branch of electrical sciences that studies the
Compatibility (EMC)	unintentional generation, propagation, and reception
	of electromagnetic energy with reference to the
	unwanted effects (Electromagnetic interference (EMI))
	that such energy may induce.
Electromagnetic	A disturbance generated by an external source that
Interference (EMI)	affects an electrical circuit by electromagnetic
	induction, electrostatic coupling, or conduction. The
	disturbance may degrade the performance of the
	circuit or even stop it from functioning.

Η

Field of View (FOV)	The diameter of the acquired images displayed across
	the image matrix.

Helical	A CT acquisition where the x-ray tube and scanner move continuously during scanning, yielding a data set in the form of a helix. Also referred to as spiral.
Hospital Information System/Radiology Information Systems (HIS/RIS)	A Radiology Information System (RIS) is the core system for the electronic management of imaging departments. The major functions of the RIS can include patient scheduling, resource management, examination performance tracking, examination interpretation, results distribution, and procedure billing. RIS complements Hospital information systems (HIS) and Picture Archiving and Communication System (PACS) and is critical to efficient workflow to radiology practices.
Hounsfield Unit (HU)	The unit of the CT number scale assigned to each pixel to quantify relative attenuation.

Glossary

Interscan delay time	Minimum amount of time that must transpire between end of one scan and initiation of next scan. Interscan delay times include idle time between scans to allow tube cooling.
lodine Map	lodine Map images display ONLY pixels that contain lodine, any pixels that do not contain lodine are replace with black pixels.
Iterative Bone Correction (IBC)	A feature build into the reconstruction software, which performs a correction on every Axial image the scanner produces, including both primary series from a scan as well as secondary reconstruction images. Current IBC settings were chosen to provide optimal correction for standard medical imaging; however, the setting can be customized as needed.

Kernel	A mathematical filter applied to raw data during CT
	image reconstruction to remove blurring artifact
	inherent to back-projection. Also referred to as an
	algorithm.

Liquid Crystal Display	A form of visual display used in electronic devices in
(LCD)	which a layer of a liquid crystal is sandwiched between
	two transparent electrodes. The application of an
	electric current to a small area of the layer alters the
	alignment of its molecules, which affects its
	reflectivity or its transmission of polarized light and
	makes it opaque. An LCD is used on the scanner and is
	a touch screen.

Μ

mAs	Tube current-time product: The product of tube
	current and exposure time per rotation, expressed in
	units of milliampere seconds (mAs).
Matrix	Two-dimensional grid numbers arranged in rows and
	columns.
Maximum Intensity	The multiplanar reformation technique that displays
Projection (MIP)	only the maximum pixel value along a ray traced
	through the object to the viewers assumed
	perspective in front of the viewing monitor.
Mean Transit Time (MTT)	A common measurement during CT perfusion studies
	of the brain. Refers to the average transit time, in
	seconds, needed for blood to pass through a given
	region of brain tissue.

L

milli amperage (mA)	Tube current: the number of electrons accelerated
	across an x-ray tube per unit time, expressed in units
	of milliampere (mA).
Modality Performed	A mechanism for modalities to pass information about
Procedure Step (MPPS)	the imaging performed back to the HIS/RIS or PACS.
Modality Worklist	Schodulad (but not vat coonned) nations list
Manager	Scheduled (but not yet scanned) patient list.
Motion artifact	Voluntary and involuntary patient motion during CT
	scan, appearing as a streak artifact on image; ghosting
	or blurring of image.
Multiplanar Reformatting	The process of displaying CT images in a different
(MPR)	orientation from the one used in the original
	reconstruction. Allows for reformation of images in
	planes that would otherwise be difficult or impossible
	to acquire with CT. Requires only image data. Raw
	data is not utilized.

Noise	Random statistical variations in the signal. Can be
	quantum noise, electronic noise due to lost signal, or
	artifact noise. Manifests itself as overall graininess of
	the reconstructed image.

Ρ

Partial volume artifact	Occurs when an object is only partly positioned within a voxel or is much smaller than the overall voxel volume. The object's attenuation is not accurately represented by the pixel value. Overlapping reconstructions further reduce partial volume artifacts.
Patient coordinates	References are as follows:
	X left to right.
	Y anterior to posterior.
	Z head to feet.
PCD	Photon Counting Detectors
Peak kiloVoltage (kVp)	The penetrating power of the photons coming from
	the x-ray tube.
Picture Archive and	Stores medical information, including 2D images, and
Communications Systems	3D medical images. All modern PACS setups will work
(PACS)	with DICOM.
Pitch	In Helical mode, refers to the speed of the scanner
	movement over the table as the scanner rotates.
Pixel	A single, picture element of image matrix.
Post reconstruction	Prescribing the reconstruction parameters after scan acquisition.

Projection	View of anatomical cross-section from a particular
	vantage point.
Prone	Patient lying on stomach.
Protocol	Prescribes the acquisition and reconstruction
	parameters to be used for a scan.

Q

Quality Assurance (QA)	Procedure of performing periodic specified tests or
	measurements to assure that a set quality level, as
	specified by system manufacturer, has not been
	compromised.

R

Radiation Safety Officer (RSO)	The person within an organization responsible for the safe use of radiation and radioactive materials as well as regulatory compliance.
Radio Frequency Interference (RFI)	Also called Electromagnetic Interference (EMI), is an unwanted disturbance that affects an electrical circuit due to electromagnetic radiation emitted from an external source.
Raw data	A transmission measurement obtained by the detectors used to mathematically reconstruct the CT image.
Reconstruction filter	Used to ensure accurate anatomical image reconstruction. Also allows for either spatial resolution or low-contrast-resolution enhancement.
Region Of Interest (ROI)	Provides a quantitative analysis of the Hounsfield values of a specific anatomic area. A graphic outline in the shape of a circle is placed over an area on the image. Software calculates the average CT number in HU within the ROI.
Resolution	A scan time, per slice, in Axial mode, only.
Retrospective reconstruction	Reconstruction performed after the initial prospective reconstruction. Multiple retrospective reconstructions of raw data are possible, with changes to display FOV, kernel, slice thickness, etc.

S

Scan delay	The time between the initiation of contrast agent
	administration and CT data acquisition. The chosen
	scan delay determines the phase of contrast
	enhancement for a given CT acquisition.
Scan protocol	A list of scanner-load parameters used to perform an
	x-ray exposure.

Scan types	Axial, Helical, Dynamic, Reference, and Scout.
Scout	Digital survey radiograph acquired by the CT system
	for the purpose of prescribing the cross-sectional
	acquisition. Similar to a conventional radiograph, the
	scout is produced by translating the scanner over the
	patient without tube or detector rotation. Also
	referred to as topogram or scanogram.
Series	A set of images acquired in a scan.
Slice spacing (Spacing)	The distance between the center of one CT slice and
	the center of the next slice.
Slice thickness	The dimension of a constructed CT slice along the
	longitudinal direction of acquisition (Z axis).
Spatial resolution	The ability of a CT imaging system to display fine
	details, separately. Given in units of line pairs per
	centimeter (lp/cm).
Supine	Lying on back.

Temporal resolution	The ability of a CT system to freeze motion and
	provide an image – free of blurring.
Test Bolus	Scan mode used to measure the contrast transit time
	using a small injection of contrast media.
Threshold	The CT number (Hounsfield Unit (HU)) where bolus
	tracking tool will trigger the system to begin the scan.
Time Attenuation Curve	A graph of the contrast enhancement versus time. TAC
(TAC)	is used to determine blood flow rate in seconds for
	contrast timing.
Time delay	Monitoring delay: Time from injection to the start of
	monitoring scans.
Transverse plane	Perpendicular to direction of Z axis.

V

Т

Virtual Mono-Energetic	A series of synthesized images resulting from a
Images (VMI)	monoenergetic X-ray beam focused on a single
	energy in the range of 30 to 140keV.
Virtual Non-contrast	VNC images allow visualization of contrast enhanced
Images (VNC)	tissues after the removal of iodinated contrast
	material. Iodinated pixels are identified and replaced
	by HU values as if the iodine material is not present.
Volume Rendering (VR)	A 3D modeling technique that utilizes the entire
image or object	acquired dataset but adjusts the opacity of the voxels
	included in the 3D image according to their tissue
	characteristics.

Voxel	Abbreviation of volume element. Refers to the
	volume of tissue represented by a pixel in the matrix
	used to display the CT image.

W

Window Level (WL)	The pixel value given in Hounsfield Units (HU) at the center of the window width. Window Level controls
	the brightness (density) of the CI image.
Window Width (WW)	The range of pixel values assigned a shade of gray in
	the displayed CT image. Window Width controls the
	contrast of the CT image.

Appendix B Error Codes

Error- code number	Fault description	Explanation	Cause ¹	Resolution
0	NO_FAULT	Success. No error occurred	N/A	N/A
1	ACQUISITION_T ERMINATED	Not an error.	Acquisition has terminated normally (or user hit Cancel)	N/A
2	XRAY_TEMPER ATURE_FAULT	Temperature fault was detected at X- ray tube	X-ray Tube overheated	Allow time for X-ray Tube to cool down
3	XRAY_ARC_FAU LT	An X-ray tube arc was detected	An arc occurred in X-ray Tube or the HV Generator	Contact Technical Support for service
4	XRAY_HIGH_M A_FAULT	Monoblock high mA condition detected	Beam current exceeds set value by more than 5% for 100 ms or more	Contact Technical Support for service
5	XRAY_LOW_MA _FAULT	Monoblock low mA condition detected	Beam current is less than 95% of set value for 100 ms or more	Contact Technical Support for service
6	XRAY_LOW_KV _FAULT	Monoblock low kV condition detected	X-ray Tube voltage is less than 95% of set value for 100 ms or more	Contact Technical Support for service

¹ There may be multiple causes that require a trained service technician to conduct an analysis and repair.

Error- code number	Fault description	Explanation	Cause ¹	Resolution
7	XRAY_HIGH_KV _FAULT	Monoblock high kV condition detected	X-ray Tube voltage exceeds set value by more than 5% for 100 ms or more	Contact Technical Support for service
8	XRAY_WATCHD OG_FAULT	Monoblock watchdog timeout condition detected	Watchdog Timer was not refreshed at a high enough rate	Contact Technical Support for service
9	XRAY_POWER_ LIMIT_FAULT	[Placeholder for power limit fault]	N/A	N/A
10	XRAY_INTERLO CK_FAULT	Interlock was de-asserted to generator	E-Stop was activated	Deactivate E- Stop
11	XRAY_BELOW_ THRESHOLD	Reference detector values are reading less than threshold value	Indicates X-rays have been turned off (due to errors 2-10 above)	Contact Technical Support for service
12	A subsystem failed to respond in a timely manner. Please retry the operation	The Operating Systems are not (Recon and Tablet) are not communicating properly	The Reconstruction computer timed out when trying to execute the command	Try the operation again or restart the scanner
13	Unable to Prepare a scan. Recon is disconnected	Scan failed to prepare	Reconstruction Server not responding	Restart the scanner
14	Please verify the bore is empty and restart the Daily Calibration.	Daily Calibration could not be performed because the scanner detected something in the bore.	An object was in the bore.	Check bore, remove object, and restart Daily Calibration.

Error- code number	Fault description	Explanation	Cause ¹	Resolution
15	CANNOT_WRIT E_TRANSMIT_ QUEUE	When Data Acquisition System (DAS) views are acquired to disk during a scan, they are sent to a "Transmit Queue" from which they are sent to Recon computer over socket interface (Ethernet). If write of view to Transmit Queue fails, this error is flagged.	Likely causes of failing this write is if downstream data path is not functioning correctly (Ethernet unplugged, scanner app not connected, etc.) during a scan, and transmit queue is full and won't allow any more writes.	Contact Technical Support for service
16	INCORRECT_RO TATE_SPEED	Indicates disk detected a rotation speed error.	Happens when a problem with tick fence, or has incorrect calibration parameters.	Contact Technical Support for service
17	POSITION_ERR OR	Translate wheels did not move specified distance.	Could be caused by incorrect calibration parameters, a translate jam, an uneven floor	Contact Technical Support for service
18	VELOCITY_ERR OR	Translate velocity not as expected during a helical scan.	Could be caused by incorrect calibration parameters, a translate jam, an uneven floor	Contact Technical Support for service
19	OFFSET_CAL_F AULT	Offset calibration (done typically at beginning of every scan) has failed.	Due to bad view data from Disk Computer Assembly (DCA).	Contact Technical Support for service

Error- code number	Fault description	Explanation	Cause ¹	Resolution
20	AIR_CAL_FAULT	Air calibration failed.	An object was in the bore	Remove obstacle and perform another air calibration
21	INVALID_PROT OCOL	[Placeholder for invalid protocol error]	N/A	N/A
22	INVALID_COM MAND	[Placeholder for invalid command error]	N/A	N/A
23	INVALID_COM MAND_SEQUE NCE	An invalid command sequence was detected.	Can happen when Disk Computer Assembly (DCA) receives Start Acq command from Scanner Control app, but is not in a READY state.	Contact Technical Support for service
24	INVALID_PARA METERS	[Placeholder for invalid parameter error]	N/A	N/A
25	XRAY_COMMU NICATION_ERR OR	Disk has detected a problem with serial port connection to monoblock	HV Generator fault, Disk Control Assy (DCA) fault	Contact Technical Support for service
26	DCB_COMMUN ICATION_ERRO R	[Placeholder for DCB communication error]	N/A	N/A

Error- code number	Fault description	Explanation	Cause ¹	Resolution
27	SUBSYTEM_CO MMUNICATION _ERROR	[Placeholder for subsystem communication error]	N/A	N/A
28	RECON_ERROR	Requested protocol has been rejected by Reconstruction App.	Recon Computer error	Contact Technical Support for service
29	INSUFFICIENT_ TUBE_CAPACIT Y	When preparing for a scan, this error is flagged if tube capacity is lower than anticipated threshold value for that scan.	X-ray Tube too hot to perform prescribed procedure	Tube needs time to cool down before scan can be run.
30	INSUFFICIENT_ BATTERY_CAPA CITY	When preparing for a scan, this error is flagged if battery capacity is lower than anticipated threshold value for that scan.	Battery was not recharged per instructions	Battery needs to be recharged to minimum level by connecting it to AC outlet before scan can be run.
31	ESTOP_INTERL OCK	ESTOP button has been activated	User depressed ESTOP button	After respective issue has been resolved accordingly, user can deactivate E- STOP
32	HVG Cathode Error	N/A	N/A	Contact Technical Support for Service

Error- code number	Fault description	Explanation	Cause ¹	Resolution
33	SAFETY_INTERL OCK	[Placeholder for safety interlock error]	N/A	N/A
34	DAS_DATA_ERR OR	Disk Computer Assembly (DCA) has detected a Data Acquisition System (DAS) Data Error from DCB device.	Defective Converter Board Assembly	Contact Technical Support for service
35	DAS_CALIBRATI ON_ERROR	Tick calibration data is invalid	Disk Computer Assembly (DCA) has detected a Data Acquisition System (DAS) Data Error from Disk Control Board (DCB)	Contact Technical Support for service
36	HOME_TICK_ER ROR	Disk has detected a Home Tick Error from DCB device (no "home" pulse detected).	Typically this is a dirty/dusty Tick Fence Assembly or defective Tick Board Assy.	Contact Technical Support for service
37	TICK_ERROR	Disk has detected a Tick Error from DCB device (incorrect number of ticks counted).	Typically this is a dirty/dusty Tick Fence Assembly or defective Tick Board Assy.	Contact Technical Support for service
38	ROTATE_CONTR OLLER_ERROR	Rotate drive detected an error while performing homing operation	Base Computer Assy. Interface with rotate motor controller has a fault.	Contact Technical Support for service

Error- code number	Fault description	Explanation	Cause ¹	Resolution
39	AIR_CAL_WAR NING	Result of air calibration was out of tolerance	When requesting a new scan, currently loaded air calibration tables were generated from a previously failed air cal. Requested scan will be allowed to continue, but images may have artifacts. A pop-up window to user is presented upon this condition.	Contact Technical Support for service
40	RECON_BUSY	[Placeholder for recon busy error]	N/A	N/A
41	INSUFFICIENT_ DISK_SPACE	Not enough disk space to store information on the Recon Computer	Hard drive on Recon Computer does not have enough space for image data.	Contact Technical Support for service
42	Failed to process image. Exported image null? {true/false} imagePath null? {true/false}	N/A	N/A	Restart the scanner and try executing command again

Error- code number	Fault description	Explanation	Cause ¹	Resolution
43	Error in protocol validation. Please reboot the scanner and try again	N/A	N/A	Restart the scanner and try executing command again
44	One of the subsystems have disconnected. Please reboot the scanner	Recon Server not responding	N/A	Restart the scanner
45	Error retrieving original protocol for post recon	Recon data is missing or cannot be located	If it is an old series, the raw data may have already been deleted.	Contact Technical Support
46	Error executing protocol. Please register patient before executing protocol	N/A	Trying to perform a scan without a patient being registered due to possible lost Wi-Fi Connection after registration	Re-register a patient and try again or restart Tablet
47	HVG Latch Error message—HE Flow Switch Open (heat exchanger off)	N/A	N/A	Retry executing command first and then restart the scanner. If message does not clear, contact Technical Support

Error- code number	Fault description	Explanation	Cause ¹	Resolution
48	HVG Latch Error DAS Data Fault (Bad Detector/Modu le	N/A	N/A	Retry executing command first and then restart the scanner. If message does not clear, Contact Technical Support
49	HVG Latch Error ACQ Rotate Limit	N/A	N/A	Retry executing command first and then restart the scanner. If message does not clear, Contact Technical Support
50	HVG Filament Error	N/A	N/A	Retry executing command first and then restart the scanner. If message does not clear, Contact Technical Support
51	Protocol Rejected Battery	Not enough battery capacity needed to perform scan	Scanner batteries are low	Plug scanner in, ensure breaker is flipped on.
52	XBT_Error (Disk HW related)	X-rays were terminated due to a software anomaly	Scans that result in X-ray Below Threshold (XBT) can result in missing images due to system being allowed to complete scan	Contact Technical Support for service

Appendix C Emergency Service Kit Procedure

Introduction

Purpose

The Purpose for this document is to explain how to use the Emergency Jack Kit in case the OmniTom fails during transport (either from failing battery or other failure with the transport components).

Lifting the OmniTom

If the OmniTom is in TRANSPORT mode (scanner raised) proceed to Section 3.1 If the OmniTom is in SCAN mode (scanner lowered) proceed to Section 3.2

In Transport Mode

Jacking Up the OmniTom



Figure 281: Jack Components



Figure 282: Closeup of Release Valve

- 1) Place the jack near one end of the scanner.
- 2) Make sure the jack is in its lowest position by putting the jack bar over the release valve and twisting it counterclockwise. The lifting toe of the jack should be level with the stabilizing feet.
- 3) Put the spacer block (50-03556-001) on the jack lower tab.



Figure 283: Placement of space block

4) Make sure the release valve is then tightened by using the jack bar and twisting it in a clockwise direction.



Figure 284: Using the release valve

5) Slide the jack and spacer under the rugged bumper, making sure the spacer is still lined up with the jack.



- **CAUTION** Confirm the system is fitted with ruggedized bumpers before attempting to jack the system up by the bumpers. Systems with non-ruggedized bumpers will not support the weight of the scanner and will be damaged by doing so.
- 6) Insert the jack bar back into the connector.



Figure 285: Inserting the jack bar

7) Using the jack bar, slowly start jacking up the side of the scanner.



Figure 286: Proper placement of Jack

8) Once the Omniwheels on the side you are jacking up have lifted off the ground with enough room for the dollies, slide the dollies under the Omniwheels (one under each Omniwheel).



Figure 287: Move the dollies into place



CAUTION Lift only as high as required to slide the dollies under the Omniwheels.

Positioning the Dolly

1) Make sure the dolly is lined up tight with the Omniwheel.



CAUTION If not placed properly, the Omniwheel could slide off the dolly while trying to move the scanner.



Figure 288: Correct placement of Omniwheel on dolly

Lowering the OmniTom onto the dolly

- 1) Once the two dollies are in place, remove the jack bar
- 2) Put the jack bar over the release valve and very slowly start to twist the bar counterclockwise until the jack starts to lower



CAUTION This needs to be done slowly. If release too quickly the scanner will drop onto the dollies, possibly displacing them.

Repeat steps 3.1.1 through 3.1.3 for the other side of the scanner.

In Scan Mode

Jacking up the OmniTom



Figure 289: Jack Components

- 1) Place the jack near one end of the scanner.
- 2) Make sure the jack is in its lowest position by putting the jack bar over the release valve and twisting it counterclockwise. The lifting toe of the jack should be level with the stabilizing feet.



Figure 290: Using the release valve

- 3) Make sure the release valve is then tightened by using the jack bar and twisting it clockwise.
- 4) Slide the jack under the rugged bumper.



- **CAUTION** Confirm the system is fitted with ruggedized bumpers before attempting to jack the system up by the bumpers. Systems with non-ruggedized bumpers will not support the weight of the scanner and will be damaged by doing so.
- 5) Insert the jack bar back into the connector.



Figure 291: Inserting the jack bar

6) Using the jack bar, slowly start jacking up the side of the scanner.



Figure 292: Jacking up the Scanner



CAUTION Lift only as high as required to slide the dollies under the Omniwheels.

7) Once the Omniwheels on the side you are jacking up have lifted off the ground with enough room for the dollies, slide the dollies under the Omniwheels (one under each Omniwheel).

Positioning the Dollies

1) Make sure the dolly is lined up tight with the Omniwheel.



CAUTION If not placed properly, the Omniwheel could slide off the dolly while trying to move the scanner.



Figure 293: Correct placement of Omniwheel on dolly



Figure 294: Proper placement of dolly

Lowering the OmniTom onto the Dolly

- 1) Once the two dollies are in place, remove the jack bar.
- 2) Put the jack bar over the release valve and very slowly start to twist the bar counterclockwise until the jack starts to lower.



CAUTION This needs to be done slowly. If release too quickly the scanner will drop onto the dollies, possibly displacing them.

Repeat steps 3.1.1 through 3.1.3 for the other side of the scanner.

Moving the OmniTom

Once the scanner has been placed on the dollies it is ready to move out of the room.

The scanner weight is roughly 1,700lbs. It will need at least two people to move the scanner.

If the scanner was in Scan Mode, the clearance under the scanner is very small; approximately 5mm. Keep this in mind if you need to move the scanner over uneven surfaces such as a door threshold or into/out of an elevator.



CAUTION The Emergency Lift Kit is not designed to be used to move the scanner long distances nor be loaded on an elevator. The dolly wheels are small and could get stuck in the gap between the elevator floor and hallway floor.



CAUTION Do not use the drive bar to push or pull the scanner.

Appendix D Revision History

Rev	ECO #/Date	Author	Description
00	ECO-007358 2024/12/11	Keith A. Kaser	Initial Release (Based off OmniTom Elite manual (1-NL5000-060 Rev17)
01	ECO-007423 2025/01/27	Keith A. Kaser	Updated cover photo. Updated screen shots. Updated Intended Use of the System. Updated International Distributer contact information. Updated Part numbers and pictures. Updated Product Marking Plate. Updated Product Description under System Overview. Added note to wait 30 seconds before powering system on after a power-off. Updated Physics sections in Chapter 7. Added information about copying and reusing a protocol. Removed reference to Manual Patient registration since it was replaced by Manual Add option. Updated Daily Calibration and QA section. Added Ability to Re-use Scouts if changing protocol after scout. Added Noise Reduction for Axial scans. Added ability to type radius of the ROI. Added Using Bed Adapter Safety Straps section to Chapter 13. Added Appendix C Emergency Service Kit Procedure.

Appendix E Varex Imaging Data





Stationary Anode X-Ray Tube



Product Description

The MCS-640 is a 140 kV, air cooled stationary anode metal ceramic x-ray source. This source is specifically designed for Imaging Applications.

X-Ray Tube Specifications

Maximum Peak Voltage	140 kV
Anode to Ground Cathode to Ground	
Focal Spot - IEC 60336 Small	1.0 W x 1.0 L
Cooling Medium	Water/Glycol
Maximum Continuous Rating Small	400 W with 0.6 gpm coolant flow
Target Material	Tungsten-Rhenium
Target Angle	5°
Radiation Coverage	
X-Ray Tube Assembly Permanent Filtration .	2.0 mm Be

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Dimensions are for reference only

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SECTION B-B

NOTE: 1. FULL BEAM COVERAGE

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MCS-640 Filament Characteristics

WARNING

Beryllium windows transmit a very high level of long wavelength X-radiation, which can injure human tissue. Injury may occur from even very short exposures to the primary X-ray beam. Follow all precautions necessary to avoid radiation exposure to humans.

The radiation dose rate cannot be accurately measured with conventional radiation measurement instruments. Radiation intensity in each installation will vary, and calibration must include the effects of long wavelength X-radiation.

Fumes from beryilium metal (or its compounds) as well as dust can be hazardous if inhaled. During use, corrosion products may occur on the beryilium window, but these should not be scraped off, machined, or otherwise removed. Tube unit disposal should conform to federal, state, and local regulations governing beryilium.



Manufactured by Varex Imaging Corporation

Specifications subject to change without notice.

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