# NL4000 BodyTom<sup>®</sup> Elite User Manual

1-NL4000-060 Revision 19



Introduction Page 1 of 427

# **NeuroLogica Corporation**

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## Intended use of the system

The NL4000 BodyTom Elite CT system is intended to be used for x-ray computed tomography applications for anatomy that can be imaged in the 85cm aperture. The CT system is intended to be used for both pediatric and adult imaging and as such has preset dose settings based upon weight and age.

The CT images can be obtained either with or without contrast. BodyTom Elite CT systems can be used for low dose lung cancer screening. The screening must be performed in compliance with the approved and established protocols as defined by professional medical societies.

\* Please refer to clinical literature, including the results of the National Lung Screening Trial (N Engl J Med 2011; 365:395-409) and subsequent literature for further information.

#### 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

## **Proprietary rights**

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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 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.

Any information or descriptions contained in this manual may not be reproduced and released to any of the general public or used in conjunction with other professional instruction without written consent of NeuroLogica Corp., USA – a subsidiary of Samsung. Direct any written inquiries to the appropriate address found in the section "Contact information" on page 25.

Unauthorized copying of this user manual may not only infringe copyright but also reduce the ability of NeuroLogica Corp. to provide accurate and up-to-date information to users: limited and restricted operators and administrators.

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, write to and include chapter title and page number:

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

If you have any questions about faults or errors on the system, battery or charge issues, or mechanical issues with the scanner, contact a **Technical Representative**.

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If you have questions about the clinical use of your system, building protocols, creating MPRs, imaging artifacts, creating a clinical workflow or process, logging in or access issues, and general usage of the system, contact **Customer Service**.

EC REP		
Winckels Medical Devices Expertise	Europe Bergerweg 18 6085 AT Horn The	Tel: +31 (0)475 582285 Fax: +31 (0)475 582278
Devices Expertise	Netherlands	Tax. 131 (0)473 382278
<b>Australian Sponsor</b>	Level 8/15 Talavera	Mobile: +61 (0)412 563 016
	Road	Tel: +61 (0)2 8114 1535
	PO Box 646	Fax: +61 (0)2 8114 1599
	North Ryde NSW 2113	Customer Service:
	Australia	1-800 060 168
<b>Brazilian Authorized</b>	2 <sup>nd</sup> floor, 515 Rua	Tel: 55-11-3371-1500
Distributor	Peixoto Gomide	
	Sao Paulo	
	Brazil	

## 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 facility 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 content 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 radiologic technologists, physicians, radiologists, and other medical specialists.

Users should be trained professionals who are certified to operate such systems *before* scanning or diagnosing patients. This training must include medical and x-ray education, as well as NeuroLogica applications training.

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Everyone that 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 NL4000 BodyTom Elite has the following essential performance factors mitigated by design:

- Over Radiation protection
- Re-scan prevention
- Stray Radiation exposure prevention
- Diagnostic performance

#### About this user manual

The instructions in this user manual describe how to use the NeuroLogica NL4000 BodyTom Elite Computed Tomography (CT) system, manufactured by NeuroLogica Corp.

The manual applies to both old and new colors of BodyTom Elite.

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, as well as 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 from 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.

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## 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.

**CAUTION** Indicates a hazardous situation, which if not avoided *could* 

result in minor or moderate injury.

## Conventions used in this user manual

Table 1: Conventions used in this user manual

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 <b>Customize</b> and then click <b>System</b> .
Bold	When content refers to commands, windows, screens, dialog boxes, popups, tabs, buttons, options, keyboard keys, statuses, and modes, these items appear in <b>bold</b> 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 the action <i>in</i> 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.
Note	The appearance of a note is as such:

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Convention	Use
	Note Indicates additional information to help you operate this product.
Hyperlink (an electronic cross- reference)	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.  A hyperlink is a quick way to go to another area of the user manual with a simple click. Hyperlinks appear like this: "Understanding the types of users" on page 101. In this case, hover the mouse pointer over the gray hyperlink text. Hold the Ctrl key on your keypad and click the mouse button. After you click the hyperlink, the hyperlink takes you to the referenced area in the user manual.
Click vs right-click	In this user manual, click means to press the left mouse button. This user manual never says 'left click' as it is assumed that is the traditional way to click; however, it does point out when to right click the mouse button.

# Understanding the use of "you" in this user manual

Unless specifically noted, the implied "you", in this user manual, is the user/operator. It is assumed users/operators are certified and medically trained personnel, qualified to use these systems.

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

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

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## **Active and inactive objects**

When a menu command, option, button, tab, field, is gray, the item is not active or enabled. When an item is gray, it can mean additional or required tasks must be completed first or you do not have permission to access that option. An active menu command, option, button, tab, and field means you can use the item to perform an action. Active items are green and/or highlighted.

## Applicable versions of BodyTom Elite

This user manual is applicable to all versions of BodyTom Elite, despite the color variations.



Figure 1: Current cover (left) and former covers (middle & right)

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# **Chapter 1** Compliance and Safety Requirements

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

# IEC classification and symbols



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

Type B equipment provides an adequate degree of protection against shock, 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 E

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.

BodyTom Elite Computer Tomography systems comply with Class I- Type B equipment as defined in IEC 60601-1 standard.

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

The BodyTom Elite CT scanner is patient-environment equipment.

Table 2: Applicable symbols

Symbol Symbols	Description	
$\sim$	Alternating current	
	Protective earth (ground)	
\display \d	Functional Earth	
<u> </u>	Caution: consult accompanying documents	
4	Caution: risk of electrical shock	
	Electrostatic sensitive devices	
<b>†</b>	Type B equipment	
A	X-ray warning	
	X-ray source assembly emitting	
	Non-ionizing radiation	
	Warning: laser in use	
LASER RADIATION DO NOT STARE INTO BEAM CLASS 2 LASER PRODUCT	Warning: Laser Radiation Do Not Stare into Beam Class 2 Laser Product	
Max Power Output: 1mW Wavelength: 650m Complies with IEC 60825-1.2014, 3rd ed (2014-05) 50:03815-001ex00	Laser Output and Standards Information Label	
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	
	Emergency switch	
	Crush warning	

Symbol	Description
	Foot/toe crush warning when lowering machine
<b>†</b>	System up
<b>+</b>	System down
-25 ·c (50 ·c) (40 ·c)	Temperature limits
$\frown$	Keep away from rain for packaging
XXX er	Humidity limit for packaging
	Warning: battery charging
-	Fuse usage
i	Refer to instruction in user manual/booklet
MD	Medical Device Symbol
	Legal Manufacturer Symbol
<b>C</b> € <sub>0413</sub>	CE Mark or Conformité Européenne ; number below CE represent Notified Body number
c Classified us Intertek	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: 24hours a day, seven days a week.

Table 3: Operating environment

Operating	
Ambient temperature	15º C to 35º C (59ºF to 95º F)
Relative humidity	20% to 80% (non-condensing)
Altitude	0-3048m (0-10,000 ft.)
Storage	
Temperature	-25º C to 70º C (-13º F to 158º F)
Relative humidity	20% to 85% (non-condensing)
Transport	
Temperature	-20º C to 60º C (-4º F to 140º F)
Relative humidity	20% to 85% (non-condensing)
Powering system	
Time period prior to powering the system	24 hours <sup>1</sup>
Floor	
Flatness	<+/1 0.120in. (3mm) per 10ft. (3.048m)
Recommended minimum scan area	10ft. x 15ft. (3.048m x 4.572m)

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

Table 4: System operating parameters and specifications

Table Heyerem eperating par		
	120V~	240V~
Phase	Single	Single
Voltage range	100-120V~±10%	208-240V~ ±10%
	Nominal voltage for U.S. 120, 240	
Circuit protection	20 amps	30 amps
Facility outlet	NEMA 5-20R	NEMA 6-30R
	Outline of outlet:	

<sup>&</sup>lt;sup>1</sup> 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.

Phase	120V~ Single	240V~ Single
	W <sub>G</sub> O	
Frequency	50 or 60Hz	50 or 60Hz
Battery capacity	Fully charged, 12 hours typical	
Typical usage	110-120V~ 60Hz	230-240V~ 50Hz
Wiring	125V, 2 pole, 3 wire grounding	250V, 2 pole, 3 wire grounding
Main power supply's apparent resistance	0.105Ω	
Heat dissipation (when system is not idle)	55kW	

## 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 the red **EMERGENCY STOP** button immediately in case

of abnormal or unexpected motion.



**WARNING** Verify scanner is on its centipedes (fully down position) prior

to positioning patient at scanner entrance.



**WARNING** Make sure all extremities are not under scanner while

lowering or raising it.



**WARNING** In the case of a single pendant failure, the additional pendant

is available for use to prevent loss of system function.



**WARNING** Always keep patient in view. Ensure that the patient can be

seen when the operator is near the scanner control panel and **EMERGENCY STOP** button. Never leave 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 correct location and consider wearing personal radiation protective equipment.
- A radiation safety plan in the working area boundaries, to optionally 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, radiation report telephone number.

## Site specification

**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~
Operating frequency	50Hz-60Hz
Apparent resistance of supply mains at 120VAC	0.3 ohms
Operating current at 120VAC	13 amps
Heat dissipation	1672 watts



**CAUTION** For domestic purposes, scanner can be powered using either 120V~ or 240V~. If the scanner is using 120V~, the facility's circuit must be capable of providing 20 amps (single phase). If the scanner is using 240V~, the facility's circuit must be capable of providing 30 amps (single phase). If other devices are connected to the same circuit, the facility's circuit breaker may trip and, therefore, prevent the scanner from being ready when needed.

Table 6: Battery operating parameters

Operating voltage	480 to 585 VDC
Output current (peak)	100 amps

## Site specification for enclosed CT room

Table 7: Site specification

Issue	Comment
Receiving area	Secured
Dacking material and waste	Near availability of a trash receptacle for
Packing material and waste	dunnage
Room dimensions for use	12ft. x 15ft. room with a finished level floor;
Rooff differsions for use	recommended the room be well lit
Power availability	120VAC/20amp wall outlets (2x)
Floor flatness	<± 0.120in. (3mm) per 10ft.
Floor strength	Site must be able to support product weight

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

- NeuroLogica advises complying with local regulations and/or site recommendations as specified by the facility physicist or certified representative for the following:
- There should be enough space inside the CT room. The area should not be less than 30m<sup>2</sup>. Any side should not be less than 4m. Leaving any unnecessary items inside the room is prohibited.
- Carpet and soft material cannot be used on the floor. There should not be obstacles on the floor. Ensure flat surface area no less than 12ft. x 15ft. level degree <±3mm per 10ft.
- Appropriate protection measures should be taken to protect staff and to ensure annual-dose-rate is less than 0.25mSv (equals to 5 uSv weekly).

### Hazardous substances

Table 8: Hazardous substances

Substance/material	≅ Weight/system
Lead	7.69 kg (17.0lbs.)
Cadmium	0.036kg (0.079lbs.)
Mercury	Okg (Olbs.)
Hexavalent chromium	Okg (Olbs.)
PolyBrominated Biphenyls (PBB)	<0.46kg (1lb.)
PolyBrominated Diphenyl Ethers (PBDE)	<0.46kg (1lb.)

## Part numbers and product-marking plates

Table 9: Core-system-component part numbers and product-marking plate locations

Component	Part number	Product-marking plate locations
	0-NL4000-001	Near the main input plug or on
BodyTom Elite gantry	0-NL4000-002	the side of the system. See
	10-00345-0001	Figure 2 below.
BodyTom Elite	40-00157-000	On the back of the
workstation		workstation.
QA phantom	10-00268-001	On the back of the phantom.

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

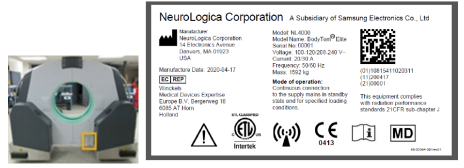


Figure 2: Product-marking plate on scanner

Table 10: Core-system component dimensions

Component / mode		Size L x W x H	
BodyTom Elite	Scan	256.54cm x 103.1cm x 199.6cm	
	Scan	101in. x 40.58in. x 78.57in.	
	Tuenenent	256.54cm x 103.1cm x 205.7cm	
	Transport	101in. x 40.58in. x 81.00in.	
	Bore	85cm	
		33in.	



Figure 3: Scanner dimensions including drivebar

Table 11: Workstation dimensions

Component	Size (inches)	Size (centimeters)	Weight	Weight
	L x W x H	L x W x H	(lbs)	(kg)
BodyTom Elite system Workstation Cart	26.3 x 24.4 x 79.8	66.8 x 62.0 x 202.7	207	94

## 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 nonmedical 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, and so on. 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 is required to instruct technologists, patients, and other people who may be around this equipment to fully comply with the above regulation.

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

### Focal spot

Nominal size is: ~1.2 x 1.4mm

Size limit is: ~1.2 to 1.7mm width and ~1.4 to 1.9mm length.

Testing standard for reference is as follows:

IEC 60336:2005

Focal spot centering is within 1mm of center of bolt pattern. Maximum motion due to gravity in X, Y, and Z axis is 0.1mm.

Maximum motion from anode rotation is 0.1mm.

Maximum motion from anode heating in X axis is 0.1mm. Maximum motion from anode heating in Z axis is 0.3mm.

## Anode input power

The maximum, anode, cooling rate is 8,750W (12,250 HU/sec).

The maximum, anode, heat dissipation is 3,400W (4,760 HU/sec).

The nominal, anode, input power is 42kW.

Continuous anode input power when applied at the nominal, x-ray, tubevoltage is 150kV, 23mA.

### **Filtration**

Table 12: Filtration

Tube Voltage (kV)	100	120	140
Half-value layer (aluminum equivalent)	6mm	7mm	8mm
Filters consist of 0.0014in. [0.036mm] of copper and 0.086in. [2.18mm] of aluminum, along with a variable thickness bowtie filter made from Teflon®.			

X-ray tube's total filtration of irremovable layers is 5.7mm of equivalent aluminum.

## Source to Detector distance (SID)

The SID value is 1041.9mm.

## **Compliance statement**

Note All editions and years of revisions for standards noted in this chapter are static as of Revision 15 of this user manual (and Revision 02 of the BodyTom Workstation User Manual), which has been incorporated into this user manual to include the scanner and the workstation.

The BodyTom Elite system complies with the regulatory requirements of the following:

- AAMI ES60601-1 Issue: 2005 Version Medical Electrical Equipment, Part 1: General Requirements for Basic Safety and Essential Performance; Amendment No. 2: 2010/05/30.
- CAN/CSA-C22.2 No. 601.1-M90(R2005) Issued: 2003/11/01 Medical Electrical Equipment – Part 1: General Requirements for Safety; General Instruction No. 1: 1990, Supplement 1: 1994, Amendment 2: 1998, General Instruction No. 2: 2003.
- CENELEC EN 60601-1 2nd Edition, Medical Electrical Equipment Part 1: General Requirements for Safety, includes Amendment A1:1993 and A2:1995.
- CENELEC EN 60601-1 3rd Edition, Medical Electrical Equipment Part 1: General Requirements for Safety.
- CSA C22.2#60601-1 Issued: 2008/02/01 Ed 3 Medical Electrical Equipment Part 1: General Requirements for Basic Safety and Essential Performance.
- IEC 60601-1 Issued: 2005/01/01 Ed 3 Medical Electrical Equipment Part 1: General Requirements for Basic Safety and Essential Performance.

- IEC 60601-1-1 (2000) 2nd Edition: Medical Electrical Equipment, Part 1: General Requirements for Safety Collateral Standard: Safety Requirements for Medical Electrical Systems.
- IEC 60601-1-2:2014 Ed4.0 Medical Electrical Equipment Part 1-2: General Requirements for Basic Safety and Essential Performance Collateral Standard: Electromagnetic Disturbances
- IEC 60601-1-3 (2008), Medical Electrical Equipment Section 1-3: General Requirements for Safety. Collateral Standard: General Requirements for Radiation Protection in diagnostic X-ray Equipment.
- IEC 60601-1-4 (2005), 3rd Edition Consolidated Edition, Medical Electrical Equipment Part 1-4: General Requirements for Safety – Collateral Standard: Programmable Electrical Medical Systems.
- IEC 60601-1-6 Issued: 2008/12/08 Ed 2 Medical Electrical Equipment Part 1-6: General Requirements for Safety. Collateral standard: Usability.
- IEC 60601-2-28 (1993) Medical Electrical Equipment, Part 2: Particular Requirements for the Safety of X-ray Source Assemblies and X-ray Tube Assemblies for Medical Diagnosis.
- IEC 60601-2-32 (1994) Part 2-32, Particular Requirements for Safety sections 2.32 Specification for Associated Equipment of X-ray Equipment.
- IEC 60601-2-44 (2009) Medical Electrical Equipment -Part 2-44: Particular Requirements for the Safety of X-ray Equipment for Computed Tomography.
- IEC 60825-1:2007 Safety of Laser Products Part 1: Equipment Classification, and Requirements 2nd Ed.
- International Electrotechnical Commission (IEC) International Standards Organization, when applicable.
- Intertek Testing Service (ITS), an independent testing laboratory.
- Medical Device Good Manufacturing Practice Manual issued by the FDA (Food and Drug Administration). Department of Health, USA.
- 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 the unintentional generation, propagation, and reception of electromagnetic energy with reference to the unwanted effects (Electromagnetic Interference (EMI)) 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.

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 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, or noise, mitigation, and hence EMC 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.

## **Electromagnetic Interference (EMI)**

**Electromagnetic Interference (EMI)**, also called **Radio Frequency Interference (RFI)** is 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 BodyTom Elite 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 (see Table 14 on page 78).
- Immunity Compliance level and recommendations to maintain equipment clinical utility.

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.

Table 13: Acronyms and abbreviations

Acronym and	Definition		
abbreviation	Bernitton		
AEC	Automatic Exposure Control		
CBF	Cerebral Blood Flow		
CBV	Cerebral Blood Volume		
СТ	Computed Tomography		
СТА	CT Angiography		
СТР	CT Perfusion		
CTDI <sub>vol</sub>	Volume Computed Tomography Dose Index		
CTDI <sub>w</sub>	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		
EMI	Electromagnetic Interference		
FOV	Field Of View		
HIS	Hospital Information System		
HU	Hounsfield Unit		
IBC	Iterative Bone Correction		
MAR	Metal Artifact Reduction		
MIP	Maximum Intensity Projection		
MPPS	Modality Performed Procedure Step		
MPR	Multi-Planar Reformation, sometimes referred to as		
IVIFIX	Multi-Planar Reconstruction		
MTT	Mean Transit Time		
PACS	Picture, Archiving, and Communication System		
QA	Quality Assurance		
RIS	Radiology Information System		
RSO	Radiation Safety Officer		
RFI	Radio Frequency Interference		
SCP	Service Class Provider		
SCU	Service Class User		



**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 14: Emission declaration for BodyTom Elite systems

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

Emissions test	Compliance	Electromagnetic environment guide
RF emissions CISPR 11	Group 1	BodyTom Elite systems use RF energy only for internal function. Therefore, RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.
RF emissions CISPR 11	Class A	BodyTom Elite systems are predominantly intended for use in non-domestic
Harmonic emissions, IEC 61000-3-2	Class A	environments, and not directly connected to the Public Mains Network. BodyTom Elite systems are predominantly intended
Voltage fluctuations/flicker emissions, IEC 61000-3-3	Complies	for use (for example, in hospitals) with an appropriate power supply (see operation manual) and recommended shielding for portable use.

Table 15: EMC Immunity declaration for BodyTom Elite systems

BodyTom Elite systems are intended for use in the electromagnetic environment specified below. The customer or user of an BodyTom Elite system should assure that it is used in such an environment.

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Immunity test	IEC 60601-1- 2 test level	Compliance level	Electromagnetic environment guidance
Electrostatic	±8kV contact	±8kV contact	Floors should be wood, concrete, or ceramic tile. If floors are
discharge (ESD) IEC 61000-4-2	±2kV, ±4kV, ±8kV, 15kV air	±2kV, ±4kV, ±8kV, 15kV air	covered with synthetic material, relative humidity should be at least 30%.
Electrical fast transient/burst IEC 61000-4-4	±2kV for power supply lines ±1kV for input/output lines	±2kV 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	±1kV line-line ±2kV line- ground	±1kV line-line ± 2kV line- ground	Mains power quality should be that of a typical commercial or hospital environment.
Voltage dips, short interruptions,	0% UT; 0.5 cycle At 0°, 45°, 90°, 135°, 180°, 225°, 270° and 315°	0% UT; 0.5 cycle At 0°, 45°, 90°, 135°, 180°, 225°, 270° and 315°	Mains power quality should be that of a typical commercial or hospital environment. If the user
and voltage variations on	40% UT; 5 cycles	40% UT; 5 cycles	of a BodyTom Elite system requires continued operation during power interruptions, it is
power supply input lines IEC 61000-4-11	70% UT; 25 cycles	70% UT; 25 cycles	recommended that the BodyTom Elite system be powered from its internal batteries.
	0% UT; 250 cycles	0% UT; 250 cycles	
Immunity test	IEC 60601-1-2 Test Level	Compliance level	Electromagnetic environment guidance.
Power frequency (50/ 60Hz) magnetic field IEC 61000- 4-8	30 A/m, 50Hz or 60Hz	30 A/m, 50Hz or 60Hz	Power-frequency magnetic-fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.
Clause 8.10	Table 14	Per Table 14	IEC 60601-1-2:2014

BodyTom Elite systems are intended for use in the electromagnetic environment specified below. The customer or user of an BodyTom Elite system should assure that it is used in such an environment.

used in such an er			
Immunity test	IEC 60601-1- 2 test level	Compliance level	Electromagnetic environment guidance
Conducted RF IEC 61000-4-6	3 Vrms 150kHz to 80MHz	V1 = 3 Vrms	Portable and mobile RF communications equipment should be used no closer to any part of an BodyTom Elite system, including cables, than recommended separation distance calculated from the equation appropriate for transmitter frequency.  Recommended separation distance: See Table 16.
Radiated RF IEC 61000- 4-3 (alternative method: IEC 61000-4-21)	3 Vrms 80MHz to 2,5GHz	E1 = 3 V/m	$d = [\frac{3.5}{V_1}]\sqrt{P}$ $d = [\frac{3.5}{E_1}]\sqrt{P}$ $80 \text{ MHz to } 800 \text{ MHz}$ $d = [\frac{7}{E_1}]\sqrt{P}$ $800 \text{ MHz to } 2.5 \text{ GHz}$ Where P is maximum power rating in watts and D is recommended separation distance in meters. Field strengths from fixed transmitters, as determined by an electromagnetic site survey, should be less than compliance levels (V1 and E1). Interference may occur in vicinity of equipment marked with the following symbol: $((\mathbf{Q}))$

Note: The wireless receiver operates within the following bands.

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2.412 to 2.462 GHz (11 channels)
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- 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 very difficult to grapple with issues related to EMC. It may take a variable amount of time and cost to identify issues causing interference.

General countermeasures of electromagnetic interference with other equipment:

- Electromagnetic interference may be alleviated by positioning other equipment far from system.
- Electromagnetic interference may be mitigated by changing relative location (installation angle) between system and other equipment.
- Electromagnetic interference may be eased by changing wiring locations of power/signal cables of other equipment.
- Electromagnetic influence may be reduced by altering the power-supply path of other equipment.
- Electromagnetic environment specified (see Table 15 on page 48).

Table 16: Recommended separation distances

Recommended separation distances between portable and mobile RF communications equipment and the BodyTom Elite system

BodyTom Elite system is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The user of the BodyTom Elite system can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the BodyTom Elite system as recommended below, according to the maximum output power of the communications equipment.

Rated maximum output Power	150kHz to 80MHz	80MHz to 800MHz	800MHz to 2,5GHz
(P) if transmitter Watts (W)	Separation distance meters <sup>2</sup>	Separation distance meters <sup>2</sup>	Separation distance meters <sup>2</sup>
0.01	.12	.12	.23
0.1	.38	.38	.73
1	1.2	1.2	2.3
10	3.8	3.8	7.3
100	12	12	23

<sup>&</sup>lt;sup>2</sup> Separation distance according to frequency of transmitter (m)

Recommended separation distances between portable and mobile RF communications equipment and the BodyTom Elite system

BodyTom Elite system is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The user of the BodyTom Elite system can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the BodyTom Elite system as recommended below, according to the maximum output power of the communications equipment.

Rated	150kHz to	80MHz to	800MHz to
maximum	80MHz	800MHz	2,5GHz
output Power (P) if transmitter Watts (W)	Separation	Separation	Separation
	distance	distance	distance
	meters <sup>2</sup>	meters <sup>2</sup>	meters <sup>2</sup>

For transmitters rated at a maximum output power not listed above, the separation distance is estimated using the equation in the corresponding column, where P is the maximum output power rating of the transmitters in Watts (W) according to the transmitter manufacturer.

**Note** At 80MHz and 800MHz, separation distance for higher frequency range applies.

**Note** These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects, and people.

Table 17: Test specifications for ENCLOSURE PORT IMMUNITY to RF wireless communications equipment

Test Frequency (MHz)	Band <sup>a)</sup>	Service <sup>a)</sup>	Modulation <sup>b)</sup>	Max Power (W)	Distance (m)	Immunity Test Level (V/m)
385	380- 390	Tetra 400	Pulse Modulation <sup>b)</sup> 18Hz	1.8	0.3	27
450	430- 470	GMRS 460,	FM <sup>c)</sup>	2	0.3	9

Test Frequency (MHz)	Band <sup>a)</sup>	Service <sup>a)</sup>	Modulation <sup>b)</sup>	Max Power (W)	Distance (m)	Immunity Test Level (V/m)
		FRS 460	±5kHz deviation 1 kHz sine			
710 745 780	704- 787	LTE Band 13,17	Pulse Modulation <sup>b)</sup> 217Hz	0.2	0.3	9
810 870 930	800- 960	GSM 800/900 TETRA 800, iDEN 820, CMDA 850, LTE Band 5	Pulse Modulation <sup>b)</sup> 18Hz	2	0.3	28
1720 1845 1970	1700- 1990	GSM 1800; CMDA 1900; GSM 1900; DECT; LTE Band 1, 3, 4, 25; UMTS	Pulse Modulation <sup>b)</sup> 217Hz	2	0.3	28
2450	2400- 2570	Bluetooth, WLAN,	Pulse Modulation <sup>b)</sup> 217Hz	2	0.3	28

Test Frequency (MHz)	Band <sup>a)</sup>	Service <sup>a)</sup>	Modulation <sup>b)</sup>	Max Power (W)	Distance (m)	Immunity Test Level (V/m)		
		802.11 b/g/n,						
		RFID 2450,						
		LTE Band 7						
5240		WLAN 802.11 a/n	Pulse					
5500	5100- 5800		802.11 Modulation b)	802.11	Modulation b)	0.2	0.3	9
5785								

### 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 BodyTom Elite 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 16 on page 83) between 150kHz and 2.5GHz, will reduce disturbances recorded at the image level, but may not eliminate all disturbances; however, 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) is put 2.3 meters apart from the BodyTom Elite system (to avoid image interference risks).

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

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

### Installation recommendations

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

BodyTom Elite system should not be used adjacent to or stacked with other equipment; if adjacent or stacked use is necessary, the BodyTom Elite 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

To avoid interference on the BodyTom Elite system, static-field limits from the surrounding environment are specified. The static field is specified less than <1 Gauss around the unit.

# Electrostatic discharge environment and recommendations

- 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 groundreference.

### **Facility IT-NETWORK**

The BodyTom Elite system utilizes the IT-NETWORK for the customer (as applicable) to communicate with the modality worklist and for supplemental, image-storage space. As part of the installation, the applicable IT-NETWORK is reviewed to create the appropriate setup to 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, 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

Review this material before using the system and observe basic, commonsense safety rules when operating this scanner.

### General safety considerations and statements

Review the following before using the system (Scanner and Workstation (as applicable)) to 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 or workstation if it appears damaged or fails.
- Wait for qualified personnel to correct any problem.

**Note** The scanner is provided with a video-camera monitoringsystem to help navigate the unit while being transported within a facility.

<u>^</u>	WARNING	Modification of this equipment is <i>not</i> allowed.
<u>^</u>	WARNING	Equipment maintenance of non-medical electrical equipment should not be performed in the patient environment.
<u>^</u>	CAUTION	All non-medical electrical equipment will comply with relevant IEC and ISO safety standards.
<u>^</u>	CAUTION	Federal law restricts the use of this device without a prescription by a physician.
<u>^</u>	CAUTION	Always store and/or use unit in a well-ventilated area. Keep air pollution to a minimum. Always keep floor clean.
<u>^</u>	CAUTION	Do not touch parts of non-medical electrical equipment in patient environment and patient simultaneously.
<u> </u>	CAUTION	For disposal of any material emanating from the system; follow local regulations.
<u> </u>	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.
<u> </u>	CAUTION	It is the user's responsibility to make sure that after installation or subsequent modification, the system complies with the requirements of collateral standard IEC 60601-1.
1	WARNING	Installation of this product is performed in accordance with Installation Manual (1-NL4000-059). All installation processes and qualified personnel are outlined in that document.
$\triangle$	WARNING	Proper disposal of batteries is required to ensure compliance with environmental safety guidelines. Contact authorized NeuroLogica representative for instructions.
<u> </u>	WARNING	Observe safety-exposure factors and operating procedures to protect patient from physical harm during contact with this x-ray scanner.
<u> </u>	WARNING	Observe safety requirements to prevent excessive dose exposure to patient and/or operator.
$\triangle$	CAUTION	Improper system (including workstation) usage could endanger patients and/or users and void the warranty if not operated correctly.
$\hat{\Lambda}$	CAUTION	Should the workstation encounter a computer related virus, be sure to contact Technical Support for assistance with removing said virus from the equipment.



**CAUTION** Radiation dose exposure to patients should not exceed

maximum of 1Gy CTDI.

<u>^</u>

**CAUTION** For proper disposal of material at equipment's end-of-useful

life; contact NeuroLogica for instructions.

WARNING Equipment in which protection against electric shock relies on

basic insulation only, should not be used in this system.

WARNING If the system fails to move due to loss of power, the patient

can be easily removed from the scanner by moving the

patient bed.

Four, foot crush hazard labels are affixed to the scanner in four places, above the four soft bumpers. The following shows a safety label:



Figure 4: Identifying the scanner's safety label(s) – foot-crush-hazard label(s)

# Laser safety

There are four lasers used with the BodyTom Elite system as indicated in Figure 5 on page 59: 1 laser (Coronal) at position 1, 1 laser (Sagittal) at position 2, and 1 external and 1 internal laser (Transverse or Axial) at position 3. Laser 2 is mounted internally (fixed) to the disk assembly, which spins within the system's bore. Therefore, the lasers output light is always aimed at and rotating within the bore itself.



WARNING Viewing the laser output with certain optical instruments (for

example, eye loupes, magnifiers, and microscopes) within

100mm may pose an eye hazard.



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.

This statement will be a label placed onto the system by the Driver Side.

### **Laser parameters**

- Lasers 1 and 3 (see Figure 5):
  - Wavelength = 650nm
  - Output Power = 1mW
- Laser 2 (see Figure 5):
  - Wavelength = 650nm
  - Output Power = 4mW

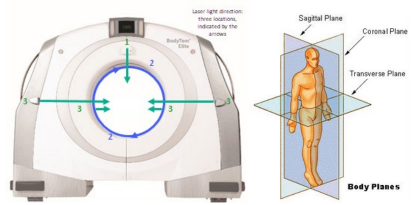


Figure 5: Laser aperture's direction



**CAUTION** 

Instruct the patient to close his/her eyes before 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. *Do not* 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 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 workstation. 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.



### CAUTION

To prevent healthcare provider injuries, a single healthcare professional should not move the scanner and workstation. Although one person can drive the BodyTom Elite when moving the scanner about the facility, NeuroLogica recommends two people move the scanner (lengthwise, only) to ensure no collisions occur when maneuvering through tight hallways and around corners. Be especially cautious when moving the system about an inclined floor.



### WARNING

To prevent involuntary movement, do not position scanner on an incline while in **Transport** mode.



**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** Contact **Technical Support** for assistance when movement is required on an incline.

**Note** Be sure there are no obstacles in front of the scanner while you move the scanner.

> If the system needs to be moved over a threshold it is critical that the scanner be oriented so that it is driven in the forward or reverse direction. The scanner does not have the capability of moving laterally over thresholds.



**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 by using control panel switches or pendant controls while observing patient.



**CAUTION** The scanner is equipped with a video camera to help the operator prevent collisions when transporting system to different locations that could otherwise result in personal injury or facility damage.



**CAUTION** Do not station or operate the system on an uneven floor. The flatness requirement is 0.12in. over 10ft. or 3mm over 305cm.



**CAUTION** Prior to transporting the scanner, verify that power cord is unplugged from wall to avoid damage to cord and outlet, and avoid tripping. Verify that the ethernet cable is unplugged from the workstation to avoid damage to the cable and connector.

### Floor level

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.



**WARNING** To separate the device from AC power simply disconnect the power cord from the wall and turn off the main breakers, located on the same side of the scanner as the power cord.



**WARNING** Access to the main breaker is critical for safety. Do not position the scanner so that the access to the breaker is diminished.



**CAUTION** 

Check to ensure the AC outlet is working properly before plugging in the system's AC power cord. NeuroLogica recommends using a dedicated outlet for powering the BodyTom Elite system, only.



WARNING

To prevent electrical shock, do not connect items that are not specified as part of the system, including the workstation.



**WARNING** To prevent electrical shock, do not remove the covers from the equipment. The covers protect the user and the patient from moving parts or 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 apply serviceable 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 to prevent short-circuiting or possible electrical shock.



WARNING

Never position the mobile system and/or workstation 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, threeconductor 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 qualified personnel 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 (for example, the workstation, AC power cord, and so on) 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 an 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, and so on, 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** All systems within the patient environment will provide the

same level of safety as medical equipment complying with IEC

60601-1.



**CAUTION** To help prevent tripping hazards, use care in the arranging of

any cords (for example, AC cord, ethernet cable, and so on)

when connecting to the system/workstation.



**CAUTION** To prevent damaging electrical outlet cords, check to ensure

they have been removed and properly stored before

transporting the scanner.



**WARNING** The BodyTom Elite 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.



**CAUTION** Use the **Universal Serial Bus** (**USB**) terminal located near the

**EMERGENCY-STOP** (E-STOP) button for archiving to USB, *only*. **Do not** use the USB terminal located near the E-STOP for

connecting any other device to equipment.



**CAUTION** For proper disposal of material at the end of the useful life of

the equipment, contact NeuroLogica for instructions.

# Mechanical safety



**WARNING** In case of unwanted movement or motion, press the

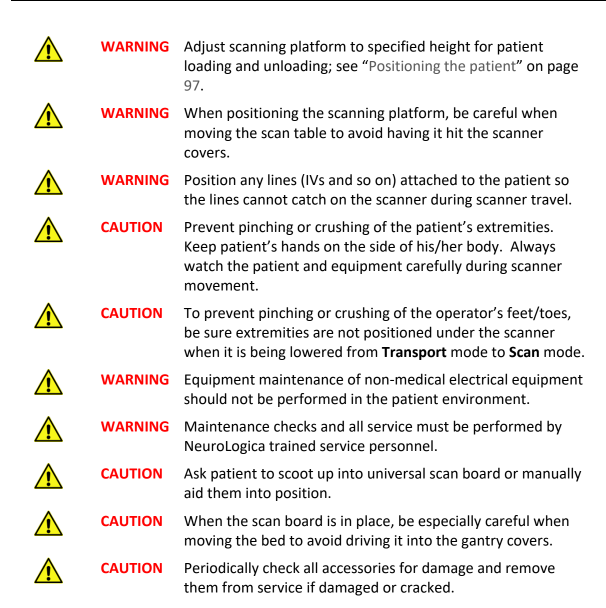
**EMERGENCY STOP** (E-STOP) button.



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 (see "Positioning the

patient" on page 97).



# Radiation safety

Two **Dangerous to patient and operator** labels are affixed to both sides of the scanner, just above the operator controls. See Figure 6 on page 65.



Figure 6: Dangerous-to-patient/operator safety-warning label location (left) and label (close-up, right)

**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 posed by 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 to prevent exposure to unwanted radiation. **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.

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

- The scanner is in the **Scan** mode position.
- The workstation is connected.
- The **START** button is activated when the patient is registered, the protocol is selected, the **Begin** button is clicked, and the protocol is prepared. The **START** button on the scanner's control panel illuminates when the scanner is ready to begin. See Figure 8 on page 67 to identify the START button.

## 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 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 BodyTom Elite 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 isolate 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, below the control panel.

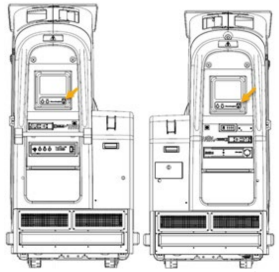


Figure 7: BodyTom Elite E-STOP locations (right and left)



Figure 8: Close-up of the scanner control panel and the E-STOP button

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

# **Battery safety and information**

The **System battery capacity** icon shows an indication of the scanner's 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; there are 145 lithium-ion batteries in the scanner; 144 are used for scanning and the remaining battery is used for moving the scanner while in transport mode.



**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 25.

## **Battery replacement and disposal**



WARNING Battery replacement is to be performed by authorized and

trained NeuroLogica service personnel, to ensure proper

disposal of hazardous material.



WARNING Dispose batteries in accordance with federal, state, and local

regulations.



**WARNING** Do not incinerate batteries.



WARNING Contact an authorized NeuroLogica representative for

appropriate, product-disposal instructions.

### Scanner battery capacity

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



Figure 9: Scanner battery capacity icon

## Run time operation

During normal, run-time operation, the battery capacity is being calculated one time, per second. The **battery capacity** indicator is updated on the scanner's **display screen** as well.

Note The battery capacity indicator on the scanner's display screen is displayed in 5% increments when above 10%. When 10% and below, the displayed capacity is in 1% increments. That is, above 10% capacity, the displayed value is always rounded to the closest multiple of 5 (for example, 93 gets rounded to 95, 42 gets rounded to 40, 47 gets rounded to 45, and so on).

### **State changes**

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

Low voltage alarm state

When **battery capacity** goes below 25%, a periodic alarm will sound. It will remain in this state until the battery capacity has gone back up to 27% or higher. The **Start** and **Cancel** buttons when scanning will not illuminate if the battery is too low.

Low voltage lock-out state

When the battery capacity goes below 1%, the scanner screen-display buttons are disabled and starting a scan is prohibited; for example, the ability to move the scanner and certain protocol buttons are disabled. It will remain in this state until the battery capacity has gone back up to 2% or higher. The low voltage alarm will continue to be active in this state.

### **Predictive scanning**

Before each scan, battery usage for that scan is predicted based on the selected load factors (for example, kV, mA, scan time) and is compared against the available battery capacity. If there is not enough battery capacity to scan, a popup appears on the workstation 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.

### Workstation

The **Workstation battery capacity** icon shows an indication of the workstation's battery capacity. On the workstation's main screen, place the mouse pointer over the battery icon to see the capacity of the battery, ranging from 0 to 100%. The user should always check the screen to verify the status of the batteries.



Figure 10: Workstation battery capacity icon

Note: If the scanner's display screen is black, the system is not charging and/or the batteries are permanently damaged. A service call is required.



**CAUTION** In newer workstations, the workstation reports battery capacity on the workstation remote power display (under the monitor). The battery system is designed to be replaced by authorized and trained NeuroLogica service personnel, only.



**CAUTION** The workstation will not report the proper battery capacity and status if a network connection is not made.



**CAUTION** The system can only be charged from a correctly rated wall outlet. A rating information plate is located on the product system label (lower backside panel or lower left side panel, see Figure 2).



**CAUTION** The system (including the workstation) 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 reports to be 0%, permanent battery damage can occur.



**CAUTION** The power cord selection must not be less than 110v/12A (USA) and 220v/7.5A (EU and Asia), 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.

## Scanner X-ray tube capacity

The percentage of the tube capacity required for a scan =  $((kV \times mA \times scan))$ time(s))/180000) x 100%. Approximately 0.11% capacity is regained each second during cooling.



Figure 11: Scanner X-ray tube capacity icon

### 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 service personnel. Service personnel use Service manual (1-NL4000-062) to effectively perform needed service and preventive maintenance and inspection of the system. See "Contact information" on page 25 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 service personnel at NeuroLogica Corp. See "Contact information" on page 25 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** NeuroLogica recommends that a quarterly preventive maintenance be conducted by NeuroLogica's service personnel/trained facility bioengineer.

Instructions for replacing serviceable parts are identified in the Service Manual (1-NL4000-062).

# Cybersecurity



**WARNING** Upon detection of a cybersecurity threat to the system or workstation, 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 goahead 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 scanning.

## Personnel privileges and terminology

## Qualified operator

The operator 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 CT system. See "Understanding the types of users" on page 101 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 User ID 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 CT system. This privilege level allows use of all clinical protocols to scan the patient.

# **Protocol privileges**

Protocol privileges are granted to a qualified healthcare professional, 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 or modify clinical protocols on the CT system. A healthcare professional with protocol privileges does not necessarily need scanning privileges on the CT system.

### Administrative privileges

Administrative privileges are granted to qualified healthcare professionals 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 CT system 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 need scanning privileges or protocol privileges on the CT system.

#### Clinical operation

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

# **Clinical scanning**

CT system operation that involves scanning of patients.

# **Clinical protocol**

A protocol on the system intended for use on patients.

#### Kernel

The kernel or filter 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

The BodyTom Elite CT system lets you scan patients in a room or ward, an Emergency Room (ER), Operating Room (OR), Radiology, and procedure rooms.

This chapter provides a brief introduction to the BodyTom Elite system. You will learn how to use the BodyTom Elite system (to see an illustration, see Figure 12 on page 75) – in subsequent chapters.

Note Whether you turn on the scanner or the workstation first (see Chapter 4) does not matter; however, it is advised to power up the BodyTom Elite system hardware first, to allow time for the scanner to warm up.

# **BodyTom Elite system**

The BodyTom Elite is a 32-slice, mobile, battery-operated CT scanner and software system with **Axial**, **Helical**, and **Dynamic**, capabilities.

The BodyTom Elite core system consists of the scanner, the workstation, and the phantom holder. Consider the following:

- The scanner and workstation communicate using a wireless connection. They communicate using an ethernet connection, if necessary.
- The BodyTom Elite workstation is a computer with custom software
  that allows the user to employ pre-defined, system protocols or
  devise unique protocols for performing patient studies. It also allows
  the user to update patient information and store images. The
  viewing portion of the BodyTom Elite workstation allows the user to
  view images in more detail and includes tools to help facilitate
  diagnosis by a physician.
- The maximum scout length is 2000mm.
- The scanner can scan a slice-thickness of 1.25mm, 2.5mm, 5.0mm, and 10mm in **Axial** mode.
  - In Axial mode, the BodyTom Elite scans 10mm of anatomy with each rotation.
  - o The maximum scan-range in **Axial** mode is 900mm.
- The scanner can scan a slice-thickness of 1.25 x 0.625, 1.25 x 1.25, 2.5 x 1.25, 2.5 x 2.5, 5.0 x 2.5, and 5.0 x 5.0 in Helical mode.
  - In Helical mode, the BodyTom Elite scans 32mm of anatomy at a pitch of 0.8.

- The maximum scan-range in **Helical** mode is 2000mm.
- In **Dynamic** mode, the BodyTom Elite scans 40mm of anatomy.
- The scanner is compatible with surgical navigation, HIS, RIS, and PACS.



Figure 12: BodyTom Elite system configuration

- 1 QA stand and phantom
- 2 BodyTom Elite or gantry
- 3 BodyTom Elite workstation
- 4 Bed with patient near bore

For information on cleaning and storing the scanner, see "Cleaning the scanner and workstation" on page 357 and/or "Storing the system" on page 359.

# Overview of the scanner control panels

Control panels appear on the right and left side of the scanner. The scanner control panel allows the operator to start and stop a scan. All motion and x-ray generation can also be quickly stopped using the EMERGENCY STOP button. After the patient is registered and you select a protocol, you must start the scan from the scanner control panel, which is located under the scanner's display screen. The operator control panel lets you power on the scanner, lift, and lower the scanner, and lock scanner functions.

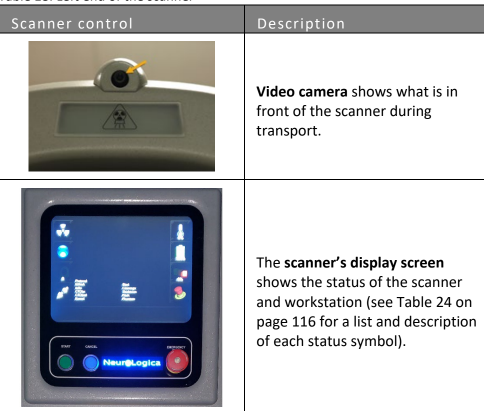
**Note** To determine where personnel should stand during a scan, consult with the hospital physicist. See "Scatter radiation" on page 244.

#### Controls on the left end of the scanner



Figure 13: Left end of the scanner

Table 18: Left end of the scanner



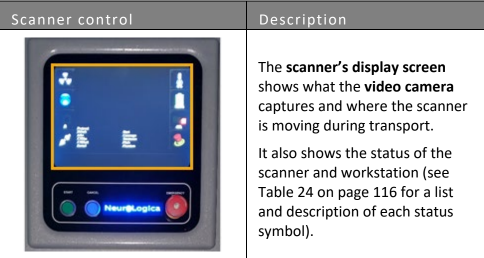
# Scanner control Description **START** and **CANCEL** buttons (to the left) and the EMERGENCY-STOP (E-STOP) button (to the right) on the scanner control panel. **Ethernet port** to connect to the workstation, and four additional data-access ports on the data interface panel. 120VAC/20A and 240VAC/30A plugs. The **pendant**; see "Overview of the pendant" on page 81 for more information.

# Controls on the right end of the scanner



Figure 14: Right end of the scanner

Table 19: Right-end of the scanner



# Description **START** and **CANCEL** buttons (to the left) and the EMERGENCY-STOP (E-STOP) button (to the right) on the scanner control panel. Drive bar and enable bar (arrow). Key lock, the Power-ON and **Power-OFF** buttons for scanner, and the Rocker-Switch-Lift Up and **Down** buttons on the operator control panel. The **pendant**; see "Overview of the pendant" on page 81 for more information.

# Identifying operator control panel buttons



Figure 15: Operator control panel buttons and indicators

Table 20: Operator control panel buttons and indicators

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Operator control panel buttons and indicators	Name	Description
	Rocker- Switch-Lift <b>Up</b> and <b>Down</b> buttons	Press and hold the <b>UP</b> or <b>Down</b> Rocker-Switch-Lift button to raise or lower the scanner.  Lowering the scanner to floor level, makes the scanner ready to scan. Raising the scanner, makes the scanner ready for transport.  When the button is inactive, it is dim; when the button is active it is illuminated.

# Overview of the pendant

The pendant lets you move the scanner, turn on and off the lasers, zero reference the scanner, and program scan and rest positions for the scanner. See Table 21 for a list of what each button activates.

Table 21: Pendant buttons

Pendant	Button	Description	Action
	POWER	POWER	Illuminates when power is supplied to pendant.
	*	LASER	Turns on all three positional lasers. While the lasers are on, the scanner spins for the internal laser to be seen within the scanner opening.
		GO TO SCAN PLANE	Moves the scanner forward approximately 30cm. This is the distance between the internal and external lasers.

Pendant	Button	Description	Action
ROWER  LASER SCAN PLANE  ZZERIO  REFERENCE  GANTRY  GANTRY  SET MEMORY  SCAN REST  POSITION POSITION		ZERO REFERENCE	Sets the scanner to zero before starting a scout or a scan.
	<b>(</b>	MOVE BACKWARD (slow)	Pressing and holding moves the scanner backward 10mm per second.
	<b>&gt;</b>	MOVE FORWARD (slow)	Pressing and holding moves the scanner forward 10mm per second.
	<b>®</b>	MOVE BACKWARD (fast)	Pressing and holding moves the scanner backward 60mm per second.
	<b>&gt;&gt;</b>	MOVE FORWARD (fast)	Pressing and holding moves the scanner forward 60mm per second.
	<b>(2)</b>	SET MEMORY	Allows the user to program <b>Scan</b> and <b>Rest</b> positions for the scanner.
NeureLogica		SCAN POSITION	Moves the scanner to the Scan Position saved using the Set Memory feature.
		REST POSITION	Moves the scanner to the Rest Position saved using the Set Memory feature.



DANGER

Store the pendant in its holder when not in use to prevent accidental and/or unintentional contact by patient and/or users.



Figure 16: BodyTom Elite remote-control pendant

The scanner's position appears on the **positional display** on the front of the scanner. You can use the pendant to zero reference the scanner. The display shows a positive or negative positional number.



Figure 17: Scanner's positional display

**Note** The two light panels on either side of the scanner's position indicate x-ray is active, when lit. The scanner produces an audible alert during scanning.

#### Overview of the workstation



Figure 18: Workstation with leaded-glass shield (optional installation)

The **workstation** is an accompanying part of the scanner; it is the computer and control unit that operates most functions of the system. All basic information related to the workstation (for example, operating distance, warnings and cautions, connectivity, functionality, etc.) appear in Chapter 4 Basic Workstation Operations. The workstation includes the computer, monitor, ethernet connections, and the remote controls. The workstation also includes the **Uninterruptable Power Supply (UPS)**. The workstation can be installed with an optional leaded-glass shield (shown in Figure 18) for additional protection.

The workstation enables you to easily move it wherever you need it to go. The workstation is designed to let you navigate in and out of elevators, over doorway thresholds, or on any type of floor including carpet, with ease.

The workstation can be set up either wirelessly or hardwired to the scanner. The administrator makes sure wireless is enabled before you create a wireless connection between the workstation and the scanner (with **System Configuration > Scanner Setup**). See "Remote Support Setup" on page 134 to learn how the administrator enables the wireless connection.

Note Wireless connections can be slower than a hardwired connection. If an unexpected delay or disconnection, due to environmental/bandwidth interference, occurs when using wireless, it is recommended to hardwire an ethernet cable to the scanner for continued communication.

After respective peripherals are properly plugged in, make sure all applicable power switches are in the on position before using the system.

For any devices connected to an AC outlet, make sure the outlet is providing required power.



**CAUTION** When not in use, the scanner and workstation should always be plugged into power outlets to ensure maximum efficiency.

> The system should be stored in an area with limited access to prevent inadvertent damage.

See "Powering on and off the BodyTom Elite system" on page 90.

**Note** Be sure to keep the workstation plugged in when it is not in use to charge the battery. Charge time while off is ~ (approximately) 2 hours; while in use, charge time is ~ 8 hours.



**CAUTION** Verify that the ethernet cable is unplugged from the workstation to avoid damage to the cable and outlet during transport.

**Note** Before using the workstation, be sure to read and understand how to clean and maintain it. See "Cleaning the scanner and workstation" on page 357.

# Workstation types

There are two types of workstations available: Granite and Dell. For more information on powering on and off the workstation, see "Using the workstation" on page 101.

Note The Granite workstation does not allow the user to wait for the computer to shut down before communicating to the workstation to shut down; this is because the workstation sends a toggling signal to the computer. If the computer is off, it will turn back on. The workstation must already be in the process of shutting down by the time the computer is fully turned off.

The product safety coverage of the specified workstation (Safety Certified to IEC 60950 standards) was evaluated and deemed acceptable for use with the BodyTom Elite to appropriate IEC 60601-1 and/or IEC 60601-1-1 harmonized national standards.

The workstation will be kept outside the patient environment as defined by IEC 60601-1-1. BodyTom Elite is suitable for use inside patient environment.



**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 19, 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.

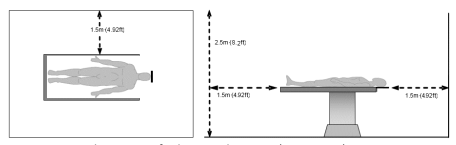


Figure 19: Workstation safe distance location (two views)

# Workstation remote power controls

The following are found on the workstation:

- Remote power controls
- Microphone and controls
- Speaker and controls



Figure 20: The remote power display See "Using the workstation" on page 101.

Note The microphone and speaker are plugged in to the USB port on the back of the monitor. If the port is changed, you will need to reboot the workstation.

#### **Workstation UPS**

The workstation uses an **Uninterruptable Power Supply** (**UPS**) to supply power to the workstation for approximately (~) 6-8 hours when the workstation is unplugged. The **UPS** is located inside the workstation. This feature allows the new workstation to run when it is not plugged in to a wall outlet.

#### Workstation considerations before use

Before using the workstation, consider the following:

- If a problem is detected with the workstation, make sure repairs or adjustments are made to it *before* using it.
- Make sure the workstation operates easily and freely, and all parts work smoothly.
- Check for excess noise, vibration, or a change in the ease-of-use.
  - Noise, vibration, or change in ease-of-use can be signs of a problem and a need for servicing.
- Be sure to read its warnings carefully and completely *before* using the workstation. Do not attempt to service the workstation. Only skilled service personnel are permitted to service the workstation. See "Hazard information" on page 56.



**CAUTION** Users are not to perform service or maintenance on the system at any time. This includes battery maintenance.



**CAUTION** Failure to heed these warnings may cause injury to the user, to others, or damage to the equipment.

**Note** NeuroLogica recommends that a quarterly preventive maintenance be conducted by NeuroLogica's service personnel/trained facility bioengineer.

> Instructions for replacing serviceable parts are identified in the Service Manual (1-NL4000-062).

# Keyboard and mouse

The workstation comes with a keyboard and a mouse (1 for left-mouse) button; 2 right-mouse button).



Figure 21: Workstation keyboard and mouse

# Workstation power cord

The power cord lets you power up the workstation.

Consider the following:

Plug style will vary depending on factory installed elements based on geographic location and voltage requirements.



CAUTION

The power cord selection must not be less than 110v/12A (USA) and 220v/7.5A (EU and Asia), 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.

# Parts that potentially come into contact with the patient

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

- The BodyTom Elite system, especially the painted, external surfaces of the system's covers.
- Universal transfer board, if purchased.

# **Chapter 3** Basic Scanner Operations

Basic scanner skills include powering on and off the scanner, learning how to use and navigate the operator and scanner control panels, how to use **E-STOP**, and how to use the Rocker-Switch-Lift **UP** and **DOWN** buttons to lift and lower the scanner.

Scanning basics you should know before scanning a patient include how your system should be setup, how to position the scanner and the patient before the scan, and how to start a scan from the operator control panel.

**Note** It is recommended that the scanner is on for at least 60-90 minutes prior to performing the daily air calibration or scanning patients.

It is recommended practice that the scanner is plugged in and turned on even when it is not in use.

# Powering on and off the BodyTom Elite system

The BodyTom Elite is not intended to be turned on and off; however, if the system should lose power, *it is advised* to power on the scanner first, to allow time for the scanner to power up.

#### 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 only power source
  that allows the scanner to perform scans.
- 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 centipedes during scanning.



Figure 22: AC cord and storage on scanner (120V left plug in or 240V right plug out)



Figure 23: Scanner's power cord receptacle for 120VAC and 240/VAC

Be sure to keep the workstation plugged in when it is not in use to charge the battery. Charge time while off is  $^2$  hours; while in use, charge time is  $^8$  hours.

# Checking a connection between the workstation and the scanner

To check if a wireless connection exists between the scanner and the workstation, look for the **Wireless connection** icon on the scanner's display screen:

**Note** You must be logged into the workstation before this icon is visible on the scanner's display screen.

If the workstation is connecting to the scanner by hardwire, check if the supplied ethernet cable is connected between the workstation and scanner.

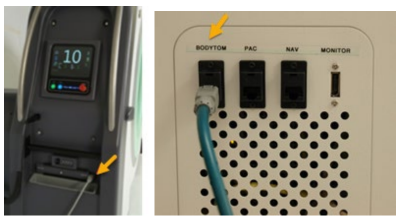


Figure 24: Scanner hardwired to the workstation with an ethernet cable to dataaccess ports

The administrator makes sure wireless is enabled when creating a wireless connection between the workstation and the scanner (with **System Configuration > Scanner Set U**p, through the workstation). See "Remote Support Setup" on page 134.

Note Make sure that the wireless setup in System Configuration has been done before making a wireless connection between the workstation and the scanner. See "Scanner Setup" on page 133 and "Remote Support Setup" on page 134.

Wireless connections can add lag time when compared with a hardwired setup. If an unexpected delay or disconnection occurs when using wireless, it is recommended to hardwire an ethernet cable to the scanner for continued communication.

# Moving and transporting the scanner

To move the scanner, the scanner must be in **Transport** mode. The Rocker-Switch-Lift **UP** and **DOWN** buttons are located on the side of the scanner, on the operator control panel. These **UP** and **DOWN** buttons prepare the scanner to move up for transporting or down for positioning the scanner before scanning a patient. To set the scanner in **Transport** mode, go to the operator control panel and follow the procedure below.

Note If the scanner has been calibrated for multiple rooms, you must select the room prior to scanning to ensure the correct floor-calibration file is loaded. See "Selecting a room for the BodyTom Elite" on page 188.



#### **CAUTION**

Before transporting the scanner, verify that the ethernet cable is unplugged from the workstation 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.

1. Press the **Transport** button.



Figure 25: Transport button on the operator control panel

2. Press and hold the Rocker-Switch-Lift **UP** button to raise the scanner off the centipedes for transport.



Figure 26: Rocker-Switch-Lift UP (top) and DOWN (bottom) button

**Note** Hold the top Rocker-Switch-Lift **UP** button until the scanner is completely raised and on its castor wheels. The top Rocker-Switch-Lift **UP** button illuminates when it is in the up position as shown in Figure 26.

3. Grip the drive bar with both hands.



Figure 27: Drive bar front



Figure 28: Drive bar

Note The drive bar is equipped with an enable bar. The enable bar is a darker soft rubber material located on front of the drive bar. While holding the drive bar, squeeze the rubberized enable bar to activate system motion. If you let go of the enable bar, the scanner stops.

4. Move the scanner.

The following points explain how to hold the **drive bar** to move it forward, backward, left, and right:

- Push forward with both hands and use equal pressure to move the scanner forward.
- Push with the right hand to turn the scanner left; push with left hand to turn the scanner right.
- Pull back on the **drive bar** to move the scanner in reverse.

Note A three-point driving technique is required to turn in smaller spaces.



CAUTION

NeuroLogica recommends two people move the scanner within the facility: one to steer and another, in front of the scanner, to insure there are no obstacles. Two people ensures there are no collisions while maneuvering through tight hallways and around corners.



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



**WARNING** When transporting, use the video camera and scanner's display screen as guides to avoid hitting objects.



CAUTION

To prevent potential for injury from overbalancing and/or tipping, do not attempt to turn this system on an incline during transportation.



WARNING

**Do not** move the system right or left if transporting on an incline becomes necessary. Always keep the system in a straight motion. Contact **Technical Support** for assistance when movement is required on an incline.



CAUTION

The BodyTom Elite scanner is larger than most medical equipment. Therefore, NeuroLogica recommends proper training and practice.

#### Drive direction of scanner



Figure 29: Scanner drive direction (right side view)

# Safety bumper system

The scanner's transport system is equipped with an active, **safety-bumper** system. Each safety bumper is electronically controlled to terminate motion in the direction in which the system is moving when the bumper is activated. If a bumper is pressed due to a collision, the drive system is disabled in that direction. All other directions are still enabled to allow you to reposition the scanner away from any impact area. The activation force needed to trigger the bumper system is approximately 7lbs.



Figure 30: Bumper system

# Positioning the scanner before a scan



**WARNING Never** raise or lower the scanner when the patient is positioned in the system's bore. Always move the patient and table away from the bore before raising or lowering the system itself.

1. Ensure the scanner is in **Scan** mode.

To move the scanner, see "Moving and transporting the scanner" on page 92. See also "Performing a scan" on page 264.

- 2. To lower the scanner, press and hold the Rocker-Switch-Lift **DOWN** button until the scanner is completely lowered on its centipedes. The Rocker-Switch-Lift **DOWN** button illuminates when the scanner is completely lowered on the centipedes.
- 3. Align the patient and table with the scanner, ensuring the patient is in the center of the Field Of View (FOV), also ensure that the scanner will not collide with the 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 centipedes.

4. Make sure the patient scan table is locked.



**CAUTION** Make sure to lock the patient bed or scan table to prevent it from moving during the scan.

5. Adjust the scan table height so that the patient is centered within the bore.



Figure 31: Patient centered in bore (height positioning)

# Positioning the patient using the laser lights



WARNING

Before scanning, position the patient in such a way that extremities, hair, life-support equipment, etc. have sufficient clearance to prevent contact with scanner and or when used with accessories and options, such as head frames, scan boards, etc.



WARNING

Make sure the patient is supported properly when positioned (both height and alignment) to prevent injury during scanning.



WARNING

Make sure the foot pedal brake on the scan table/bed is engaged to prevent it from moving during the scan.



WARNING

**Never** raise or lower the scanner buttons on the operator control screen when a patient is positioned in the system's bore. **Always** slide the scan table away (by disengaging its brake) from the system 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





**CAUTION** 

If the patient becomes nauseated or is unable to be still (motionless), stop the scanner immediately using the **E-STOP** button.

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

The scanner provides a rotating laser-light to guide you to properly position the patient. The laser indicates the center of the scan plane. The laser light is always centered on the actual x-ray beam. For multi-slice protocols, this means that the laser light will indicate the middle position of all simultaneous acquisitions being acquired. There is no offset between the laser-light plane and the actual scan-plane. The accuracy of the position of the laser plane, with respect to the scan plane, should be +/- 2mm.

There are two sets of laser lights: one set of external and one set of internal. The external set contains the following: 1 **Coronal** positioning, 2 **Transverse** laser, 1 mid-**Sagittal** laser. The internal set contains lasers affixed to the x-ray tube; the lasers rotate around the patient to designate internal zero-point reference. Keep the following in mind:

- To adjust vertical or horizontal positions, use table and bed controls, only.
- To adjust Z axis, walk the scanner to position using the pendant.

See "Overview of the pendant" on page 81 to see how buttons act.

- 1. Position the patient on the bed.
- 2. Align the bed to the scanner and make sure there is sufficient clearance around the patient by positioning the patient in the center of the **FOV**.



Figure 32: Phantom positioned in center of FOV

3. Use the pendant positioning buttons to center the patient within bore.



Figure 33: Pendant use for positioning lasers upon patient

- 4. On the pendant, press the **LASER** button to turn on all positional lasers. The following three lasers are available to help with positioning.
  - 1 set of external, lateral-side-positioning lasers (**Coronal** positioning for table height).
  - 1 external, crosshair-positioning laser (top, to designate the mid-Sagittal plane).

 1 external crosshair positioning laser (side-to-side to designate the Transverse plane).



Figure 34: Positioning lasers upon patient See the laser precautions in "Laser safety" on page 58

5. Use the scanner's positional display to identify position of the scanner.



Figure 35: Positional display

- 6. Consider the following:
  - 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.

### **Operating the E-STOP button**

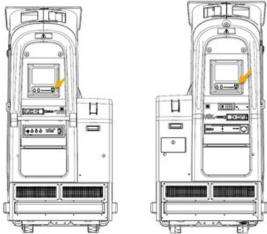


Figure 36: BodyTom Elite E-STOP locations (right and left)



Figure 37: E-STOP button on the scanner control panel on both the left and right sides of the scanner

- 1. Press the **E-STOP** button to perform the following:
  - Stop system (if it loses control).
  - Stop all system motion and x-ray.
  - Remove power to the gantry drives and x-ray system.
  - If the scanner 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 after pressing the **E STOP** button.

# **Chapter 4** Basic Workstation Operations

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

Note Whether you turn on the scanner or the workstation first (see Chapter 4) does not matter; however, it is advised to power up the BodyTom Elite system hardware first, to allow time for the scanner to warm up.

# Understanding the types of users

There are three **User Levels** available on the workstation: administrator, limited operator, and restricted operator. User ID's 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 it's configurations. Can
Administrator	create protocols, User ID's, and passwords, as well as
	access all functions of the system.
	Modified access to the system. Users with Limited
Limited	access can modify protocols during scanning but
	cannot create and save protocols; has no access to
	system configurations.
	Users with Restricted access can scan with the
Restricted	system but are unable to make any changes to
	protocol parameters while scanning, they also have
	no access to system configurations.

# Using the workstation

The workstation uses an **Uninterruptable Power Supply** (**UPS**) to supply power to the workstation for ~6-8 hours when the workstation is unplugged. The workstation includes a computer, the remote power display, and a microphone and speaker.

# Identifying the workstation's remote power display

The **Power On** button on the remote power display allows you to power on the Granite workstation.



Figure 38: Workstation remote power display

The following describes the action of each workstation power-control button.

Table 22: Workstation power-control buttons

Workstation power- control button	Button or indicator name	Description
00000	UPS battery level	Shows the battery usage; each LED represents 20 percent of battery power.
$\sim$ •	Power-On	When the system has power, the LED light illuminates.
! 0	Alarm	When the system is at or less than 20 percent battery power, an alarm sounds, and the LED light illuminates to warn you.
மு	Power-On and Power-Off	Press and hold the <b>Power-On</b> / <b>Power-Off</b> button for 3-5 seconds to turn on and off the workstation.
<b>*</b>	Mute	Press this button to silence the alarm.

# Identifying the microphone, speaker, and controls



Figure 39: Microphone, speaker, and controls

Table 23: Speaker control buttons

Microphone button	Name	Description
<b>A</b>	Microphone mute	Press the <b>Mic Mute</b> button to mute the microphone.  NeuroLogica recommends using the <b>Mute</b> button located on the bottom right of the workstation screen.
<b>4</b> ×	Mute	Click the <b>Mute</b> button, on the bottom right of the workstation screen to mute the microphone and speaker.
	Speaker	The <b>Speaker</b> button appears on the bottom right of the workstation screen when the speaker is activated to hear the patient.
+	Volume	Press the "-" button on the left side of the speaker to decrease the volume, press the "+" button on the right side of the speaker to increase the volume. The speaker has illuminated volume lights to indicate volume level.

# Powering the workstation

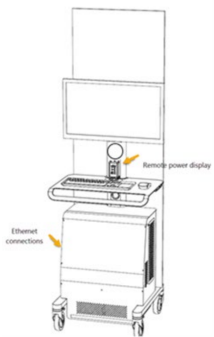


Figure 40: Remote power display on workstation

**Note** Depending on the workstation, the **Power-On** and/or **Power-Off** button may not be in this location.

### To power up a Granite workstation

 Press and hold the Granite workstation's Power-On/Power-Off button on the workstation's remote power display until the green light blinks.

### To power down a Granite workstation

1. Press and hold the Granite workstations **Power-On/Power-Off** button on the workstations remote power display until the green light blinks.

**Note** The Granite workstation does not allow you to wait for the computer to shut down before communicating to the workstation to shut down; this is because the workstation sends a toggling signal to the computer. If the computer is off, it will turn back on. The workstation must already be in the process of shutting down by the time the computer is fully turned off.

#### To power up a Dell workstation





Figure 41: Dell workstation power on button

#### To power down a Dell workstation

- 1. Click File
- 2. Click Shutdown Computer.
- 3. On the Shutdown Computer? popup box, click Yes.

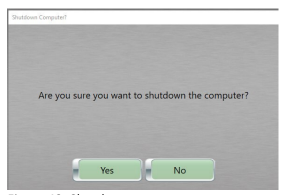


Figure 42: Shutdown computer popup

# Logging in to the workstation

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

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

User ID
Administrator
Password

Login Shutdown

1. Click the User ID dropdown and select your User ID.

Figure 43: User ID dropdown box



Figure 44: User ID dropdown list

2. Click in the **Password** field and type your password. Passwords are case sensitive.



Figure 45: Password text box

3. Click the **Login** button.

If the User ID and/or password are invalid, a prompt appears to reenter the information.

Note You have a limited number of login attempts before the system locks the account. An administrator can unlock the account. See "System and User Configuration and Setup" on page 121 for more details about unlocking an account.

When the User ID and password are verified, you are logged into the system.

The main screen to the workstation software appears with the **Patient Registration** tab open.



Figure 46: Patient Registration tab

4. Verify that the correct User ID appears at the top center of the screen.



Figure 47: User ID, current date, and time

If the correct User ID *does not* appear, contact your supervisor or administrative user to verify the account.

#### Locking and unlocking the workstation

The **Lock** button indicates whether your system is in **Lock** or **Unlock** mode. This button is a toggle, if the workstation is locked, the **Unlock** button shows, if the workstation is unlocked, the **Lock** button shows.

To prevent unwanted personnel from accessing the system, you should lock the workstation if you intend to leave the area for any period. When you lock the workstation, it remains **on**, but no one can access it without supplying a User ID and password.

#### Using Lock if you need to step away from the workstation

1. Click the **Lock** button located at the top-and center of the workstation screen.



Figure 48: System Lock button





Figure 49: Lock/Unlock System popup to lock the workstation

- 2. Enter the User ID and password by selecting the option from the dropdown and entering information in the **Password** field.
- 3. Do one of the following:

- Click the Lock button to lock your system.
   When you select this option the Lock button changes to the Unlock button.
- Click the **Cancel** button to return to your work.

## Using Unlock to view your work

1. Click the Unlock button.



Figure 50: Unlock button

## The Lock/Unlock System popup appears.



Figure 51: Lock/Unlock System popup to unlock the workstation

- 2. Enter the User ID and password by selecting the options from the dropdown and entering information in the **Password** field.
- 3. Do one of the following:
  - Click the Unlock button to unlock your system.
     When you select this option the button changes to the Lock button.
  - Click the **Cancel** button to return to your work.

# Navigating around the workstation's main screen

The workstation screen shows similar information, even if the screen elements change when you perform different actions. These constant screen elements are as follows:

Main menu options	Appears in the top left corner of the screen and shows the <b>File</b> , <b>Tools</b> , <b>Customize</b> , and <b>Help</b> commands.
User, date, and time	Appears in the center of the screen and displays a greeting for the user, the signed-in user's name, the current date, and time.
Scanner and workstation status	Appears in the top right corner of the screen and displays status information for both the scanner and workstation.

The main screen to the workstation software always opens with the **Patient Registration** tab.

#### Brief overview of the main menu

The main menu provides you access to the basic functions from the commands: File, Tools, Customize, and Help.

The main menu appears on every screen and is always located in the same location, regardless of what you are doing.



Figure 52: Main menu

Note To select commands, click the first command and the subsequent commands. For example, if you see

Customize > System that means, click Customize from the main menu and then click System.

#### Brief overview of the File menu

When you log off, restart, or shutdown the application and/or the workstation, you must use your User ID and password to log back in.

# Logging off the system

1. Click File from the main menu.



Figure 53: File menu

The File dropdown appears.



Figure 54: File > Log Off

2. Click **Log Off** from the dropdown to shut down the software without shutting down the workstation.

The Login popup appears.



Figure 55: Login popup

This is also the login and shutdown portal. You can login or shutdown the workstation from this popup.

## Restarting the application and/or computer

You can restart the computer in two ways: restarting the application only or restarting both the application and the computer.

1. Click **File** from the main menu.



Figure 56: File dropdown menu

- 2. Perform one of the following:
  - Click Restart Application from the dropdown to restart only the application software.
  - Click **Restart Computer** to restart both the application and the computer.

The following **Restart Application** or **Restart Computer** popup appears.





Figure 57: Restart Application or Restart Computer popup

- 3. Perform one of the following:
  - Click the Yes button to restart the workstation.
  - Click the No button to return to the screen.

## Shutting down the computer

1. Click **File** from the main menu.



Figure 58: File dropdown menu

- 2. Click **Shutdown Computer** from the dropdown. The **Shutdown Computer** popup appears.
- 3. Perform one of the following:
  - Click the Yes button to shut down the workstation.
  - Click the No button to return to the screen.

#### Brief overview of the Tools menu

This menu provides you with tools to store and print, set up protocols, and test your system to ensure it is operating as specified.



Figure 59: Tools dropdown menu

- 1. Click **Tools** from the main menu.
- 2. Click one of the following from the dropdown:

Store/Print Queue	The <b>Store/Print Queue</b> displays the status of studies being archived. You will learn more
	about how to store to various media later in
	this user manual; see page 121.
Protocol Manager	Allows users with <b>Administrative</b> privileges
	to create, modify, delete, and/or upload
	protocols to the scanner. You will learn
	more about how to use Protocol Manager
	later in this user manual; see page 189.
	The tool that verifies the system is at its
Quality Assurance	optimum performance. You will learn more
(QA)	about QA later in this user manual see page
	222.

#### Brief overview of the Customize menu

This menu provides you with tools to set up the system as well as define user profiles.



Figure 60: Customize dropdown menu

1. Click **Customize**, then one of the following sub commands from the drop-down list:

System	Allows users with Administrative privileges to customize site-related settings; see "Chapter 5 System and User Configuration and Setup" on page 121.	
User	Allows you to customize layouts in the system as well as set the password. See "Chapter 5 System and User Configuration and Setup" on page 121.	
Select Room	Allows you to identify and select the room the scanner will be used in. See "Selecting a room for the BodyTom Elite" on page 188.	

## Getting Help from the Help menu

NeuroLogica Help includes an online user manual and information about the system. It also provides remote support from NeuroLogica **Technical Support** for file transfer, remote upgrades, or system review and support. When you enter a six-digit number, **Technical Support** will take control of the system to retrieve files or review the issue in question.



Figure 61: Help dropdown menu

## Getting an online user manual

To open a .pdf version of this user manual:

- 1. Click **Help** from the main menu.
- Click User Manual from the dropdown list.A PDF version of this manual will be opened.

## **Getting remote support**

- 1. Click **Help** from the main menu.
- Contact NeuroLogica Technical Support. See "Contact information" on page 25.
- 3. Click **Remote Support** from the dropdown list. The **Support Connection** window appears.

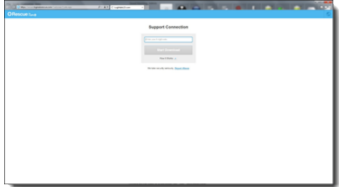


Figure 62: Support Connection browser window

When connected to **Remote Support**, a NeuroLogica **Technical Support** representative will supply a six-digit code to start a remote support session, which allows the support representative to review your system and troubleshoot the issue.

#### Getting information about the product and NeuroLogica

To get additional information about the product and NeuroLogica:

- 1. Click **Help** from the main menu.
- 2. Click **About** from the dropdown list.



Figure 63: About Us popup

The following information is found:

Version(s)	Identifies the current software versions for the
	system.
Licensed To	Identifies who the product is licensed to.
Station AE	Identifies the title for your workstation (for PACS
Title	purposes).
Licensed	Identifies any licensed packages available on the
Packages	system.

# Getting to know the status bar

The status bar appears in the top-right portion of the screen. The status bar provides a quick view of the systems current state. Details for the icons on the status bar are in the tables below.



Figure 64: Scanner and workstation status bar

Table 24: Status bar icons

Status bar	Status bar icon	Status description
icon	Radiation status	Identifies x-ray as on or off. The icon changes from a gray/black icon when x-ray is off, to a rotating yellow/black icon when x-ray is on.
	System state	Identifies the system's current state. The orb changes color depending on the state the system is in. See Table 25 on page 118 for a list of the different orb colors and system states they identify.
	Scanner position	Identifies the system's current position relative to its zero reference.
8	System E-STOP status	Identifies when <b>E-STOP</b> is engaged. The icon will flash when <b>E-STOP</b> is pressed.
٩	System tube heat status	Indicates the current X-Ray tube heat status. The values are color coded as follows:  Blue 0% - 25%  Yellow 26% - 50%  Orange 51% - 75%  Red 76% - 100%

Status bar icon	Status bar icon name	Status description
	Scanner 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%
	System air freshness status	Indicates the air freshness status; it is recommended that an air calibration be performed:  • Every eight (8) 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%  After calibration it returns to 100%.

Status bar icon	Status bar icon name	Status description
		Indicates the remaining workstation battery capacity available. The capacity values are color coded as follows: Green 100% - 21% Yellow 20% - 11% Red 10% - 0%
	Workstation battery capacity status	You will be prompted to plug the workstation into an outlet to charge if the battery capacity is low; a scan cannot complete when the battery capacity is 10% or lower.
		When the workstation reaches the red capacity range, the system will shut down. A message informs you that the system will shut down due to low battery.
		The lightning bolt icon signifies that the workstation is currently charging and goes away when unplugged.
()	Image storage space status	Indicates the available disk space for image storage. The available space values are color coded as follows:  Green 100% - 51% Yellow 50% - 20% Red 19% - 0%

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.

Table 25: System state orbs

Orb	Color	State
	Dark gray	The system is in an unknown state.
	Light gray	The system is powering up or down.

Orb	Color	State
	Dark purple	The system is busy.
	Purple	The system is completing air calibration.
	Light purple	The system is archiving.
	Blue	The system is idle.
	Green	The system is ready to perform a scan.
	Light yellow	The system is planning.
	Dark yellow	The system is preparing.
	Light orange	The system is reconstructing.
	Dark orange	The system is scanning.
	Pink	The system is not ready.
	Red	The system is in fault.

#### The workstation tabs

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



Figure 65: Workstation tabs to perform a patient examination

The tabs include active tabs which will be green, and inactive tabs which will appear gray. The active tabs are **Patient Registration**, **Post Reconstruction**, and **Patient Browser**. The **Acquisition** and **Viewing** tabs require additional steps to be performed before they become active. The following actions are available in each tab:

Patient	Allows you to register a patient either manually or
Registration	from the hospital's database sites.

Acquisition	Allows you to select a protocol and perform the
	examination. This tab is inactive until a patient is
	registered.
Post	Allows you to manipulate raw data in different
Reconstruction	parameters and settings after your scan is
	completed.
Patient Browser	Allows you to view, manipulate, and archive scans
	already performed.
Viewing	Allows you to view patient images This tab is
	inactive until a study is loaded from Patient Browser.

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.

# Chapter 5 System and User Configuration and Setup

A user with administrative privileges must set up the BodyTom Elite system configurations for other users. System configuration is used to set up the scanner 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.

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 administrator 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 **Customize > System** and provides a brief description of each.

Table 26: System configuration tabs

Tab name	Description
General Settings	Allows the administrator to set system configurations such as hospital name, wireless settings, and dose report settings.
User Accounts	Allows the administrator to create and edit user accounts and permissions.
DICOM Servers	Allows the administrator to set up DICOM servers for archiving, such as PACS or HIS/RIS.
DICOM Setting	Allows the administrator to view and configure DICOM tags for HIS/RIS, MPPS, and Patient, Study, Series, and Image modules.
Audio Configuration	Allows the administrator to upload default audio files with protocols; also lets the administrator record, play, and remove audio files.
Dose Configuration	Allows the administrator to set up dose notifications, dose alerts, and configure dose limits for specific scans.
Windowing Presets	Allows the administrator to view and modify kernel and window width and window level presets.

Tab name	Description
Audit Trail Viewer	Allows the administrator to view and log all changes as well as actions in the system, including user logins, patient registrations, and series updates.
Image	Allows the administrator to view and modify how
Orientation	images are oriented in the system.
Filter Kernels	Allows the user to activate custom kernel options for both Axial and Helical scans.

## Setting user accounts

Only a user with administrative access can update another user's account, add a new user, delete a user, and lock or unlock a user's access.

- Click Customize > System from the main menu.
   The System Configuration dialog box appears.
- 2. Click the User Accounts tab.

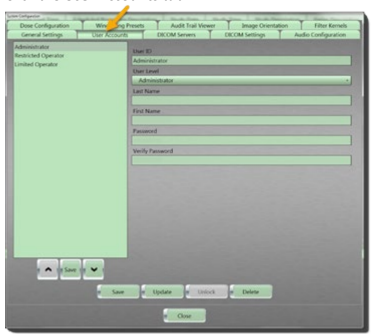


Figure 66: User Accounts tab

- 3. For the **User ID** field, enter the User ID name.
- 4. For the **User Level** field, enter one of the following user levels:

	Full access to the system and it's configurations.
Administrator	Can create protocols, User ID's, and passwords,
	as well as access all functions of the system.

	Modified access to the system. Users with
Limited operator	Limited access can modify protocols during
	scanning but cannot create and save protocols;
	has no access to system configuration.
	Users with Restricted access can scan with the
Restricted	system but are unable to make any changes to
operator	protocol parameters while scanning, they also
	have no access to system configuration.

- 5. For the **Last Name** field, enter the user's last name.
- 6. For the **Enter First Name** field, enter the user's first name.
- 7. For the **Password** field, enter the user's password.

**Note** The password must contain 8 to 12 characters, and must include one number, one symbol and one letter.

8. For the **Verify Password** field, re-enter the user's password to confirm the password.



Figure 67: User account fields filled in

Click the **Save** button.The user is added to the list.

The **Save Aborted** popup appears if your password does not meet the rule for passwords. If this is the case, return to the step above, and fulfill the password rule.

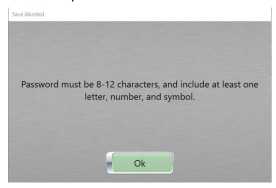


Figure 68: Save aborted popup message – Password requirements

10. Click the **Close** button to exit.

## Setting or updating the user's information

- 1. Click **Customize > System**, from the main menu. The **System Configuration** dialog box appears.
- 2. Click the User Accounts tab.
- 3. Select a user from the list of users.



Figure 69: List of users

4. Modify the user's information; for example, password, or user's first and last name.

**Note** The password must be 8-12 characters and include at least a letter, number, and symbol.

The user's changes take effect after clicking the **Update** button. It is recommended that you log off and log back on and check that the password is working.

5. Click the **Update** button to keep the change(s).

The **Update Aborted** popup appears if your password does not meet the rule for passwords. If this is the case, return to the step above, and fulfill the password rule.

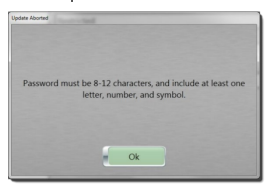


Figure 70: Update Aborted popup message – Password requirements

6. Click the Close button to exit.

## Unlocking the user

- 1. Click **Customize > System** from the main menu. The **System Configuration** dialog box appears.
- Click the User Accounts tab.
- 3. Select the user to unlock from the list of users in the panel.

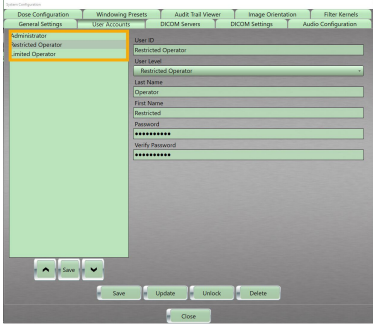


Figure 71: List of users not selected

4. Click the **Unlock** button.

The user's changes take effect after clicking the **Update** button. It is recommended that you log off and log back on and check that the password is working.

## Deleting the user

#### Note The administrator user cannot be deleted.

- 1. Click **Customize > System** from the main menu. The **System Configuration** dialog box appears.
- 2. Click the **User Accounts** tab.
- 3. Select the user to delete from the list of users.

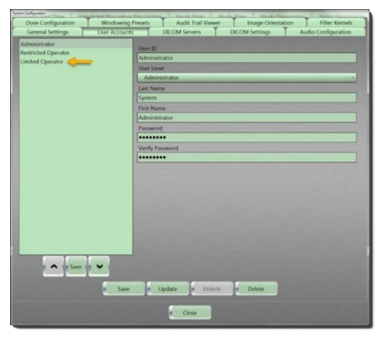


Figure 72: List of all available users

- Click the **Delete** button.
   The **Action Succeeded** popup box appears.
- 5. Click the **Ok** button in the **Action Succeeded** popup.
- 6. Click the **Close** button to exit.

# Modifying the order of the user in the accounts list

- 1. Click **Customize > System** from the main menu. The **System Configuration** dialog box appears.
- 2. Click the User Accounts tab.
- 3. Select the user order to modify from the list of users in the panel.
- 4. Click the **Down** arrow to move the user down the list.

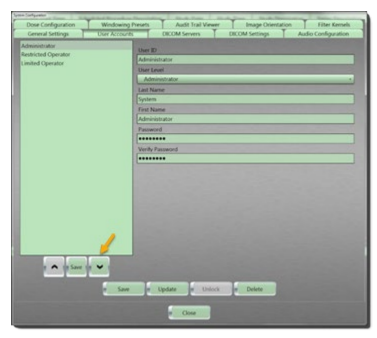


Figure 73: Down arrow

5. Click the **Up** arrow to move the user up the list.

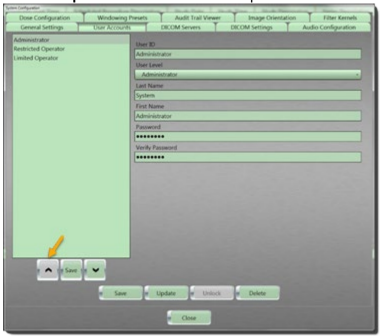


Figure 74: Up arrow

The **Up** arrow will not activate until you move down the list of users.

6. Click the **Save** button under the user list to keep the new user list order.

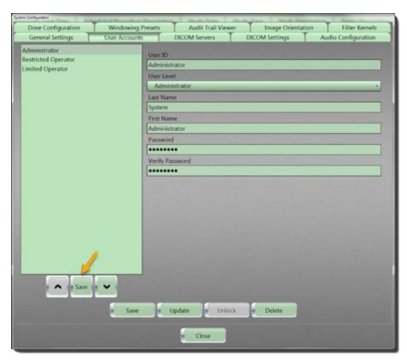


Figure 75: Save button for list order

- 7. Click the **Save** button next to the **Update** button to keep other changes.
- 8. Click the **Close** button to exit.

## Assigning general settings

Perform the following to set how the hospital, workstation's application, scanner, and remote support are configured.

- 1. Click **Customize > System** from the main menu. The **System Configuration** dialog box appears.
- 2. Click the **General Settings** tab.

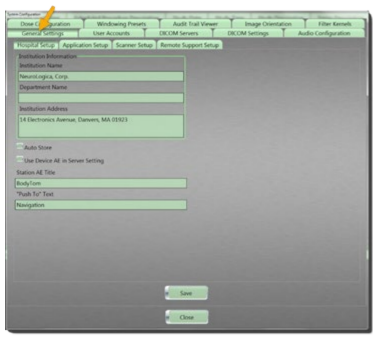


Figure 76: General Settings tab

3. The following tabs are available to the administrator, *only*:

Hospital Setup	Sets up hospital information specific to the site.
Application	Sets up application information the user will
Setup	experience.
Scanner Setup	Sets up scanner IP address information.
Remote Support	Sets up IP address information to allow
Setup	NeuroLogica remote support.

The following sections provide detailed information related to the options available in the **General Settings** tabs.

# **Hospital Setup sub tab**

- 1. Click **Customize > System** from the main menu. The **System Configuration** dialog box appears.
- 2. Click the **General Settings** tab.
- 3. Click the **Hospital Setup** sub tab.

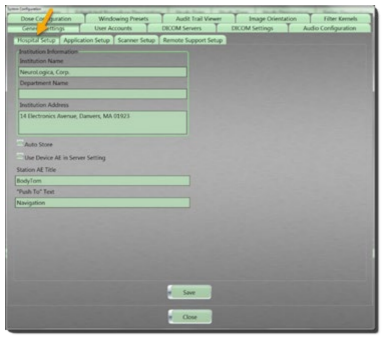


Figure 77: General Settings > Hospital Setup subtab

Enter the institution name in the Institution Name field.

#### **Note** The name appears on all images.

- 5. Enter the department name in the **Department Name** field.
- 6. Enter the institution address in the **Institution Address** field.
- 7. Click the following options that are applicable:
  - When Auto Store is selected and you Finalize a scan, the system will automatically send the images to the Default PACS server defined on the DICOM Servers tab. (see System Configuration > DICOM Servers > Servers). The status of your export will appear in the bottom right of the Patient Browser screen.
  - Click the Use Device AE in Server Setting option to apply the BodyTom Elite AE title tag to DICOM when the operator archives the image data to PACS.
    - If enabled when sending images to PACS, this option will include the BodyTom Elite AE setting as a DICOM tag.
- 8. Enter the system name (for example BodyTom Elite) in the **Station AE Title** field.
- Enter the name you want to call the archive option in the "Push To" Text box.

- 10. Click the **Save** button to keep your changes. The **Save Successful** popup appears.
- 11. Click the **Ok** button in the **Save Successful** popup.
- 12. Click the Close button to exit.

## **Application Setup subtab**

- 1. Click **Customize > System** from the main menu. The **System Configuration** dialog box appears.
- 2. Click the **General Settings** tab.
- 3. Click the **Application Setup** sub tab.

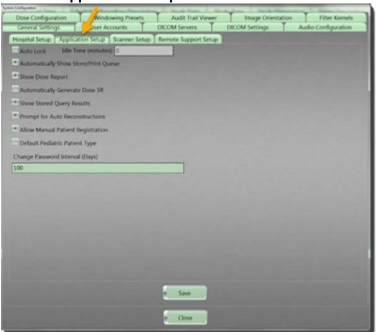


Figure 78: General Settings > Application Setup subtab

4. Click and/or enter information for the following options that apply:

Auto Lock	The <b>Auto Lock</b> feature can be used to automatically lock the workstation screen after a user defined <b>Idle Time</b> is reached.
Automatically Show Store/Print Queue	Allows the user to automatically display the <b>Store/Print Queue</b> status when images are set to archive to a network device.
Show Dose Report	Displays the <b>Dose Report</b> on the screen when the <b>Finalize</b> button is clicked.

	A dose report will not be generated until
	the operator clicks the <b>Finalize</b> button
	on the <b>Acquisition</b> tab.
Automatically Congrets	Generates a <b>Dose SR</b> (Structured Report)
Automatically Generate	along with the dose report when the
Dose SR	Finalize button is clicked.
Show Stored Query Besults	Displays the <b>Stored Results</b> at the
Show Stored Query Results	bottom of Patient Registration.
	Allows the user to automatically start
Prompt for Auto	any additional reconstructions added to
Reconstruction	a scan series after the exam is
	completed.
Allow Manual Patient	Allows the user to manually register a
Registration	patient.
Default Pediatric Patient	If selected, the <b>Protocol Manager</b> will
Туре	default to the pediatric protocols.
Change Password Interval	Sets the number of days before a
(Days)	password change is required.

- 5. Click the **Save** button to keep your changes.
- 6. Click the **Close** button to exit.

# **Scanner Setup subtab**

- 1. Click **Customize > System** from the main menu. The **System Configuration** dialog box appears.
- 2. Click the **General Settings** tab.
- 3. Click the **Scanner Setup** sub tab.

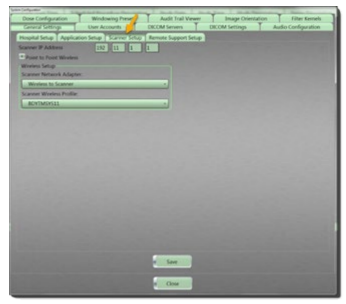


Figure 79: General Settings > Scanner Setup subtab

4. Click and/or enter information for the following options that apply:

-	9 1 11 7
Scanner IP address	Sets the scanner's IP address in the field(s).
	Sets up wireless information regarding the connection from the workstation to the
	scanner.
Point to Point Wireless	For <b>Scanner Network Adapter</b> , enter the
	adaptor, for example, Wireless to Scanner.
	For <b>Scanner Wireless Profile</b> , enter the
	wireless identifier in the field.

- 5. Click the **Save** button to keep your changes.
- 6. Click the Close button to exit.

## **Remote Support Setup subtab**

- 1. Click **Customize > System** from the main menu. The **System Configuration** dialog box appears.
- 2. Click the **General Settings** tab.
- 3. Click the **Remote Support Setup** subtab.

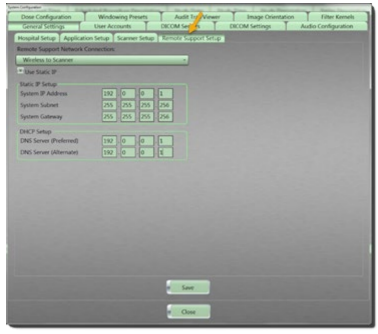


Figure 80: General Settings > Remote Support Setup subtab

- 4. Click the **Remote Support Network Connection** dropdown to select one of the following network connections:
  - Wireless to Scanner
  - Wired to Network
- 5. Click the **Use Static IP** option to enter the **Static IP Setup** data for the following:
  - System IP Address
  - System Subnet
  - System Gateway
- 6. Enter the **DHCP Setup** to enter IPs for the following:
  - DNS Server (Preferred)
  - DNS Server (Alternate)
- 7. Click the **Save** button to keep your changes.
- 8. Click the **Close** button to exit.

#### **Managing DICOM servers**

**Digital Imaging Communication in Medicine** is the definition of the acronym **DICOM**. **DICOM** servers are used to export images from the scanner. The **System Configuration > DICOM Servers** tab allows a user with administrative rights to access all the **DICOM** devices connected to the scanner.

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

**Note** You must have administrative privileges and be logged in as an administrator to access and modify DICOM servers.

Incorrect changes to the DICOM servers may make the system inoperative.

**DICOM** servers are set up by the **field-service engineer** and the appropriate IT person at the hospital.

- Click Customize > System from the main menu.
   The System Configuration dialog box appears.
- 2. Click the **DICOM Servers** tab.



Figure 81: DICOM Servers tab

#### The following tabs appear:

	Lists existing servers based on type:
	Store and Worklist
	Store: Identifies a storage server.
	Worklist: Identifies servers in a database you can
Servers	query from.
	Also displays server details and options, with
	controls for saving, updating, deleting, and echoing
	servers.
PACS List	Displays a list of <b>PACS</b> by <b>Server Name</b> , <b>Type</b> , and <b>In</b>
	<b>List</b> – to send to by default.
Options	Displays controls for PACS Options and HIS/RIS
	Options.



Figure 82: DICOM Servers tabs

3. Go to the following sections to assign specific actions to the **DICOM** server.

# Assigning a server as a store or worklist server

- 1. Click **Customize > System** from the main menu. The **System Configuration** dialog box appears.
- 2. Click the **DICOM Servers** tab. The **Servers** tab is active.



Figure 83: DICOM Servers > Servers tabs

3. Click one of the following options:

Store	A storage server, typically a <b>PACS</b> server that archives images and patient information. The images and data can later be imported from the server to another system or the same system. It is partly a backup and partly a waypoint for transferring data from the system the scans were acquired on to another system for viewing.
Worklist	A database of patient information that can be queried to generate a list of patients based on name, modality, procedure date, and other variables. These patients are usually hosted on a server within the facility and the information can be imported from the server when trying to acquire all a patient's information before a procedure is run. This eliminates the need to manually enter patient information at the time of the procedure.

- 4. Under **Details**, enter the server name in the **Server Name** text box.
- 5. For the **HIS/RIS** and **PACS Network Connection**, click the dropdown to identify the following:
  - Wireless to Scanner
  - Wired to Network
- 6. Click the **Point to Point Wireless** check box if it applies.

- 7. Click the Wireless Profile dropdown to select the appropriate profile.
- 8. In the **Server AE Title** text box, enter the server AE title.
- 9. In the **Device AE Title** text box, enter the BodyTom Elite AE title.
- 10. In the **Server IP Address** text boxes, enter the server IP address.
- 11. In the **Port** text box, enter the port identifier.
- 12. In the **Connection Timeout** text box, enter the number of seconds before the connection timeout is activated.
- 13. Click the **Use Static IP** option to identify the following:
  - System IP Address
  - System Subnet
  - System Gateway
- 14. To set the server as a default server, click the **Default Server** check box.
- 15. To set the server as the default surgical navigation server, click the **Default Navigation** check box.
- 16. To enable the storage commitment, click the Enable Storage Commit check box to send a message back to system that confirms the storage was successful; it is an extra confirmation from PACS that the images were received.
- 17. To gather responses, click the **Listen for Responses** check box. For every image that is sent, the system will wait for acknowledgement before sending the next image.
- 18. To gather 12-bit images, click the **12-bit Images** check box.
- 19. Click the **Save** button, to keep your work.



Figure 84: Action Succeeded popup message – Server saved The new server should appear in the list box to the left.

- 20. Click the **Ok** button.
- 21. Click the **Close** button to exit.

## Modifying a server

- 1. Click **Customize > System** from the main menu. The **System Configuration** dialog box appears.
- 2. Click the **DICOM Servers** tab.
- 3. Click server type: **Store** or **Worklist**.
- 4. Select a server from the list.
- 5. Modify the parameters.

  See "Assigning a server as a store or worklist server" on page 137.
- 6. When all your changes are made, click the **Update** button.

  A message appears that explains the update was successful and includes the update(s).



Figure 85: Action Succeeded popup message – Server updated

- 7. Click the **Ok** button.
- 8. Click the **Close** button to exit.

#### **Echoing a server**

- Click Customize > System from the main menu.
   The System Configuration dialog box appears.
- 2. Click the **DICOM Servers** tab.
- 3. Click a server type: Store or Worklist.
- 4. Select the server to echo from the list.
- Click the **Echo** button.The status of the server appears.



Figure 86: Echo Successful and Echo Failed popups

- 6. Click the **Ok** button.

  If the echo was unsuccessful, determine why and repeat step 5 until you are successful.
- 7. Click the **Close** button to exit.

## **Deleting a server**

- 1. Click **Customize > System** from the main menu. The **System Configuration** dialog box appears.
- 2. Click the **DICOM Servers** tab.
- 3. Click a server type: **Store** or **Worklist**.
- 4. When the server you want to remove is available, click the server from this list.

- Click the **Delete** button.
   The server is removed from the list; the **Action Succeeded** popup appears.
- 6. Click the **Ok** button.
- 7. Click the **Close** button to exit.

## Moving a server up and down the server list

- 1. Click **Customize > System** from the main menu. The **System Configuration** dialog box appears.
- 2. Click the **DICOM Servers** tab.
- 3. Click a server type: **Store** or **Worklist**.
- 4. Select the server to move up or down the list.
- 5. Click the **Up** arrow to move the server up the list; click the **Down** arrow to move the server down the list.



Figure 87: Up and Down arrows to move up and down server list

6. Click the **Save** button to save the server order.



Figure 88: Save button

The **Action Succeeded** popup appears.

- 7. Click the **Ok** button.
- 8. Click the **Close** button to exit.

# Saving DICOM servers to a PACS list

- 1. Click **Customize > System** from the main menu. The **System Configuration** dialog box appears.
- 2. Click the **DICOM Servers** tab.
- 3. Click the **PACS** List tab to view available servers.

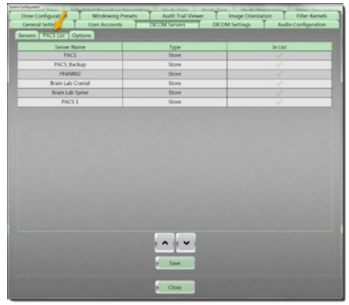


Figure 89: DICOM Servers > PACS List tab

- 4. Double-click the light-gray checkmark under In List. Each checkmark adds the server to the PACS listing. The checkmark turns green when active.
- 5. Click the **Save** button.

The PACS List Saved popup appears.



Figure 90: PACS List Saved popup message – PACS saved

- 6. Click the **Ok** button.
- 7. Click the **Close** button to exit.

## **Selecting PACS options**

1. Click **Customize > System** from the main menu. The **System Configuration** dialog box appears.

- 2. Click the **DICOM Servers** tab.
- 3. Click the **Options** tab.

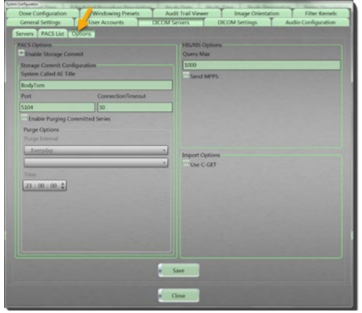


Figure 91: DICOM Servers > Options tab

- 4. Under **PACS Options**, click the **Enable Storage Commit** to verify that patient information and data archived to the **PACS** server was received.
  - If **Enable Storage Commit** is not selected, it is assumed and accepted that the data was received.
  - If **Enable Storage Commit** is selected, the workstation sends a request to the PACS server to verify that the data was received.
- 5. If the **Enable Storage Commit** check box is selected, perform the following:
  - Enter the appropriate title in the System Called AE Title text box.
  - Enter the port identifier in the Port text box.
  - Enter the number of seconds before a connection timeout in the Connection Timeout (secs) text box.
  - Enter 30 seconds or 60 seconds.
- 6. Under **Purge Options**, make selections based on the understanding that any studies archived to the **PACS** server are deleted from the workstation on a regular basis, depending on the selected interval; identify the following:

Purge Interval	Select one of the following from the dropdown:  Everyday, Weekly, or Monthly. When you select the first Purge Interval dropdown and select Weekly or Monthly, the inactive dropdown is active to let you select the day of the week or the
	first of the month.
Time	Changes the hour, minute, and second interval; use the buttons to increase those time elements.  23:00:00 Figure 92: Time (increase and decrease time) arrows

- 7. Under the **HIS/RIS Options**, enter the maximum number of results sent back from a query worklist in the **Query Max** text box.

  There is no maximum limit.
- 8. Click the **Send MPPS** check box to apply a service that allows a modality to better coordinate with image storage servers by giving the server a list of objects to send before or while sending such objects.
- Under Import Options, click the Use C-GET check box to pull information from a PACS server when importing from the server (as opposed to archiving to it).

The administrator sets this to pull from **PACS** from anywhere, so the machine does not have to be set up as a reliable destination on the **PACS** machine. **PACS**, typically needs to equate a computer's IP address with an AE title; however, **C-GET** accepts that the calling IP is a legitimate device.

The NeuroLogica BodyTom Elite scanner automatically uses **C-Move** when importing from **PACS**. If the operator wants to use **C-GET** instead, the user can select **C-Get**.

10. Click the **Save** button.

The **PACS List Saved** popup appears.



Figure 93: PACS List Saved popup

- 11. Click the Ok button.
- 12. Click the **Close** button to exit.

# **Assigning DICOM settings**

**DICOM** settings include many kinds of settings. The administrator can add or remove optional information to be displayed using actions described in this section.

See **DICOM** standards on the **NEMA.org** website for a full list and description of **DICOM** tags.

Note You must have administrative privileges and be logged in as an administrator to access and modify DICOM settings.

Incorrect changes to the DICOM settings may make the system inoperative.

- 1. Click **Customize > System** from the main menu. The **System Configuration** dialog box appears.
- 2. Click the **DICOM Settings** tab.



Figure 94: DICOM Settings tabs (six)

3. Click the **HIS/RIS Query** tab to select the types of **HIS/RIS** query results the user will see when performing a **HIS/RIS** query. See "Selecting PACS options" and go to step 6, for more information.

Note Even numbered DICOM tags are public DICOM tags as per the DICOM standard. Odd numbered DICOM tags are vendor specific.

Green checkmarks are optional **DICOM** tags and orange checkmarks are required per the **DICOM** standard and cannot be modified.

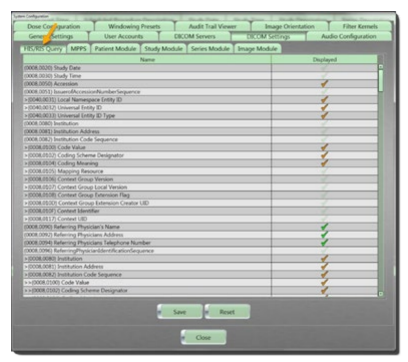


Figure 95: DICOM Settings > HIS/RIS Query

4. Click the **Modality Performed Procedure Step (MPPS)** tab to select the types of **MPPS** information the user will see.



Figure 96: DICOM Settings > MPPS

5. Click the **Patient Module** tab to select the types of **Patient Module** information the user will see.



Figure 97: DICOM Settings > Patient Module

6. Click the **Study Module** tab to select the types of **Study Module** information the user will see.

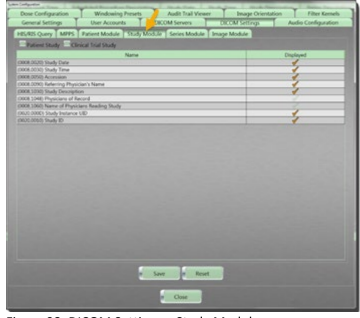


Figure 98: DICOM Settings > Study Module

7. Click the **Series Module** tab to select the types of **Series Module** information the user will see.

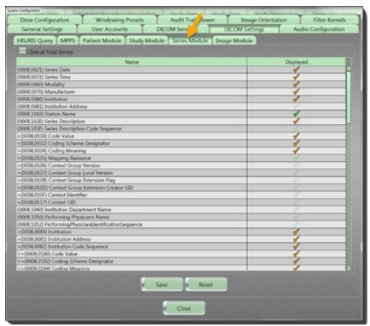


Figure 99: DICOM Settings > Series Module

8. Click the **Image Module** tab to select the types of **Image Module** information the user will see.

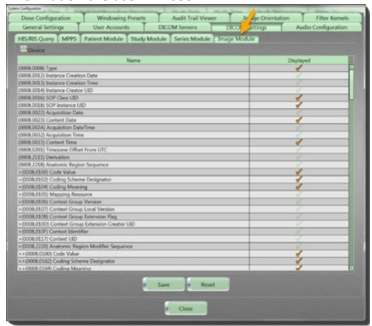


Figure 100: DICOM Settings > Image Module

- 9. Click the **Save** button to save your changes.
- 10. Click the Close button to exit.

## Assigning audio configuration

Default audio files are installed on the workstation. Audio files can be attached to protocols and sent to the scanner. Each audio file has an indication if it has been sent to the scanner.

**Note** You must have administrative privileges and be logged in as an administrator to access and modify audio configurations.

Incorrect changes to the audio configurations may make the system inoperative.

### Finding and listening to audio files

- Click Customize > System from the main menu.
   The System Configuration dialog box appears.
- 2. Click the Audio Configuration tab.

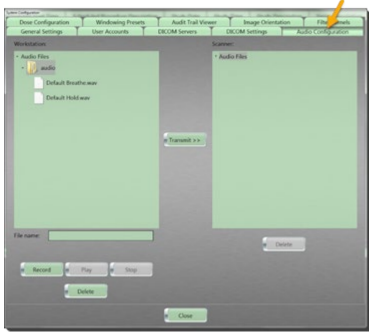


Figure 101: Audio Configuration tab

3. Review the audio files that exist on the Workstation.

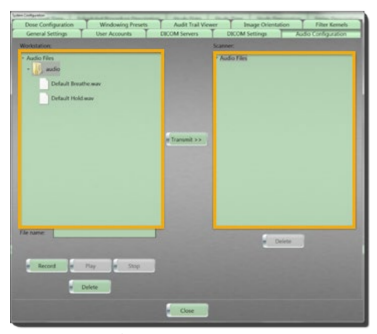


Figure 102: Audio files list

- 4. To listen to an audio file, select the name from the workstation list and click the Play button.
- 5. To exit the Audio Configuration, click the **Close** button.

# Recording and saving an audio file

- 1. Click **Customize > System** from the main menu. The **System Configuration** dialog box appears.
- 2. Click the **Audio Configuration** tab.

  The available audio files appear under **Workstation**.

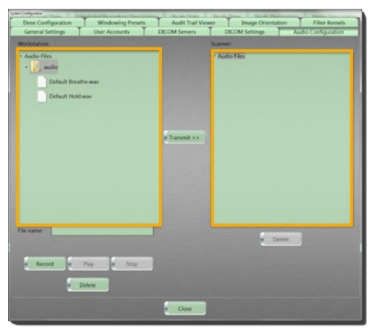


Figure 103: Audio files list

- 3. Enter the name of your new audio file in the File name text box.
- 4. Click the Record button.
- 5. Record your audio file.
- 6. Press the **Stop** button.
- 7. To review, highlight your new recording and press the **Play** button.



Figure 104: New audio file

- 8. When you like your recording, highlight the file and press the **Transmit** button to copy the file into the audio folder for your scanner protocols.
- 9. Click the Close button to exit.

### Transmitting an audio file

- Click Customize > System from the main menu.
   The System Configuration dialog box appears.
- Click the Audio Configuration tab.The available audio files appear under Workstation.
- 3. Select the audio files under **Workstation** audio files to transfer to the scanner.
- 4. Click the Transmit button.

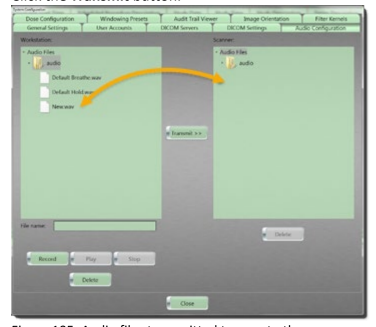


Figure 105: Audio files transmitted to save to the scanner

5. Click the Close button to exit.

# Deleting an audio file

- Click Customize > System from the main menu.
   The System Configuration dialog box appears.
- 2. Click the Audio Configuration tab.

The available audio files appear under **Workstation**.

- 3. Select the audio file you want to delete from the list.
- 4. Click the **Delete** button.
- 5. Click the **Close** button to exit.

# Assigning dose report

The **dose report** is created at the end of the scan and can be customized to include **DICOM** specific tags.

**Note** You must have administrative privileges and be logged in as an administrator to access and modify dose report settings.

- 1. Click **Customize > System** from the main menu. The **System Configuration** dialog box appears.
- 2. Click the **Dose Configuration** tab.
- 3. Click the **Dose Report** tab.

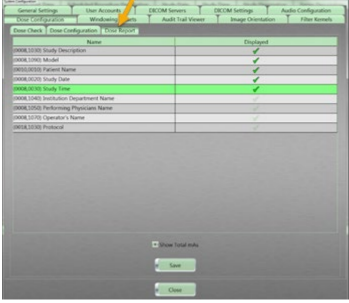


Figure 106: Dose Report tab

4. Select the **DICOM** tags you want to see in the **dose report**. A **dose report** is generated after the exam is finalized; the black area includes dose report information like the following.



Figure 107: Generated dose report

- 5. Click the **Save** button.
- 6. Click the **Close** button to exit.

## Applying dose configuration

**Note** You must have administrative privileges and be logged in as an administrator to access and modify dose configurations.

Incorrect changes to dose configuration settings 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**.

Dose Notification	Notifies the user when the planned CTDI <sub>vol</sub> and/or DLP value of a single series will exceed the defined
	value.
Dose Alert	Notifies the user when the planned CTDI <sub>vol</sub> and/or DLP value from the combination of all planned series will exceed the defined value set in <b>System Configuration</b> . Dose Alerts represent a value which would be well above an institutions established CTDI/DLP range to the given
	examination and warrant a more stringent review and consideration before proceeding.

### **Setting Dose Check**

See Appendix A on page 389 for information on protocols, CTDI<sub>vol</sub>, and DLP.

- 1. Click **Customize > System** from the main menu. The **System Configuration** dialog box appears.
- 2. Click the **Dose Configuration** tab.
- 3. Click the Dose Check tab.



Figure 108: Dose Configuration > Dose Check

4. Click one of the following **Dose Check Type** options:

Doso	Notifies the user when a pre-defined
Dose Notification	CTDI <sub>vol</sub> or DLP value will be exceeded on
	a series-by-series basis.
Dose Alert	Notifies the user when a pre-defined
	CTDI <sub>vol</sub> or DLP value will be exceeded
	from a combination of all planned series
	or scans.

Note The default Dose Alerts which are set at 1000mGy CTDI and 2000mGy\*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 should be done by qualified medical personnel.

5. Click one from the following **Scan Type** options.

All	Identifies <i>all</i> scan types.
Axial	Identifies only <b>Axial</b> scan types.
Helical	Identifies only <b>Helical</b> scan types.
Dynamic	Identifies only <b>Dynamic</b> scan types.

- 6. Define the **Dose Limit** by entering the following:
  - Enter the CTDI<sub>vol</sub> (mGy) value in the text box.
  - Enter the DLP (mGy.cm) value in the text box.
- 7. Click the Save button.

The Save Successful popup appears.



Figure 109: Save Successful popup – Dose Check successfully saved

- 8. Click the **Ok** button.
- 9. Click the Close button to exit.

# Assigning Dose Configuration to a patient protocol

**Dose Configuration** limits are used to prevent users from selecting kV or mA values that are not appropriate for the given patient types, such as pediatrics etc.

- Click Customize > System from the main menu.
   The System Configuration dialog box appears.
- 2. Click the **Dose Configuration** tab.
- 3. Click the **Dose Configuration** subtab.

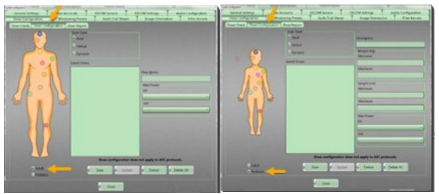


Figure 110: Dose Configuration > Dose Configuration for adult and pediatric

4. Click one of the following:

Adult	Selecting Adult shows the pre-defined adult
	protocols, stored by anatomical area.
Pediatric	Selecting Pediatric shows the pre-defined pediatric
	protocols, stored by anatomical area.
Trauma	The <b>Trauma</b> orb can be used to store protocols
	commonly used for emergency scans.

5. Click the colored orb that marks the anatomical region to apply the dose to.

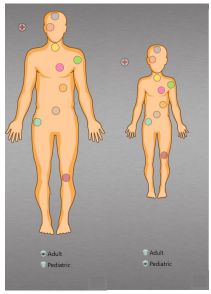


Figure 111: Anatomical orbs

6. Click a scan type from the following list:

Axial	Identifies only <b>Axial</b> scan types.
Helical	Identifies only <b>Helical</b> scan types.
Dynamic	Identifies only <b>Dynamic</b> scan types.

- 7. Enter a description for the **Dose Configuration** in the **Description** text box.
- 8. For **Pediatrics** enter the **Minimum** and **Maximum Weight** and **Length** information.
- 9. Under the **Max Power** settings, click the **kV** dropdown and select the maximum allowed kV.
- 10. Click the **mA** dropdown and select the maximum allowed mA.

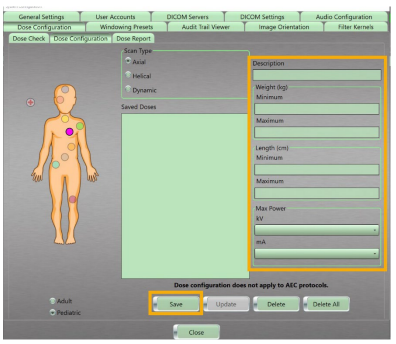


Figure 112: Pediatric Dose Configuration Parameters

11. Click the **Save** button to save your work.

If the level overlaps an existing level, you are prompted to adjust.



Figure 113: Invalid Parameter popup message – Dose setting kV already exists

If the save is successful, the **Save Successful** popup appears.



Figure 114: Save Successful popup message – Maximum dose saved

- 12. Click the **Ok** button.
- 13. Under the **Saved Doses** list box, check that your dose configuration appears, if so, go to the next step.
- 14. Click the Close button to exit.

### **Updating saved dose**

#### To modify a saved **Dose Configuration**.

See "Setting Dose Check" page 158 and/or "Assigning Dose Configuration to a patient protocol" on page 159.

- 1. Click **Customize > System** from the main menu. The **System Configuration** dialog box appears.
- 2. Click the **Dose Configuration** tab.
- 3. Select the desired **Saved Dose** from the **Saved Doses** list.

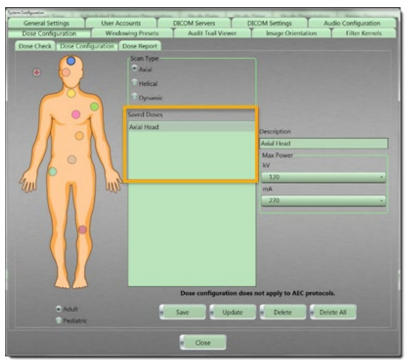


Figure 115: Saved Doses List

- 4. Modifying values causes the **Update** button to become active.
- 5. Click the **Update** button.



Figure 116: Save Successful popup message – Maximum dose saved

- 6. Click the **Ok** button.
- 7. Click the **Save** button to save your work.

**Note** If the level overlaps an existing level, you are prompted to adjust.

8. Click the **Close** button to exit.

### Deleting a saved dose limit

To remove a saved **Dose Configuration**.

- 1. Click **Customize > System** from the main menu. The **System Configuration** dialog box appears.
- 2. Click the Dose Configuration tab.

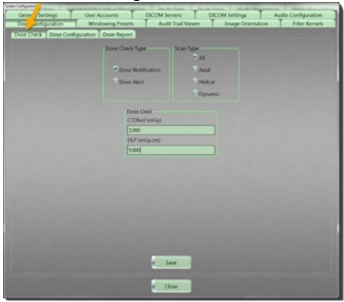


Figure 117: Dose Configuration > Dose Check tab

- 3. Click an already-saved dose, under the **Saved Doses** list.
- 4. Perform one of the following:
  - To delete a saved dose from the Saved Doses list and clear the restriction, select the dose, and click the Delete button.
  - To delete all the saved doses in the Saved Doses list and clear all restrictions saved, click the Delete All button, which returns all settings for that selection to the maximum scanner default.

Note If there are no saved doses or limits, the operator will be able to scan using the maximum 140kV and 300mA available on the scanner.

5. Click the **Save** button.

The save success message appears and, because the **CTDIvol** (**mGy**) and **DLP** (**mGy.cm**) are empty, there is no longer a limit applied.

The Save Successful popup appears.



Figure 118: Save popup message – Maximum dose saved

- 6. Click the Ok button.
- 7. Click the **Close** button to exit.

#### **Applying Windowing Presets**

Windowing presets allow you to define window width and center presets for specific anatomical locations as well as specific reconstruction kernel presets. An Administrative User can delete or update the default Window Presets as well as create new Windowing Presets.

**Note** You must have administrative privileges and be logged in as an administrator to access and modify the windowing presets.

### **Editing kernel presets**

**Note** Kernel presets are pre-installed in the system; kernel presets can be set and modified.

- Click Customize > System from the main menu.
   The System Configuration dialog box appears.
- 2. Click the Windowing Presets tab.

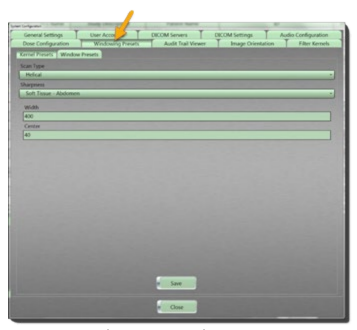


Figure 119: Windowing Preset tab

3. Click the Kernel Presets tab.

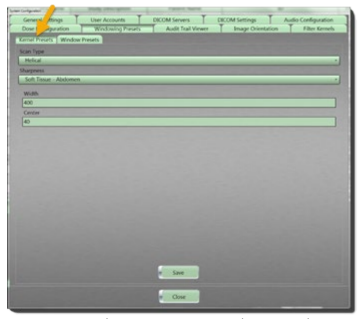


Figure 120: Windowing Presets > Kernel Presets tab

- 4. Click the **Scan Type** dropdown to select one of the following scan types:
  - Axial
  - Helical

5. Click the **Sharpness** dropdown to select a sharpness from the list. **Sharpness** is the reconstruction algorithm, and the available **Sharpness** values are based on the scan type.

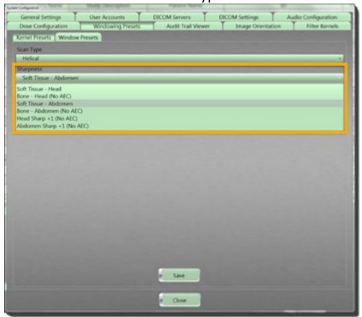


Figure 121: Sharpness dropdown

- 6. Enter the Window Width in the Width text box. Window Width describes the range of Hounsfield units, or shades of gray, displayed across the image. The Window Width controls the contrast of the image. Low Hounsfield numbers below the range are displayed as black, while High Hounsfield numbers above the range are displayed as white.
- Enter the Window Center in the Center text box.
   Window Center describes the Hounsfield number in the center of the Window Width. Window Center controls the brightness or density of the image.
- 8. Click the **Save** button to save your work. The **Action Succeeded** popup appears.



Figure 122: Action Succeeded popup message – Preset saved

- 9. Click the **Ok** button.
- 10. Click the **Close** button to exit.

## **Setting Window Presets**

Window presets allow you to define **Window Width** and **Window Center** presets for specific anatomical locations, such as bone, brain, lung, and soft tissue.

- 1. Click **Customize > System** from the main menu. The **System Configuration** dialog box appears.
- 2. Click the Windowing Presets tab.
- 3. Click the Window Presets subtab.

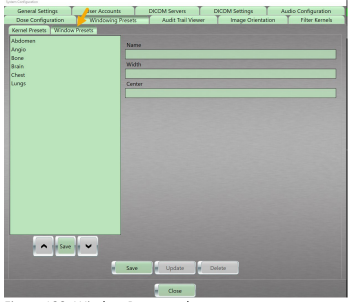


Figure 123: Window Presets tab

General Settings

User Accounts

Dicom Servers

Dicom Settings

Audio Configuration

Windowing Presets

Audit Trail Viewer

Image Orientation

Filter Kernels

Kernel Presets

Abdomen

Angio

Bone

Brain

Chest

Lungs

Center

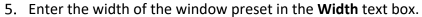
Save

Update

Delete

4. Enter the name of the window preset in the **Name** text box.

Figure 124: Window Presets > Name



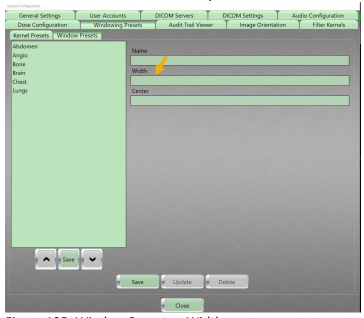


Figure 125: Window Presets > Width

6. Enter the center for the window preset in the **Center** text box.



Figure 126: Window Presets > Center

7. Click the **Save** button to save your work. The **Action Succeeded** popup appears.



Figure 127: Action Succeeded popup message – Preset saved

- 8. Click the **Ok** button.
- 9. Click the Close button to exit.

# Editing a window preset

- Click Customize > System from the main menu.
   The System Configuration dialog box appears.
- 2. Click the Windowing Presets tab.

- 3. Click the **Window Presets** subtab.
- 4. Click a preset that exists in the **Window Presets** listing.

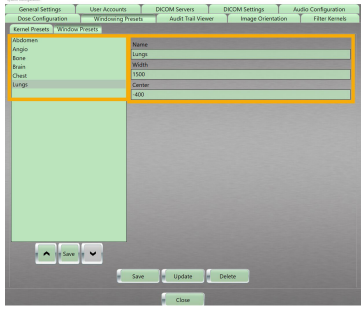


Figure 128: Listing update

- 5. To edit the preset, make your changes in the **Name**, **Width**, and/or **Center** text boxes.
- 6. Click the **Save** button to save your changes. The **Action Succeeded** popup appears.



Figure 129: Action Succeeded popup message – Preset saved

- 7. Click the **Ok** button.
- 8. Click the **Close** button to exit.

### **Deleting a preset**

- 1. Click **Customize > System** from the main menu. The **System Configuration** dialog box appears.
- 2. Click the **Windowing Presets** tab.
- 3. Select the preset.
- 4. Click the **Delete** button.



Figure 130: Action Succeeded popup message – Preset deleted

- 5. Click the **Ok** button.
- 6. Click the Save button to exit.

## **Setting up the Audit Trail Viewer**

The **Audit Trail Viewer** gives a user with administrative access the ability to view all activities performed by anyone logged into the system. This includes changes to protocols, deletion of images, as well as acknowledgement of alerts etc.

**Note** You must have administrative privileges and be logged in as an administrator to access the Audit Trail Viewer.

- 1. Click **Customize > System** from the main menu. The **System Configuration** dialog box appears.
- 2. Click the **Audit Trail Viewer** tab.

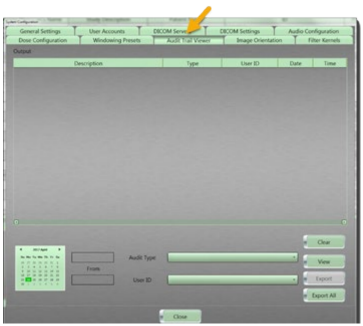


Figure 131: Audit Trail Viewer tab

- 3. From the **Audit Trail Viewer** tab, use the calendar to select a date range to view.
  - To select a single date, position the mouse pointer in the top box and click the date on the calendar to find audits for that date.
  - To select a date range, click the desired start date on the calendar.
     This will automatically populate the top box of the From range. Click the desired end date on the calendar, which automatically fills the bottom box of the From range.



Figure 132: Adding a date or a date span

4. Click the **Audit Type** dropdown to select the type of audit you are searching.

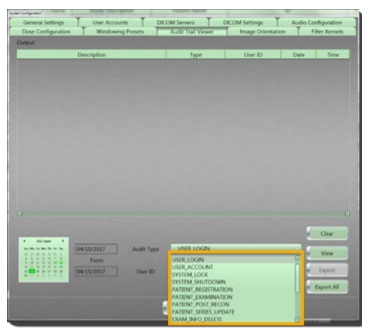


Figure 133: Audit Trail Viewer > Audit Type dropdown

5. From the **User ID** dropdown, click the type of user to track.

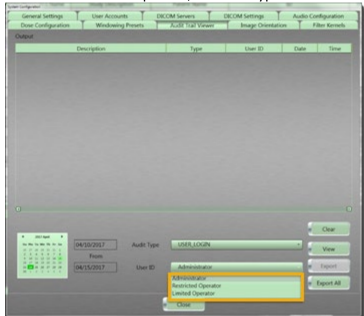


Figure 134: Audit Trail Viewer > User ID dropdown

6. Click the **View** button to see the result of audits that met your criteria.

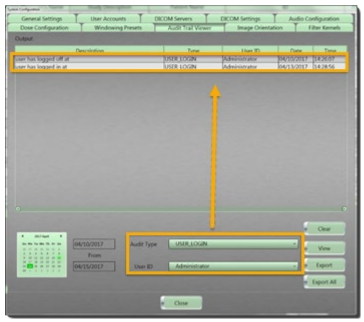


Figure 135: Audit results

- 7. Perform one of the following:
  - Click the Clear button to remove the audit results.
  - Click the **Export** button to export the audit result that you selected to the audit backup file on the system.
  - Click the Export All button to export the audit results to the audit backup file on the system.
- 8. Click the Close button to exit.

#### Setting image orientation

NeuroLogica describes patient orientation as if the viewer were looking towards the front of the gantry. 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.

**Note** You must have administrative privileges and be logged in as an administrator to modify image orientation settings.

Changes to image orientation settings will modify the displayed orientation markers on the images.

1. Click **Customize > System** from the main menu.

The **System Configuration** dialog box appears.

2. Click the **Image Orientation** tab.

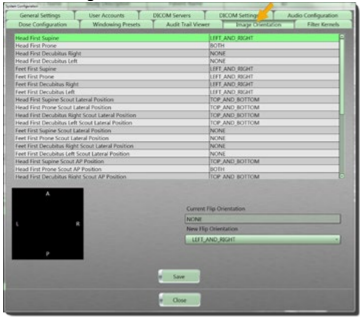


Figure 136: Image Orientation tab

The top half of the **Image Orientation** screen shows the available patient orientations. The black image orientation square represents the viewing area and shows four different orientation markers: A = anterior, L = left, P = posterior and R = right. If you do not see letters in the image orientation box, select an image orientation from the list.

- Select the appropriate orientation from the list.
   For example, select, Head First Supine. In the figure below, the highlighted selection shows the current orientation in the Current Flip Orientation field, which is not changeable; however, the New Flip Orientation lets you change the orientation.
- 4. Click the **New Flip Orientation** dropdown to select one of the following new-flip orientations:
  - NONE
  - LEFT\_AND\_RIGHT
  - TOP AND BOTTOM
  - BOTH

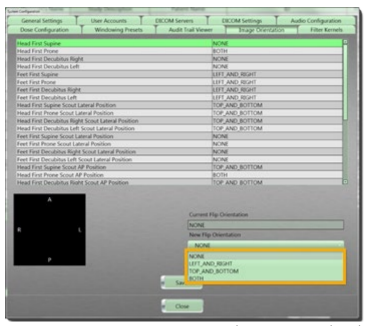


Figure 137: Image Orientation > New Flip Orientation dropdown

- 5. Click the **Save** button to save changes.
- 6. The **Settings Saved** popup appears.



Figure 138: Settings Saved popup message – Image orientation settings saved

- 7. Click the **Ok** button.
- 8. Click the Close button to exit.

## **Setting Filter Kernels**

Filter kernels allow you to activate custom kernel options for both **Axial** and **Helical** scans to control the sharpness and smoothness of the images.

- 1. Click **Customize > System** from the main menu. The **System Configuration** dialog box appears.
- 2. Click the Filter Kernels tab.

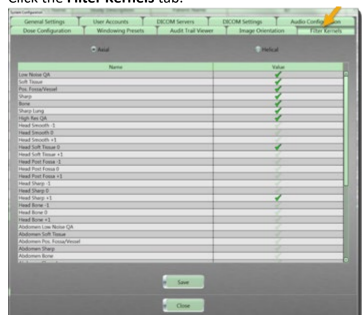


Figure 139: Filter Kernels tab

- 3. Perform one of the following:
  - To add new Axial kernels, select the Axial radio button, then doubleclick the Value cell next to the desired Axial kernel.

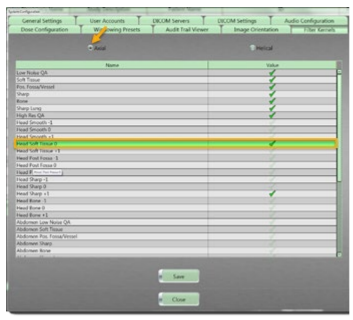


Figure 140: Selected Axial kernel

To add new Helical kernels, select the **Helical** radio button, then double-click the **Value** cell next to the desired Helical kernel.

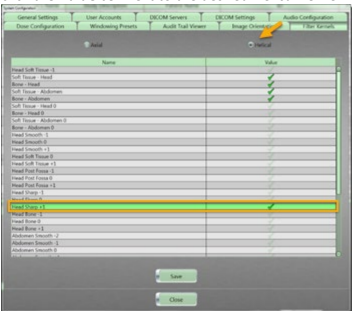


Figure 141: Selected Helical kernel

- Click the Save button.The Success popup appears.
- 5. Click the **Ok** button.
- 6. Click the **Close** button to exit.

## **User configuration**

User configuration allows users with either **Administrator** or **Limited Operator** access to change the password for their own account.

#### Updating your user account

Click Customize > User from the main menu.
 The User Configuration dialog box appears.

**Note** You must have administrative or limited operator privileges to access and modify user accounts.

Incorrect changes to user accounts may cause users to be unable to access the system.

The **Update Accounts** tab is the default selection.

- 2. Modify the following fields associated with your user account by entering relevant information:
  - Last Name
  - First Name
  - Password
  - Verify Password

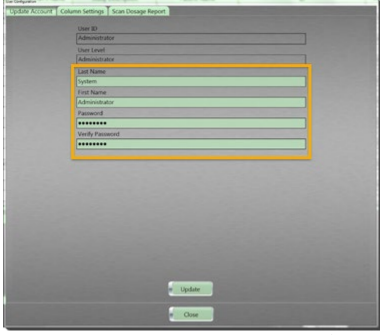


Figure 142: Last Name, First Name, Password, and Verify Password fields

3. Click the **Update** button.



4. The Update Succeeded popup appears.

Figure 143: Update Succeeded popup message – Account updated

- Click the **Ok** button.
- 6. Click the **Close** button to exit.

#### Applying column settings to HIS/RIS Query

Allows you to customize the columns of information that appear when viewing the **Hospital Information System (HIS)** and/or **Radiology Information Systems (RIS)** information that is queried.

All users can access user configuration and make changes to the column settings; however limited and restricted operators cannot make their changes to the column headings the default. Only users with administrative access can make column settings a default, using the **Make Default** option.

- Click Customize > User from the main menu.
   The User Configuration dialog box appears.
- 2. Click the Column Settings tab.
- 3. There may be no entries that appear, initially.

Note When an option is selected (for example HIS/RIS Query or Patient Browser), a table is created that lists the columns, along with a check box to indicate whether it will be displayed within the table. Required columns cannot be unchecked and are colored orange instead of the default green.

4. Click the **HIS/RIS Query** option.

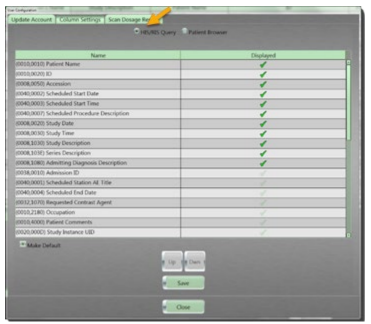


Figure 144: Column Settings dialog box with HIS/RIS Query option

5. Double click the Checkmark under **Displayed** column for the row you want to display or remove if it is already selected.

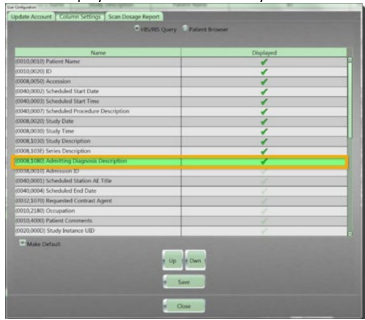


Figure 145: Column Settings with a selected query in HIS/RIS

- 6. Notice that the **Up** and **Dwn** buttons are active when a row is selected.
- 7. Click the **Up** or **Dwn** buttons to move the active selection up or down the list.

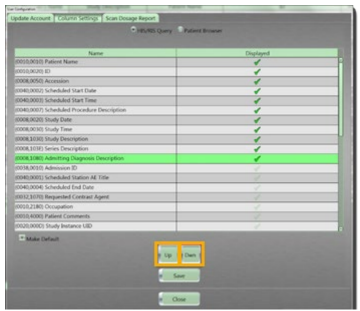


Figure 146: Column Settings with HIS/RIS Query option using Up and Dwn buttons

8. If you have administrative privileges, click the **Make Default** option to make the selected column display the default for all users.

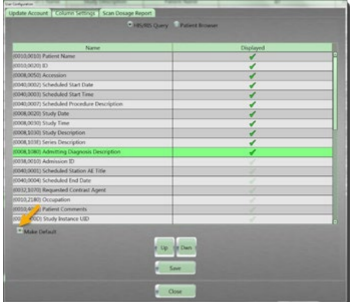


Figure 147: Make Default option

- 9. Click the **Save** button to keep changes.
- 10. Click the **Close** button to exit.

#### **Applying column settings to Patient Browser**

Allows you to configure the columns of information seen in the **Patient Browser.** 

- Click Customize > User from the main menu.
   The User Configuration dialog box appears.
- 2. Click the Column Settings tab.
- 3. Click the **Patient Browser** option.

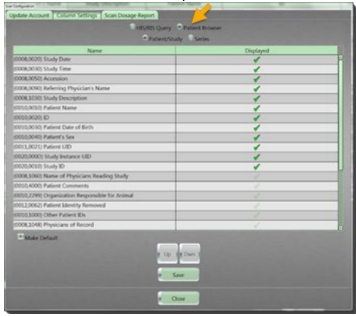


Figure 148: Column Settings with Patient Browser option

4. Click one of the following options:

	Information that appears on the top portion of the
Patient/Study	Patient Browser that defines patient specific
	information.
Series	Information that appears on the lower portion of the
	Patient Browser that defines series specific information.

- 5. Double click the Checkmark under **Displayed** column for the row you want to display or remove if it is already selected.
- 6. Notice that the **Up** and **Dwn** buttons are active when a row is selected.
- 7. Click the **Up** or **DWN** buttons to move the active selection up or down the list.

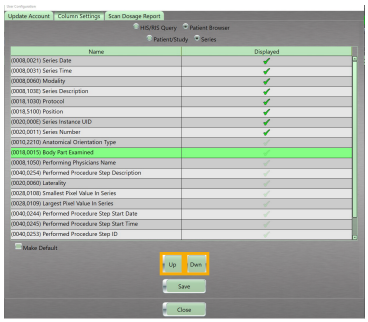


Figure 149: Column Settings with Patient Browser Series option – using Up and Dwn buttons

8. If you have administrative privileges, click the **Make Default** option to make the selected column display the default for all users.

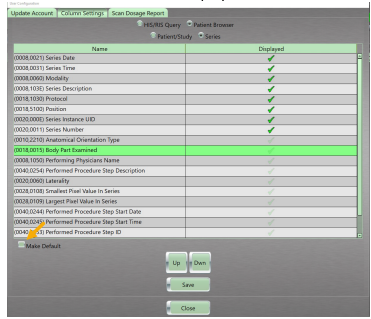


Figure 150: Make Default option

- 9. Click the **Save** button to keep changes.
- 10. Click the **Close** button to exit.

#### **Viewing Scan Dosage Report**

- 1. Click **Customize > User** from the main menu. The **User Configuration** dialog box appears.
- 2. Click the Scan Dosage Report tab.

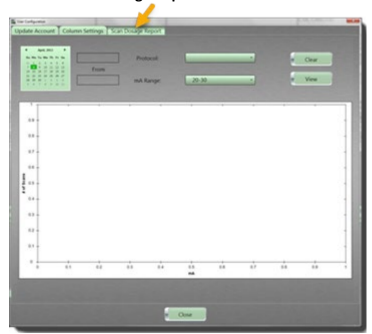


Figure 151: Scan Dosage Report tab

3. To view **Dosage Reports**, click the date or date range on the calendar.

Note If the date is left blank – all doses for all dates are retrieved.

4. Click the **Protocol** dropdown to select a protocol.

**Note** If the protocol is left blank – all doses for all protocols are retrieved.

Click the mA Range dropdown to select the mA range.
 The mA Range default is 20-30; it can be changed after data is retrieved.



Figure 152: Date, Protocol, and mA Range filled

6. Click the **View** button to display a graph showing dosages performed by the scanner using the selected filters.

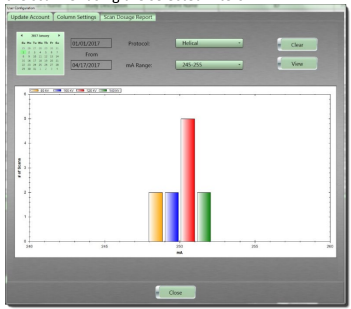


Figure 153: Scan Dosage Report results

**Note** If you adjust the mA range, the graph displays those ranges within the retrieved data.

7. To clear the filters selected, click the **Clear** button.

# Selecting a room for the BodyTom Elite

Selecting a room ensures that the correct calibration is loaded or used when the scanner has been calibrated in more than one location.

- 1. Click Customize > Select Room.
- 2. A list of the rooms available appear in the cascading menu.



Figure 154: Available rooms before moving the scanner

- 3. Click the room in which the scanner will be used.
- 4. Move scanner to the selected room.

# **Chapter 6** Protocol Manager

Note You must have administrative privileges and be logged in as an administrator, to access the Protocol Manager.

**Protocol Manager** allows a user with administrative privileges to create new protocols, modify existing protocols and delete protocols from the system. **Protocol Manager** provides three patient options: **Adult, Pediatric,** or **Trauma** • patient.

Table 27: Protocol Manager command buttons

Table 27: Protocol Manager command buttons	
Button	Action
New	Allows you to create a new protocol.
Build From	Allows you to create a new protocol from an existing protocol.
Edit	Allows you to modify protocols.
Delete	Deletes a saved protocol.
Import	Imports previously exported protocols into the workstation.
Export	Exports protocols to a media device.
Close	Closes the <b>Protocol Manager</b> dialog box.
	Moves a protocol up or down the ordered list.
Save	Saves the order of the protocol list.

Note Different patient sizes and attenuations require specific protocols where technical parameters have been adjusted according to the physical characteristics of the patient. It is recommended that you select pediatric protocols when scanning children, rather than selecting adult protocols and modifying technique factors.

### Creating a new protocol

1. Click **Tools >Protocol Manager** from the main menu. The **Protocol Manager** dialog appears.



Figure 155: Protocol Manager for adult and pediatric

2. Click one of the following:

Adult	To create and/or scan with adult scan protocols,
	which are stored by anatomical location.
Pediatric	To create and/or scan with pediatric scan protocols,
	which are stored by anatomical location.
Trauma	The <b>Trauma</b> orb can be used to store protocols
	commonly used for emergency scans.
• Adult • Pediatric	By selecting either an <b>Adult</b> or <b>Pediatric</b> patient, the
	corresponding list of saved protocols becomes
	available.

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.

This scanner comes with reference protocols for pediatric patients that vary according to age or weight. These protocols are meant to be a guide for routine scanning. An individual patient may require higher or lower doses than in the reference protocol to achieve the diagnostic goals for the patient's medical condition. These reference pediatric protocols were established as a reduction from the adult protocols to achieve the same image signal to noise. Currently there is no agreement among scientific and medical organizations as to what the proper protocol is for pediatric patients. It is the responsibility of the user to decide whether these protocols are adequate for achieving their diagnostic goals on a patient-by-patient basis.

The user can modify any of the existing protocols on the machine and/or create new protocols as deemed necessary. To create these protocols, Administrator privileges are required.

Total
Onet ACA
Onet A

3. Click the colored orb corresponding to the appropriate body part.

Figure 156: Adult and pediatric anatomical orbs, with Chest orb selected

Existing protocols in the selected Orb will appear in the **Protocol** list box as seen below. The **New** button will become active.

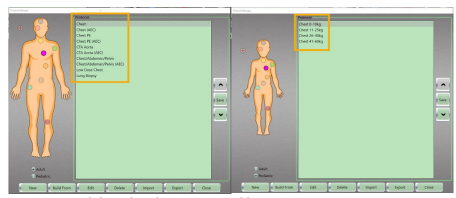


Figure 157: Adult and pediatric protocol lists

4. Click the **New** button to create a new protocol. The **New Protocol** dialog box appears.

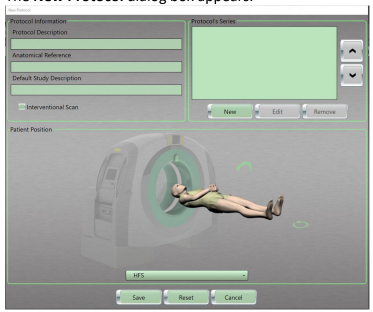


Figure 158: New Protocol dialog box

#### Enter **Protocol Information** in the text boxes:

Protocol Description	The name of the protocol as it will be displayed in protocol manager or when selecting a protocol to use for a scan.
Anatomical Reference	References the anatomy that will be scanned.
Default Study Description	This will appear as the <b>DICOM</b> image tag, this description also appears in <b>PACS</b> as a <b>Study Description DICOM</b> tag (0008,1010).

5. Under **Patient Position**, select one of the following scanning positions from the dropdown:

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

You can also move the rotating positional handles by hovering the mouse pointer over the handle and clicking to select a position.

- The arrows at the feet rotate the patient orientation from Head
   First to Feet First.
- The arrow above the patient rotates the patient orientation from Supine to Prone to Decubitus.

The **Patient Position** settings are identical whether it is for an adult, pediatric, or emergency patient.

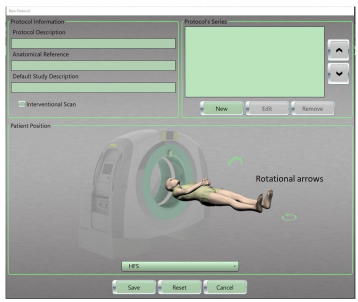


Figure 159: Patient position handles

6. Under **Protocol's Series**, click the **New** button. The **New Series** dialog box appears.

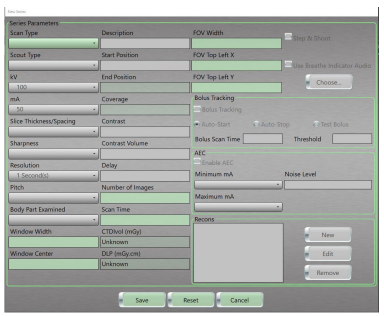


Figure 160: New Series dialog box

- 7. For **Scan Type**, select one of the following:
  - Axial
  - Helical
  - Dynamic
  - Reference
  - Scout

Note For Helical scanning, the exposed area is extended by at least ½ rotation to 1 full rotation at the start and end of the planned scan, based on the sharpness selected.

- 8. For **Scout Type**, select one of the following:
  - AP
  - PA
  - Lateral

**Scout Type** is not available for **Axial**, **Helical**, **Dynamic**, and **Reference** scan modes.

- 9. For **kV** (scan voltage), select one of the following:
  - 80 To set the scan kV to 80.
  - 100 To set the scan kV to 100.
  - 120 To set the scan kV to 120.
  - 140 To set the scan kV to 140.

**kV** is not selectable when using the **Dynamic** and **Reference** scan modes.

See "Identifying load factors" on page 235.

- 10. For mA (scan current), select the desired value from the dropdown. mA is not selectable when using the Reference scan mode. The available mA range is 30 to 300 with an increment of 5.
- 11. For **Slice Thickness/Spacing**, select the desired value from the dropdown.

**Slice Thickness/Spacing** is not available for **Scout** and **Reference** scan modes.

- 12. For **Sharpness**, select the image reconstruction kernel from the following list of kernels:
  - Low Noise QA Not for Clinical Use
  - Soft Tissue Available for Axial Scan Type only
  - Soft Tissue Head Available for Helical Scan Type only
  - Soft Tissue Abdomen Available for Helical Scan Type only
  - Pos. Fossa/Vessel Available for Axial Scan Type only
  - Sharp Available for Axial Scan Type only
  - Bone (No AEC) Available for Axial Scan Type only
  - Bone Head (No AEC) Available for Helical Scan Type only
  - Bone Abdomen (No AEC) Available for Helical Scan Type only
  - Sharp Lung (No AEC) Available for Axial Scan Type only
  - High-Res QA (No AEC) Not for Clinical Use

Sharpness is not selectable when using the **Reference** and **Scout** scan modes.

Note The Low Noise QA and High-Res QA (No AEC) options should not be used for clinical scanning.

- 13. For **Resolution**, which also refers to scan time, select one of the following options:
  - 1 Second(s)
  - 2 Second(s)

**Resolution** in only available for **Axial Scan Types.** 

- 14. For **Pitch**, which describes how fast the scanner is moving during one rotation of the x-ray tube, select one of the following options:
  - **0.4** where the scanner will move 16mm per second.
  - **0.8** where the scanner will move 32mm per second.

Pitch is only available for Helical Scan Types.

- 15. For **Body Part Examined**, select the appropriate Body Part from the drop-down menu.
- 16. For **Window Width**, enter the range of CT numbers that are distributed over the viewable gray scale of the display device or film.
- 17. For **Window Center**, enter the CT number in the center of the viewable gray scale.
- 18. For **Description**, enter the desired study description.
- 19. For **Start Position**, enter the start scan position.
- 20. For **End Position**, enter the end scan position.
- 21. **Coverage** is a calculated value that automatically fills based on the **Start** and **End** position values.
- 22. For **Contrast**, enter the type of contrast given for example
- 23. For **Contrast Volume**, enter the amount of the contrast given. **Contrast** is not available for **Reference** and **Scout** scan modes.
- 24. For **Delay**, enter the delay time that will occur after clicking the **START** button on the scanner control panel.
- 25. **Number of images** is a calculated value based on the Slice Thickness/Spacing and length of the scan.
- 26. **Scan Time** is a calculated value based on the protocol parameters selected. Scan time is affected by **Resolution**, **Pitch** and **Scan Length**.
- 27. For **CTDIvol (mGy)**, if applicable, the calculated number appears here, depending on other selections.
  - CT Dose Index Volume (CTDI<sub>vol</sub>) 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 CTDI<sub>vol</sub> is calculated differently for both the **Axial** and the **Helical** scan modes:
  - For Axial scan mode: CTDIvol = [(N x T)/I] x CTDIw
  - For Helical scan mode: CTDIvol = 1/pitch x CTDIw
- 28. **Dose Length Product (DLP (mGy.com))**, is the measure of ionizing radiation exposure during the entire acquisition of images.

  Therefore, DLP (mGy.cm) = CTDI<sub>vol</sub> (mGy) x irradiated length (cm).

29. Select the following options, if applicable. See Scanning with special features for more details.

Step & Shoot	Allows you to manually start the <b>Axial</b> scan acquisition from the workstation when scanning a patient who is unable to remain still.
Bolus Tracking	A CT angiography technique that allows you to monitor the administration of contrast to initiate the scan at peak contrast enhancement.
Enable AEC	Allows you to automatically adapt the tube current according to the patient's body habitus to achieve the specified image quality at the lowest possible dose.

30. To add a secondary reconstruction for the protocol, click the New button in the Recons section.



The **New Reconstruction** popup appears.

Figure 161: New Reconstruction popup

- 31. Complete the following in the **New Reconstruction** popup:
  - Enter a description in the **Description** text box to identify the new reconstruction.
  - Click the Slice Thickness/Spacing to select a slice thickness and spacing.
  - Click the **Sharpness** dropdown to select a sharpness from the list.
  - Enter the window width in the Window Width text box.

- Enter the window center in the Window Center text box.
- If needed, enter the FOV width in the FOV Width text box.
- If needed, enter the FOV top left x location in the FOV Top Left X text box.
- If needed, enter the FOV top left y location in the FOV Top Left Y text box.

#### 32. Perform one of the following:

- Click the Save button to save the reconstruction protocol to the list
  - The dialog box closes, and your changes are added to the Recons area.
- Click the Reset button to reset the fields to their original data.
- Click the **Cancel** button to remove your changes and return to the previous dialog box.



33. Click the **Save** button on the **New Series** dialog box.

Figure 162: Edit Series dialog box

Save Reset

CTDIvol (mGy) 16.91 DLP (mGy.cm)

Pitch

0.8

Body Part Exami

CHEST

Window Width

34. Repeat the steps 6 thru 33 to add additional scans to the protocol.

Remove

35. When all required series have been created click the **Save** button on the **New Protocol** dialog box.



Figure 163: Save New Protocol

### 36. Click the **Close** button to exit.

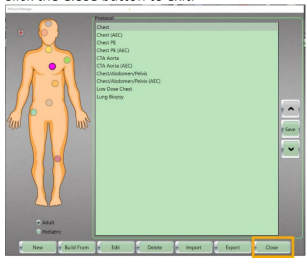


Figure 164: Close Button

# Using Build From to create a new protocol

The **Build From** button is used in **Protocol Manager** when you want to create a new protocol from an existing protocol.

- 1. Click **Tools >Protocol Manager** from the main menu. The **Protocol Manager** dialog appears.
- 2. Click one of the following:

Adult	To create and/or scan with adult scan protocols,
	which are stored by anatomical location.

Pediatric	To create and/or scan with pediatric scan protocols,
	which are stored by anatomical location.
Trauma	The <b>Trauma</b> orb can be used to store protocols
	commonly used for emergency scans.
Adult     Pediatric	By selecting either an <b>Adult</b> or <b>Pediatric</b> patient, the
	corresponding list of saved protocols becomes
	available.

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.

3. Click the colored orb corresponding to the appropriate body part.

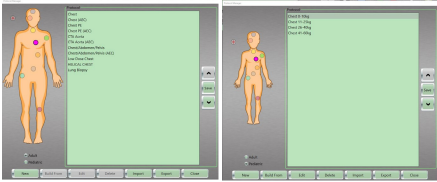


Figure 165: Anatomical orbs

4. Click the protocol you will **Build From** in the **Protocol** list. The **Build From** button will become active.

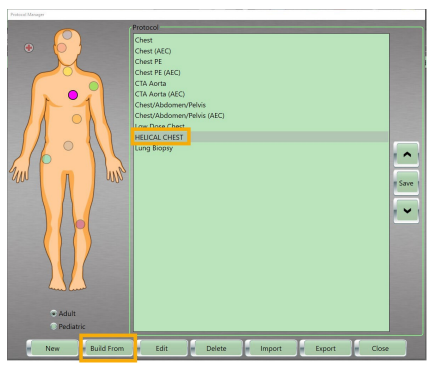


Figure 166: Build from protocol selected

5. Click the **Build From** button.

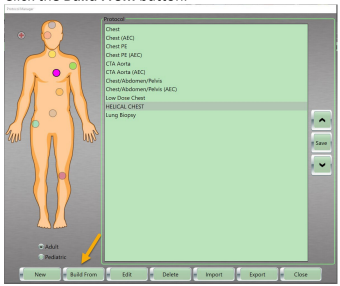


Figure 167: Build From button

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

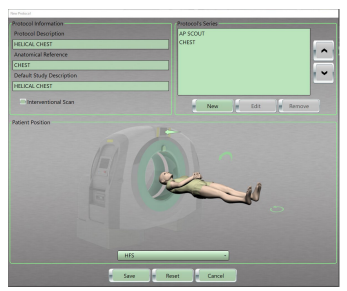


Figure 168: New Protocol dialog box

7. Modify the protocol parameters to meet your needs. Click the **Update** button on the **Edit Series** dialog box to save your changes.

Note Different patient sizes and attenuations require specific protocols where technical parameters have been adjusted according to the physical characteristics of the patient. It is recommended that you select pediatric protocols when scanning children, rather than selecting adult protocols and modifying technique factors.

See "Creating a new protocol" on page 190 to learn how the fields and options perform to make informed choices on what to change.

Note Be sure to assign the **Build From** protocol a new **Protocol Description** before you make your additional changes.

8. When all required series have been modified click the **Save** button on the **New Protocol** dialog box.

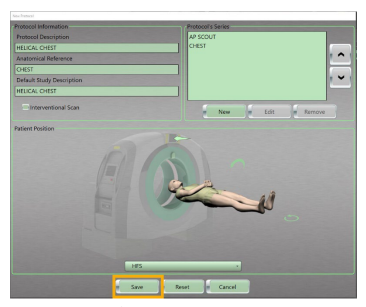


Figure 169: Build from save

9. Click the Close button to exit.

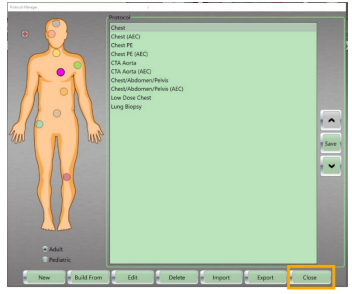


Figure 170: Build from close

# **Editing an Existing Protocol**

The **Edit** button is used in **Protocol Manager** when you want to modify the parameters of an existing protocol.

- 1. Click **Tools >Protocol Manager** from the main menu. The **Protocol Manager** dialog appears.
- 2. Click one of the following:

Adult	To create and/or scan with adult scan protocols,
	which are stored by anatomical location.
Pediatric	To create and/or scan with pediatric scan protocols,
	which are stored by anatomical location.
Trauma	The <b>Trauma</b> orb can be used to store protocols
	commonly used for emergency scans.
Adult     Pediatric	By selecting either an <b>Adult</b> or <b>Pediatric</b> patient, the
	corresponding list of saved protocols becomes
	available.

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.

3. Click the colored orb corresponding to the appropriate body part.

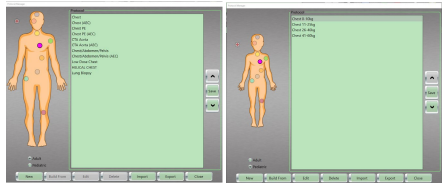


Figure 171: Edit protocol orbs

4. Click the protocol you wish to **Edit** in the **Protocol** list. The **Edit** button will become active.

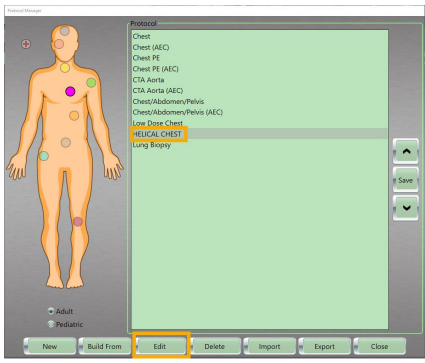


Figure 172: Edit protocol selected

#### 5. Click the Edit button.

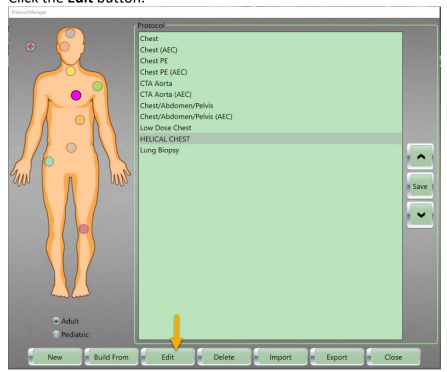


Figure 173: Edit button

Protocol Information
Protocol Description
HELICAL CHEST
Anatomical Reference
CHEST
Default Study Description
HELICAL CHEST
Interventional Scan

Patient Position

HELICAL CHEST
Update Reset Cancel

6. The Edit Protocol dialog box appears.

Figure 174: Edit Protocol dialog box

7. Modify the protocol parameters to meet your needs. Click the **Update** button on the **Edit Series** dialog box to save your changes.

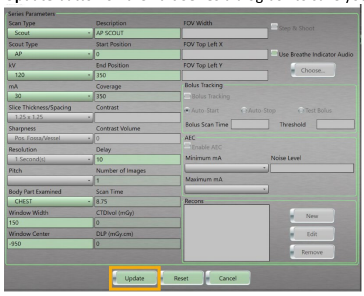


Figure 175: Edit series update button

Note Different patient sizes and attenuations require specific protocols where technical parameters have been adjusted according to the physical characteristics of the patient. It is recommended that you select pediatric protocols when scanning children, rather than selecting adult protocols and modifying technique factors.

See "Creating a new protocol" on page 190 to learn how the fields and options perform to make informed choices on what to change.

Note Be sure to assign the **Build From** protocol a new **Protocol Description** before you make your additional changes.

8. When all required series have been modified click the **Update** button on the **Edit Protocol** dialog box.



Figure 176: Edit protocol update button

9. Click the Close button to exit.

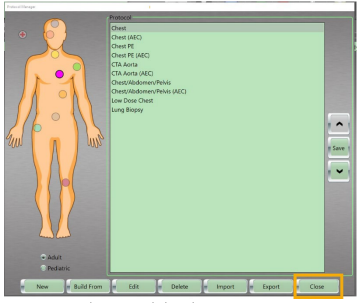


Figure 177: Edit protocol close button

## Copying and pasting protocols

To copy and paste protocols from one body part orb to another, including the Trauma orb.

- 1. Click **Tools >Protocol Manager** from the main menu. The **Protocol Manager** dialog appears.
- 2. Click one of the following:

Adult	To create and/or scan with adult scan protocols, which
	are stored by anatomical location.
Pediatric	To create and/or scan with pediatric scan protocols,
	which are stored by anatomical location.
Trauma <sup>®</sup>	The <b>Trauma</b> orb can be used to store protocols
	commonly used for emergency scans.
• Adult • Pediatric	By selecting either an <b>Adult</b> or <b>Pediatric</b> patient, the corresponding list of saved protocols becomes available.

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.

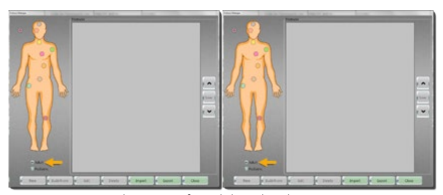


Figure 178: Protocol Manager for Adult and Pediatric

3. Click the colored orb corresponding to the appropriate body part.

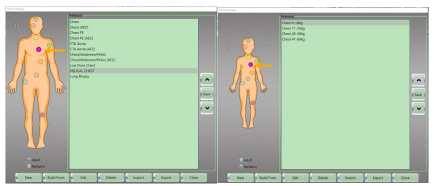


Figure 179: Anatomical orbs, in this case the chest orb

- 4. Review the protocol you would like to copy.
- 5. Highlight the protocol, right-click to see the floating menu, and click **Copy**.

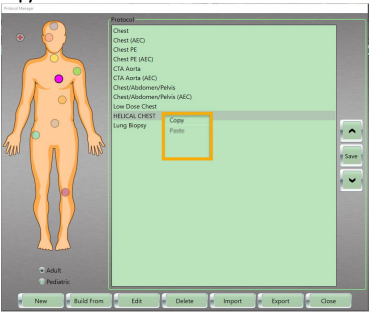


Figure 180: Copy right-click floating menu

6. Go to body part orb you want to paste the protocol to, which can include the Trauma orb.

Protocol
Abdomen Pelvis (AEC)
CTA Aorta
CTA Aorta (AEC)
Biopsy

Copy
Paste

Adult
Pediatric

New
Build From
Edit
Delete
Import
Export
Close

7. Right-click to see the floating menu and click Paste.

Figure 181: Paste right click floating menu

8. Click Close button to exit.

# **Deleting a protocol**

- 1. Click **Tools >Protocol Manager** from the main menu. The **Protocol Manager** dialog box appears.
- 2. Click one of the following:

Adult	To create and/or scan with adult scan protocols,
	which are stored by anatomical location.
Pediatric	To create and/or scan with pediatric scan protocols,
	which are stored by anatomical location.
Trauma	The <b>Trauma</b> orb can be used to store protocols
	commonly used for emergency scans.
• Adult • Pediatric	By selecting either an <b>Adult</b> or <b>Pediatric</b> patient, the corresponding list of saved protocols becomes
	available.

3. Click the colored orb corresponding to the appropriate body part. Select the protocol from list to be deleted.

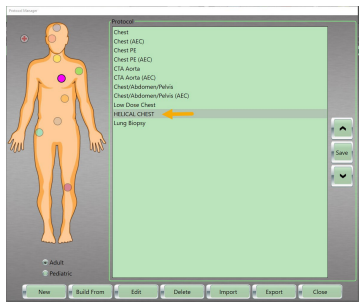


Figure 182: Protocol Manager with a protocol selected

4. Click the **Delete** button.

The **Delete Confirmation** popup appears.

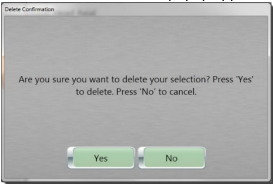


Figure 183: Delete Confirmation popup message – Yes or No to delete selection

- 5. Perform one of the following in the **Delete Confirmation** popup:
  - Click the Yes button to delete the selected protocol.
  - Click the **No** button to return to the Protocol Manager dialog box. The **Delete Confirmation** dialog box disappears, and the **Protocol Manager** dialog box appears.
- 6. Click the Close button to exit.

# Adding breathing instructions to your protocol

Default audio files are installed on the workstation. Audio files can be attached to protocols and sent to the scanner. Each audio file has an indicator whether it was sent to the scanner.

- 1. Click **Tools >Protocol Manager** from the main menu. The **Protocol Manager** dialog box appears.
- 2. Click one of the following:

Adult	To create and/or scan with adult scan protocols, which are
	stored by anatomical location.
Pediatric	To create and/or scan with pediatric scan protocols, which are
rediatric	stored by anatomical location.
Trauma	The <b>Trauma</b> orb can be used to store protocols commonly
	used for emergency scans.
• Adult • Pediatric	By selecting either an <b>Adult</b> or <b>Pediatric</b> patient, the corresponding list of saved protocols becomes available.

- 3. Click the colored orb corresponding to the appropriate body part.
- 4. Click the protocol you would like to add **Breathing Instructions** to.
- 5. Click the Edit button.

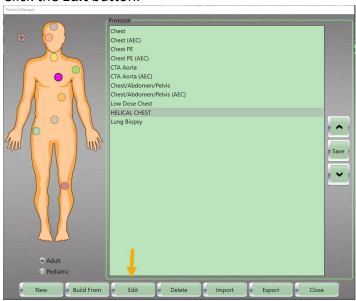


Figure 184: Edit button

The **Edit Protocol** dialog box appears.

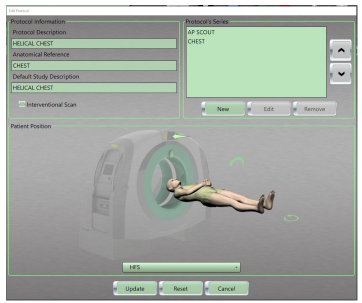


Figure 185: Edit Protocol dialog box

6. Select the **Protocol's Series** you want to add breathing instructions to.

The **Edit** button is enabled.

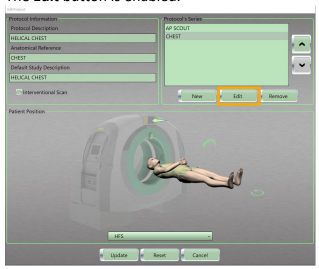


Figure 186: Add breathing edit button

7. Click the **Edit** button.

The **Edit Series** dialog box appears.

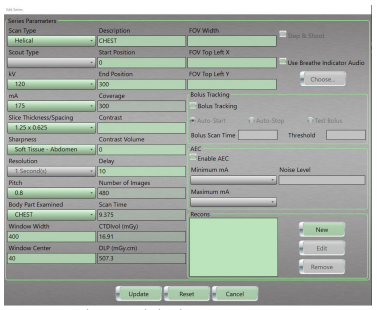


Figure 187: Edit Series dialog box

8. Click the **Use Breathe Indicator Audio** option and click the **Choose** button.

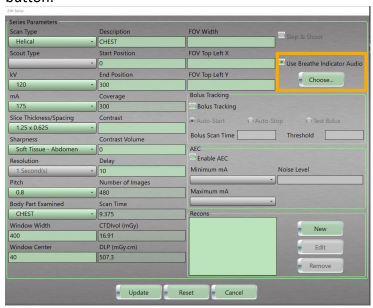


Figure 188: Use Breathe Indicator Audio option

The **Breathe Indicator Audio Files** popup appears.

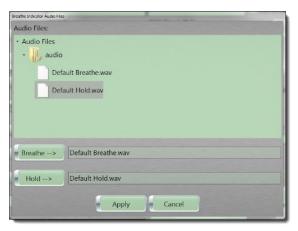


Figure 189: Breathe Indicator Audio Files popup

- 9. Select an audio file to use for the **Breathe** instruction. The **Breathe** and **Hold** buttons are activated.
- 10. Click the **Breathe** button to place the file with **Breathe** files.
- 11. Select an audio file to use for the **Hold** instruction.
- 12. Click the **Hold** button to place the file with **Hold** files.
- 13. Click the **Apply** button to keep the files you selected to use with protocols.
- 14. Click the **Close** button to exit.

## Importing protocols from a storage device

- Click Tools >Protocol Manager from the main menu.
   The Protocol Manager dialog box appears.
- 2. Click the **Import button**

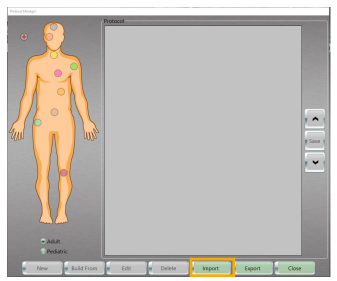


Figure 190: Import button

3. The **Select File** popup appears.



Figure 191: Select File popup

4. Double click the Drive Letter that contains the protocols you want to import.



Figure 192: Select file

5. Click the file in the **Select File** popup.



Figure 193: Import button active in Select File when file(s) selected

6. Click the **Import** button.



Figure 194: Protocols Imported popup message – Protocols imported

- 7. The **Protocols Imported** popup appears.
- 8. Click the **OK** button.
- 9. Check that the required files have been imported.
- 10. Click the **Close** button to exit.

# Exporting protocols to a storage device

- Click Tools >Protocol Manager from the main menu.
   The Protocol Manager dialog box appears.
- 2. Click the Export button.

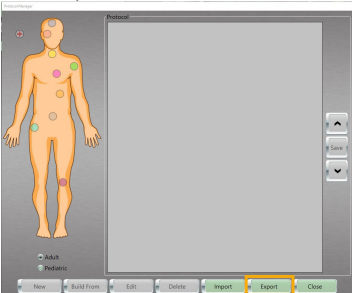
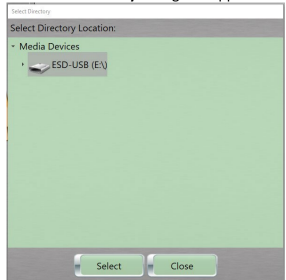


Figure 195: Export button



3. The **Select Directory** dialog box appears.

Figure 196: Select Directory popup

- 4. If more than one **Media Device** is available, select the device to use.
- 5. Click the **Select** button.
- 6. The **Protocols Exported** popup appears.



Figure 197: Protocols Exported popup message – Protocols exported

- 7. Click the **Ok** button.
- 8. Check that the exported files are exported.
- 9. Click the **Close** button to exit.

## Changing the order of protocols in the list

- Click Tools >Protocol Manager from the main menu.
   The Protocol Manager dialog box appears.
- 2. Click one of the following:

Adult	To create and/or scan with adult scan protocols, which
Addit	are stored by anatomical location.
Pediatric To create and/or scan with pediatric scan protoc	
rediatric	which are stored by anatomical location.
_	The <b>Trauma</b> orb can be used to store protocols
Trauma	commonly used for emergency scans.
• Adult • Pediatric	By selecting either an <b>Adult</b> or <b>Pediatric</b> patient, the corresponding list of saved protocols becomes available.

- 3. Click the colored orb corresponding to the appropriate body part.
- 4. Click the protocol to move up or down the list.
- 5. Click the **Up** arrow to move the protocol up the list; click the **Down** arrow to move the protocol down the list.

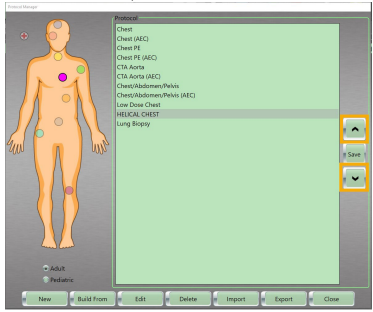


Figure 198: Changing protocol order with Up and Down (arrow) buttons

6. When you are finished ordering your protocols, click the **Save** button to save the new ordered list.

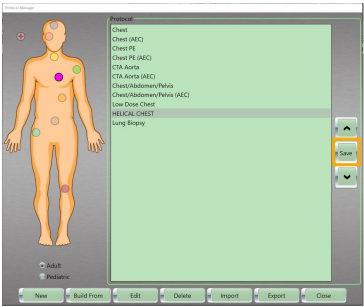


Figure 199: Protocol Save button

7. Click the **Close** button to exit.

# Chapter 7 Daily Calibration and Quality Assurance

In this chapter, you will learn how to perform a daily air calibration and use the **Quality Assurance** (**QA**) tool that verifies the system is working as specified.

Keep in mind that **before** using the BodyTom Elite system, you **must** conduct a **Quality Assurance** (**QA**) test to verify the system is working as specified. Performing a daily (air) calibration

Note NeuroLogica recommends that an air calibration is performed every 6-8 hours. If the air freshness falls below 50%, or the scanner is moved to an area with a dramatic change in humidity or temperature, perform another air calibration to ensure optimum image quality during patient scanning.

If room-temperature fluctuations have occurred, you may need to perform more than one air calibration. In addition, scanners can drift out of alignment; make sure you perform a **QA Test** with the test phantom **before** scanning a patient.

**Note** It is recommended that the scanner is on for at least 60-90 minutes prior to performing the air calibration.

It is recommended practice that the scanner is plugged in and turned on even when it is not in use.

- 1. Make sure that nothing is in the bore before starting the daily air calibration.
- Click Tools > Perform Daily Cal from the main menu.The Perform Daily Cal popup appears.



Figure 200: Perform Daily Cal popup

Colors identify previous air calibrations outcomes:

Cuan	Indicates the calibration was
Green	successful
Yellow	Indicates the calibration is soon
Tellow	to expire
Orango	Indicates the calibration has
Orange	expired
<b>Red</b> Indicates the calibration failed.	

- 3. Select one of the following options:
  - Click Select All to perform all calibration steps.
  - Click Clear All and individually select the calibration step(s) to perform.
- 4. Click the Start button.

The **Perform Daily Cal** popup appears, and the timer counts down.

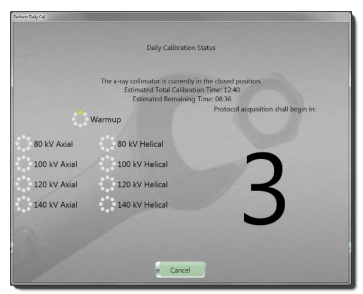


Figure 201: Perform Daily Cal popup with count down

A warmup period begins, and the countdown begins; when completed the daily calibration will perform the calibration(s) you selected.

**Note** To stop the calibration, click the **Cancel** button to end the daily (air) calibration(s).

When the calibrations are completed the **Daily Calibration Summary** will display showing the status of the steps performed.



Figure 202: Perform Daily Cal summary popup

The following are the status indicators:

Green	Identifies the calibration completed successfully.
Yellow	Identifies the calibration is in progress.
Red	Identifies the air calibration failed.

Click the Close button to exit the Perform Daily Cal popup.
 The Daily Cal icon will change to green when it reaches a 100% air freshness.



Figure 203: Air freshness icon changes as the air quality drops from green to yellow to red

#### The QA phantom overview

The **QA phantom** is a device that measures parameters that characterize image quality; these parameters are as follows:

- Uniformity
- Noise
- High-contrast resolution
- · Slice width
- Low-contrast resolution
- Sensitometry (contrast scale)

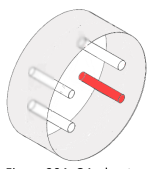


Figure 204: QA phantom

The **QA phantom** is a 20cm diameter disk consisting of a substrate made of **poly methyl methacrylate** (**PMMA**) containing specific inserts. The uniform area of the disk is used to measure uniformity and noise. Four other parameters are measured by the inserts in the substrate.

The QA phantom goes onto the phantom holder when performing a QA. See "Storing the QA phantom" on page 360.



Figure 205: Phantom holder

The **Axial** resolution wire, also called the **Modulation Transfer Function** (**MTF**) wire, is intended for measuring resolution in the **Axial** plane. Resolution is defined as the ability to distinguish small objects. It is expressed in line pairs per millimeter.

The **slice width wires** are the two inclined wires. They are intended to determine scanner resolution along the Z axis, that is, in the direction that is perpendicular to the **Axial** plane. Resolution along the Z axis is expressed in terms of slice width in millimeters. Although one wire is sufficient to measure the Z axis resolution if its position is accurately known, a second wire is included to confirm the alignment. If the alignment was incorrect, the results of the slice width test would not be accurate.

The low-contrast insert is a compound insert. It is made of two half cylinders of different materials with a known contrast difference between them. The low-contrast insert is intended to measure the contrast resolution of the scanner. The contrast resolution is the ability to measure slight differences in x-ray attenuation.

The sensitometry inserts are an air bore and cylinder made of different materials. They are intended to measure the contrast scaling of the scanner.

The QA scan protocols appear in the following table.

Scan voltage 120 kV
Scan current 200 mA
mAs 400 mAs
Scan time 2 second
Kernel Pos. Fossa/Vessel
Slice thickness 10mm

Table 28: Scan protocols used by the QA

#### **Starting Quality Assurance**

To ensure the system is at its optimum, factory-specification level, the workstation provides **QA** tools to verify the system's state and to perform image-quality verification. To maintain consistent image quality over the system's lifetime, you should establish and maintain a regular **Quality Assurance** (**QA**) program. **QA** results are stored in the **Patient Browser**. Contact your local service representative to delete **QA** results.

The **QA protocol** is shipped with the system and appears when you click **Quality Assurance** from the main menu. You cannot customize or modify the **QA protocol**.

**Note** The **QA test** should be conducted per hospital requirements.

Before you begin this section, be sure to run a fresh **Daily Calibration** on the system. See "Performing a daily (air) calibration" on page 222.

Also, before you start the **QA protocol**, make sure the **QA phantom** is available and ready to install on the phantom holder. The phantom serial number label should be facing the front of the scanner and be positioned at the top – as shown in the figure below. The red insert should be on the bottom right when facing the scanner. The position of the phantom will greatly affect the **QA** results.

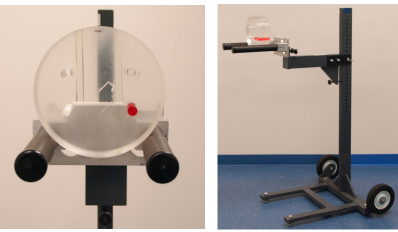


Figure 206: Phantom on the phantom holder

1. Move the QA stand to the front of the gantry, so the prongs point into the bore as shown.



Figure 207: Place QA phantom

2. Ensure the QA stand is centered in the bore using the sagittal laser as shown. If needed, adjust the prongs side to side.

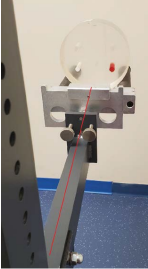


Figure 208: Proper QA stand positioning

3. Place the QA phantom on the prongs as shown.

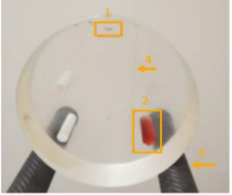


Figure 209: QA phantom positioning

- 1. The serial number label should be facing the front of the gantry and at the top.
- 2. The red insert should be on the bottom right.
- The phantom should be in the middle of the carbon fiber post/prongs.
- 4. The two wires in the phantom

4. On the pendant, press the **Laser** button and align the internal laser to the etched line in the center of the phantom.



Figure 210: Laser button See the laser precautions in "Laser safety" on page 58.

- 5. Click **Tools > Quality Assurance** from the workstation main menu.
- 6. The following **Quality Assurance** popup appears.



Figure 211: Quality Assurance popup

- 7. Click the **Prepare** button to begin the QA procedure.
- 8. The **System Ready to Scan** popup appears.



Figure 212: System Ready to Scan popup message – System is ready to begin

9. Go to the scanner and press the **START** button. The system will scan the phantom and display the **QA Results** image.



Figure 213: QA results of QA image

**Note** Items in green are passed results. Items in red are failed results. Often positional issues cause the failure; reposition your phantom and perform another scan. If you try multiple times and failures persist, call your service representative or **Technical Support**.

10. Click the **Close** button on the **QA Results** popup when finished reviewing. The image of the phantom appears.

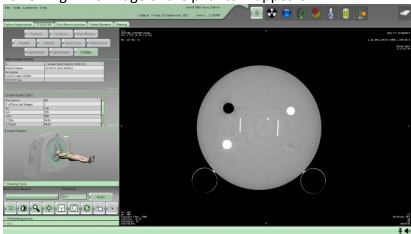


Figure 214: Phantom image

- 11. Click the **Finalize** button on the workstation to exit the protocol.
- 12. The **QA** appears in the **Patient Browser**; however, it is locked.



Figure 215: Locked QA results shown in Patient Browser

**Note** See your service representative to remove locked QA results.

### **Ensuring good image quality**

To produce consistent image quality over the system's lifetime, you should establish and maintain a regular **Quality Assurance** (**QA**) program. **QA** results are stored in the Patient Browser. Contact your local service representative to delete **QA** results.

 Compare the results to previous or optimum values and repeat these tests on a regular basis to detect changes in image quality values before any problem becomes visible. Note If you notice degradation in image quality or a change in QA values, schedule a site visit and let your service representative or imaging physicist run more detailed tests.

Early intervention could prevent a major breakdown.

**QA** begins with baseline performance data that is acquired during system installation or after the repair or replacement of an x-ray generator-assembly, collimator, detector, Data Acquisition System (DAS) or main power circuitry.

2. Compare subsequent **QA** results against the baseline.

Baseline images can be saved for a visual comparison with QA checks, but measurement values provide a more objective way to monitor quality.



Figure 216: Results of QA image after the QA test

## Identifying filtration accuracy

Values of attenuation equivalent, half-value layer, and quality-equivalent filtration are expressed as a thickness of aluminum at the minimum of 99.9% purity. Attenuation of items in the x-ray beam should not be higher than 2mm of equivalent Aluminum (AI).



**CAUTION** 

Do not put anything in the x-ray beam that exceeds 2mm of equivalent AL as it may produce adverse effects to the image.

#### Using Axial plane to determine image resolution

The method to determine resolution in the **Axia**l plane is to measure the modulation-transfer function of the scanner. A planar section of the **MTF** wire provides a point, called an **impulse**. The impulse is blurred by the imaging system, and the measurement of the blur quantifies the resolution. The blur is quantified by the **MTF** amplitude, which provides a measure of gain for a given object size in line pairs per centimeter (lp/cm).

The **MTF** is measured in two directions, called **radial** and **tangential** directions. The **radial** direction is along the line that joins the wire to the scanner isocenter. The **tangential** direction is perpendicular to the radial direction. The **MTF** along each direction produces a curve. The points at which each curve's amplitude is 50% and 10% of its amplitude at zero lp/cm are reported.

The expected results are given below.

Table 29: Modulation Transfer Function (MTF) direction

Direction	50%	10%
Radial (lp/cm)	4.7	7.2
Tangential (lp/cm)	4.7	7.2

Table 30: QA results

	Low limit	High limit
Slice width (mm)	9	11
Noise (HU)	2.5	3.5
Low-contrast resolution (mm)	4	6
Uniformity (HU)	0	5
CT of air (HU)	-1030	-950
CT of Teflon (HU)	924	1004
CT of acrylic (HU)	75	155

#### Using slice width

The method for determining the slice width for the **Axial** mode QA is to take an image of the inclined wire. The scanned section of the inclined wire is a line segment. The scanner blurs a scanned object in the **Axial** plane as well as in the direction perpendicular to it. The image of the inclined wire includes both the **Axial** plane blurring (**MTF**) of the scanner as well as the blurring in the z-direction. The slice width is determined by removing the component of in-plane blurring, by measuring the length of the wire segment and by using the known angle of wire inclination. The range is noted in QA results; see Table 29 on page 233.

#### Measuring noise

**Noise** is measured as the standard deviation of pixel values in a 1cm **Region of Interest (ROI)** at the center of the phantom. The **ROI** selection is automatic. The CTDI<sub>100</sub> center dose in a standard CTDI head phantom is 145 mGy for this scanning technique. The BodyTom Elite noise measurement is performed on a 10mm slice.

#### Measuring low contrast

**Low-contrast resolution** is measured as the difference between the mean CT values in each half of the low-contrast insert. An **ROI** is automatically selected around the low-contrast phantom and is automatically segmented into halves. Within each **ROI**, the mean pixel value is computed. The two mean values are subtracted.

The expected difference in the mean values is given in the electronic report. The low contrast should be:  $5.0 \pm 1.0 \text{ HU}$ .

## Finding uniformity

A **ROI** is automatically selected in each of five locations in the phantom. One **ROI** is at the center. Four outer **ROI**s are 60 to 70mm from the center of the phantom and spaced 90 degrees apart. A mean value is calculated in each **ROI**. The maximum difference between the means is calculated. The maximum allowable difference between the means is 3 HU.

Separate, independent measurements of the CT numbers of air, Teflon, and acrylic (see "Identifying CT contrast scale" on page 235 below) ensure that the overall CT number-scaling of the system is in order. If the uniformity portion of the QA protocol passes (maximum difference less than or equal to 3 HU), but the mean values of the five **ROI**s are all uniformly high or low in CT number, it is expected that the tests for air, Teflon, and/or acrylic will fail. See Table 39 on page 240.

#### Identifying CT contrast scale

Contrast scale represents the attenuation scaling of the scanner. The mean CT numbers of each of the sensitometry objects is calculated and reported.

#### Using performance phantoms

The phantoms (in the following sections) were used for measurement of dose and imaging performance.

#### Measuring dose

The dose is measured using the standard CTDI body phantom.

## **Identifying load factors**

Table 31: Load factors

Protocol description	kV	m A	Time (seconds)
Axial	80-140	30-300	1
Helical	80-140	30-300	1 per rotation



**CAUTION** When conducting multiple or repeat scans, ensure that the total exposure does not exceed 1Gy CTDI.

**Note** The highest x-ray tube current is 300mA and the highest x-ray tube voltage selection at this current is 140kV.

> The nominal x-ray output power is 42kW when operating at an x-ray tube voltage of 140kV and x-ray current of 300mA for 4 seconds.

The x-ray tube voltage/current tolerance is ±10%.

The nominal x-ray tube voltage is 140kV with 100cfm minimum cooling flow.

## The BodyTom Elite dose information (21 CFR 1020.33 c)

Dose is measured using standard CTDI head and body phantoms. Surface and center CTDIs were both measured. Weighted CTDI is computed using surface and center CTDIs:

$$CTDI_w = (\frac{2}{3} \ CTDI_{surf} + \frac{1}{3} \ CTDI_{cen})$$

Measured values are normalized to scan current, for example, CTDI values are in mGy/100 mAs. For any given scan protocol CTDI $_{\rm w}$  can be estimated using following equation and data from Table 32 and Table 33 on page 238:

$$CTDI_{w}(kV, m, S) = \left(\frac{m}{100.0} \cdot S\right) CTDI_{w}(kV, 100mAs) mGy$$

CTDI<sub>w</sub> can also be computed using data from Table 34 and Table 35 on page 238, and the following equation:

$$CTDI_{w}(kV, m, S) = \left(W(kV) \cdot \frac{m}{100.0} \cdot S\right) CTDI_{w}(120_{kV}) mGy$$

Where  $\boldsymbol{W}$  is the  $\boldsymbol{kV}$  relative dose ratio with respect to 120 kV.  $\boldsymbol{m}$  is the x-ray tube current in mA and  $\boldsymbol{S}$  is the scanning time in seconds. If scan kV matches measured scan voltage, then  $\boldsymbol{W}$  is equal to 1.0. For Helical scans,  $CTDI_{vol}$  is calculated as follows:

$$CTDI_{vol} = \frac{CTDI_{w}}{Pitch}$$

For **Axial** scans,  $CTDI_{vol}$  is calculated as follows:

$$\mathit{CTDI}_{vol} = \frac{\mathit{CTDI}_w}{\mathit{Scan Increment}}$$

For example, CTDI<sub>w</sub> of 2 seconds, 100 kV **Axial** head scan, with 200 mA can be evaluated as follows using data from Table 32 and Table 33 on page 238:

$$CTDI_{w}(100kV, 200mA, 2 Sec) = \left(\frac{200.0}{100.0} \times 2\right) CTDI_{w}(100_{kV}, 100mAs)$$
  
=  $4 \times (100_{kV}, 100mAs) \ mGy = 4 \times 14.4 = 57.6 \ mGy$ 

Using data from Table 34 and Table 35 on page 238, CTDI<sub>w</sub> can be computed as follows:

$$CTDI_{w}(100kV, 200mA, 2 Sec) = \left(0.626 \times \frac{200.0}{100.0} \times 2\right) CTDI_{w}(120_{kV}, 100mAs)$$
  
=  $2.504 \times (120_{kV}, 100mAs) \ mGy =$   
 $2.504 \times 23.0 = 57.59 \ mGy$ 

CTDIvol is evaluated as follows:

$$CTDI_{vol} = \frac{CTDI_w}{Scan\ Increment} = \frac{57.6}{1.0} = 57.6\ mGy$$

In another example, we can compute  $CTDI_w$  of 1 second, 120 kV **Helical** abdomen scan, with 200 mA and a pitch of 0.8. Dose can be evaluated as follows, using data from Table 32 and Table 33 on page 238:

$$CTDI_{w}(120kV, 200mA, 1 Sec) = \left(\frac{200.0}{100.0} \times 1\right) CTDI_{w}(120_{kV}, 100mAs)$$
  
=  $2 \times (120_{kV}, 100mAs) mGy$   
=  $2 \times 7.56 = 15.12 mGy$ 

Using data from Table 34 and Table 35 on page 238,  $CTDI_w$  can be computed as follows:

$$CTDI_{w}(120kV, 200mA, 2 Sec) = \left(1.0 \times \frac{200.0}{100.0} \times 1\right) CTDI_{w}(120_{kV}, 100mAs)$$
  
=  $2.0 \times (120_{kV}, 100mAs) \ mGy = 2.0 \times 7.56 = 15.12 \ mGy$ 

CTDI<sub>vol</sub> is evaluated as follows:

$$CTDI_{vol} = \frac{CTDI_w}{Scan\ Pitch} = \frac{15.12}{0.8} = 18.9\ mGy$$

#### **Body CTDIw phantom**

 $CTDI_{w}$  using CTDI body phantom is listed in the following table. Data was measured using the 32 rows collimation with the phantom placed on the phantom holder. Dose measurements were taken using raw data acquisition in **Service** mode.

**Note** Performing scans in different **Acquisition** modes can cause slight variations in measured dose.

Table 32: Body CTDI<sub>w</sub> (mGy/100mAs)

	140 kV	120 kV	100 kV	80 kV
CTDI <sub>100</sub> Center (C)	6.81	4.54	2.65	1.20
CTDI <sub>100</sub> Surface (S)	14.1	9.84	6.49	3.48
CTDI <sub>w</sub>	11.7	8.07	5.21	2.72

#### Head CTDIw phantom

Weighted average Computed Tomography Dose Index (CTDIw) using the CTDI head phantom is listed in the following table. Data was measured using the 8 rows collimation. Dose measurements were taken using raw data acquisition in **Service** mode using phantom holder.

Table 33: Head CTDI<sub>w</sub> (mGy/100mAs)

	140 kV	120 kV	100 kV	80 kV
CTDI <sub>100</sub> Center (C)	34.3	23.6	14.6	7.55
CTDI <sub>100</sub> Surface (S)	38.1	26.7	17.0	9.33
CTDI <sub>w</sub>	36.8	25.7	16.2	8.74

Normalized CTDI tables are listed below. CTDI is normalized with respect to a typical 120kV scan protocol:

Table 34: Normalized CTDI of body phantom

	140 kV	120 kV	100 kV	80 kV
CTDI <sub>100</sub> Center (C)	1.50	1.00	.584	.264
CTDI <sub>100</sub> Surface (S)	1.43	1.00	.660	.354
CTDI <sub>w</sub>	1.45	1.00	.646	.337

Table 35: Normalized head CTDI

	140 kV	120 kV	100 kV	80 kV
CTDI <sub>100</sub> Center (C)	1.45	1.00	.619	.320
CTDI <sub>100</sub> Surface (S)	1.43	1.00	.637	.349
CTDI <sub>w</sub>	1.43	1.00	.630	.340

#### The BodyTom Elite dose in air

Dose measurements were taken using raw data acquisition in **Service** mode.

Table 36: CTDI air (mGy/100mAs)

	140 kV	120 kV	100 kV	80 kV
32 rows	31.2	22.5	14.9	8.56
8 rows	47.3	33.4	21.9	12.6

Dose is measured using a typical head protocol and a typical abdomen protocol. Dose in air was also measured for repeatability over 10 scans. Average value and standard deviation are noted below:

Table 37: Mean and standard deviation of CTDI air

	8 rows	32 rows
Mean mGy	19.3596	13.18493
Standard deviation mGy	0.058716	0.017831

#### **QA** measurements

The QA phantom is typically used to monitor the scanner on site; however, the following phantoms can be used for measuring the imaging performance of the scanner.

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

## **Tube accuracy**

Table 38: Tube accuracy

kV	Tolerance
140	± 5%
120	± 5%
100	± 5%
80	± 5%

## **ACR** testing procedure

Most sites use the ACR phantom for evaluating the QA parameters of the scanner. Furthermore, each scanner is evaluated using the ACR phantom prior to shipping. Due to the special tube filtration some of the limits for CT values may be different from those set by the ACR committee. Table 39 lists the NL limits for the CT number and linearity of each insert in the ACR phantom. It also lists the limits set by ACR. The difference in CT

number is mainly due to the difference of the x-ray beam quality due to the tube filtration and the ACR committee acknowledge this in published papers3.

Table 39: The CT number and linearity of the different inserts in the ACR phantoms

P		
Insert Material	NL Limits	ACR Limits
Air	-1005 to -970	-1005 to 970
Polyethylene	-110 to -85	-110 to -85
Water	-7 to 7	-7 to 7
Acrylic	110 to 135	110 to 130
Bone	1010 to 1110	850 to 970

The scan protocols are typically selected by the site physicist in the CT manager. However, ACR recommends the use of standard head and abdomen protocols. NeuroLogica uses the protocols listed in Table 40.

Table 40: The NeuroLogica head and abdomen ACR scan protocols

Protocol	Head/Abdomen	Abdomen
kVp	120	120
mA	200	250
Time per rotation (seconds)	1	1
Dose (Weighted)	45 to 50 mGy	22 to 25 mGy
Scan FOV	59.5 cm	59.5 cm
Display FOV (minimum)	25.0 cm	25.0 cm
Reconstruction sharpness	Soft tissue	Soft Abdomen
Scan type	Axial	Helical
Z-axis collimation	10 mm	40 mm
Table increment (mm) or Table speed (mm/rot)	10 mm	32 mm/rot
Slice thickness	5 mm	5 mm
Scan time (seconds)	1	1
Slice separation	5 mm	5x5 mm
Number of images per scan	2	N/A

<sup>&</sup>lt;sup>3</sup> The ACR committee will often change their limits, as such, the limits listed in Table 39 may have been changed.

#### Identifying high-contrast resolution

The high-contrast resolution phantom is a wire placed at the center of a uniform disk. The wire provides an impulse function in the **Axial** plane when it is placed parallel to the scanner-gantry axis-of-rotation. The high-contrast resolution is measured from the **Modulation Transfer Function (MTF)**. Typical **MTF** curves are shown in the following figures. Variations of 10% may occur in measurements due to phantom placement error and measurement inaccuracies.

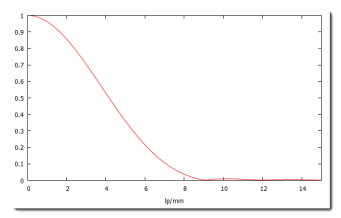


Figure 217: MTF

Table 41 lists the 50%, 20% and 10% cutoffs of the most commonly used kernels on the scanner. The cutoffs were measured using the MTF curve for each kernel like the one displayed in Figure 217.

Scan Type	Kernel	MTF50%	MTF20%	MTF10%
Axial	Soft Tissue	3.4038	4.9538	5.7187
Axial	Pos. Fossa/Vessel	3.9930	5.7816	6.7455
Axial	Sharp	6.6819	8.2549	9.2587
Axial	Bone	7.7757	9.6942	10.9681
Axial	Sharp Lung	6.1620	8.9318	11.8802
Axial	High-Res QA	7.9286	10.5069	12.4050
Helical	Bone Head	5.2370	6.5734	7.1577
Helical	Soft Tissue-Abd	6.2422	7.2252	7.7846
Helical	Soft Tissue-Head	3.0327	4.4322	5.1328
Helical	Bone-Ahdomen	5 5166	6 8081	7 4292

Table 41: The cutoffs of some of the common reconstruction kernels

## Noise, uniformity, and mean CT number of water

One of two phantoms may be used in these tests. These are Catphan® 412 or a cylindrical 20cm diameter water cylinder.

The variation in standard deviation may be ±10% due to variations between systems.

**Noise** is measured as the standard deviation at isocenter. The value is  $2.1 \pm .2 \, \text{HU}$  when the imaging protocol is 140 kV, 42 mAs and standard kernel. This protocol gives a CTDI<sub>100</sub> center dose of 160 mGy.

Noise is measured as the standard deviation of pixel values in a 1cm **ROI** at phantom's center. Range is as noted in QA results (see Table 30 on page 233).

Table 42: Uniformity and Mean CT Number using Water Phantom

Description	Noise (HU)
Body	11.9HU
(cp300 mm Water phantom)	
Head (cp200mm Water Phantom)	2.89HU

#### Uniformity and mean CT number

The mean CT number of air is -1000 HU and that of water is 0 HU. The tolerance of the mean CT number will be  $\pm 3$  HU. For mean CT numbers measured at different points of the water phantom, the maximum difference in the means will be less than 4 HU<sup>4</sup>.

An **ROI** is automatically selected in each of five locations in the phantom. One **ROI** is at center. Four outer **ROI**s are 60 to 70mm from phantom center and spaced 90 degrees apart. A mean value is calculated in each **ROI**. The maximum difference between means is calculated. The maximum allowable difference is as noted in QA results (see Table 30 on page 233).

Maximum difference between periphery **ROI**s and center **ROI** mean CT values in an image is less or equal to 4 HU. The maximum error in CT number of water is ±3 HU.

Table 43: Uniformity and Mean CT Numbers using Water Phantom

Description	Uniformity (HU)	Mean CT Number (HU)
Body	1.91	0.32
(cp300 mm Water phantom)		
Head (cp200mm Water	1.21	1.21
Phantom)		

<sup>&</sup>lt;sup>4</sup> For States that required the sites to perform water phantom testing, please contact customer service for assistance.

#### **Low-contrast resolution**

The phantom used for low-contrast resolution measurement is CTP 515 section of the Catphan 600.

The **low-contrast resolution** is 4mm rod at 0.3% contrast when the center CTDI<sub>w</sub> dose is 71 mGy. The imaging protocol is 120 kV, 300 mAs with 1 rotation, 5mm slice thickness, and using the soft tissue filter.

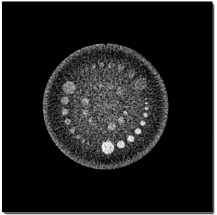


Figure 218: Catphan 515 using 120kV, 300mA, 1 rotation, and 5mm slice

#### Geometric efficiency in the Z axis direction notification

This is the ratio of the integral of the dose profile integrated over the detector width in Z divided by the total  $CTDI_{100}$ . The geometric efficiencies for the two available collimations are listed in Table 47 on page 251. The scanner has two scan modes: Axial and Helical. The Helical scan mode uses the wide collimation of 32 rows therefor it is accuracy is around  $83\pm5\%$ . The Axial scan uses the narrow collimation of 8 rows only with it is geometric efficiency of  $60\pm5\%$ . When this occurs, the operator will be given an opportunity to continue/affirm the prescribed scan. See Figure 219.

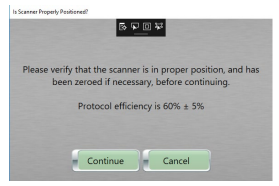


Figure 219: Geometric efficiency confirmation

#### Half-value layer

Table 44: Half-value layer

Scan voltage (kV)	100	120	140
Half value	6mm	7mm	8mm

#### Allowable variations

The following are allowable variations:

	A ±5% variation in dose may occur due to
Dose	variations between systems and measurement
	differences. The maximum variation is ±10%.
	The variation in values on the MTF curve may be
High Contrast	±10%. These will occur mainly due to phantom
Resolution	placement errors, measurement inaccuracies and
	system variations.
Noise	The variation in standard deviation may be ±10%
Noise	due to variations between systems.
	The maximum difference between ROI means in
Uniformity	an image is 4 HU. The maximum error in the CT
	number of water is ±3 HU.

Dose: Maximum variation is ±10%.

Variation in values on the **MTF** curve may be ±10%. These occur mainly due to phantom placement errors, measurement inaccuracies and system variations.

#### Scatter radiation

Reference the following radiation scatter plots identifying proper distances to protect from radiation exposure. The scatter plots provide scattered radiation dose in air-kerma, per current-time product in both standard and SI units for nominal technique of 120kV ( $\mu$ Rad/100mAs and  $\mu$ Gy/100mAs respectively). This information is given so the facility physicist and/or **Radiation Safety Officer** (**RSO**) can use these charts to calculate exposure with the following formula:

$$Stray\ radiation\ (scan\ current, scan\ voltage) = stray\ radiation\ (100, 120) \times \left(\frac{scan\ current}{100}\right) \times \left(\frac{scan\ voltage}{120}\right)^{2.3}$$

In addition, per IEC 60601-2-44, 3<sup>rd</sup> Edition, "Particular requirements for the basic safety and essential performance of x-ray equipment for computed tomography", scatter plots are provided as shown in Figure 222 on page 247 and Figure 223 on page 248 for the maximum techniques settings of 140kV and 300mA (standard and SI units respectively).

This information is specifically intended for the facility Physicist and/or an **RSO** to perform a safety and shielding analysis such as described in NCRP 147, "Structural Shielding Design for Medical X-Ray Imaging Facilities."



#### WARNING

Exposure to secondary radiation can be harmful, and scanner usage should only be done under the direct supervision of the facility's qualified **Radiation Safety Officer (RSO)** in compliance with site, local, mode, 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, and so on.

**Note** The BodyTom Elite scanners are compatible with IRR1999 and EU Directive 96/29/EURATOM.

#### Typical application environment and radiation safety

The BodyTom Elite is an advanced radiation protection mobile CT. There is an effective x-ray shielding that is equivalent of 0.75mm of lead within the gantry. Under normal circumstances, no additional radiation shielding is needed per recommendations of the facility physicist. The scanner can be used in a mobile environment and/or within an enclosed environment.

The scatter plot below shows the dose map during a normal scan:

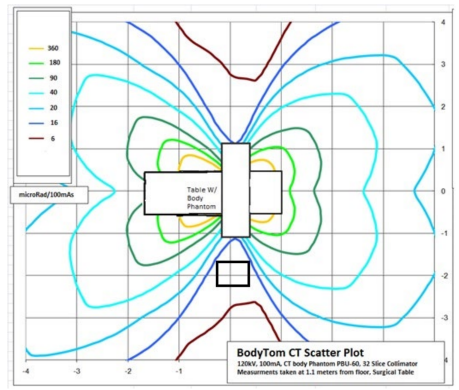


Figure 220: Scatter plot (120kV, 100mA in μRad)

Note In compliance with IEC 60601-2-44:2009, section 203.11, the above 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.

The black box (located at 0 on the X axis and -2 on the Y axis) represents an approximate (24 x 24 x 79in. or 60 x 60 x 200cm) zone of occupancy. The system in **Scan** mode stands at 78.5in., which meets the 79in. (200cm) requirement.

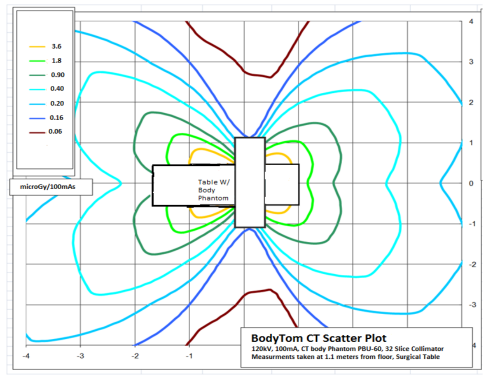


Figure 221: Scatter plot (120kV, 100mA in μGy)

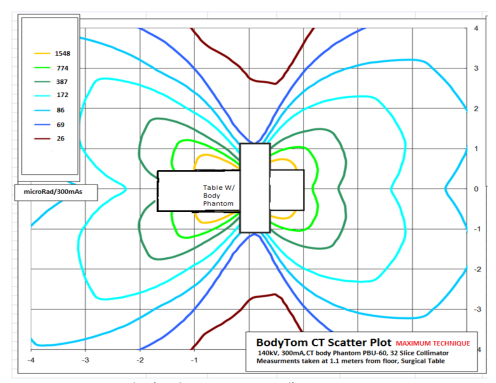


Figure 222: Scatter plot (140kV, 300mA in μRad)

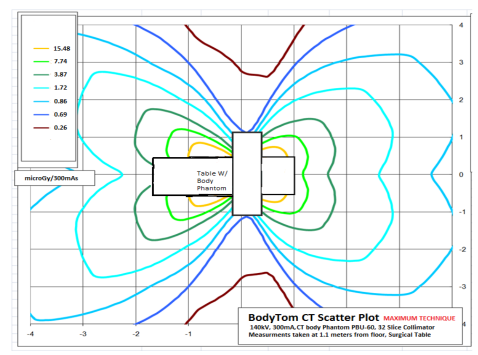


Figure 223: Scatter plot (140kV, 300mA in μGy)

Note The anatomical body phantom was placed on a scan table inside the gantry to consider scatter through patient. Measurements were made using the following scan protocol: 140kV, 200 mA, and 5 sec. The following figures show measurement points in vertical X—Y and perpendicular Y—Z planes, followed by corresponding tables detailing resulting data.

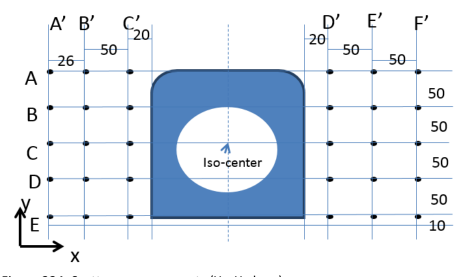


Figure 224: Scatter measurements (X—Y plane)

Table 45: Scatter measurements (X—Y plane) (μRad/100 mAs)

	A'	B'	C'	D'	Ε'	F'
А	12.2	15.3	17.7	18.3	15.7	12.7
В	11.9	14.8	16.4	16.7	16.0	12.6
С	11.4	13.6	11.1	12.9	15.2	12.5
D	9.40	10.4	8.87	9.57	12.4	10.4
Е	6.09	6.26	4.09	4.26	7.66	7.66

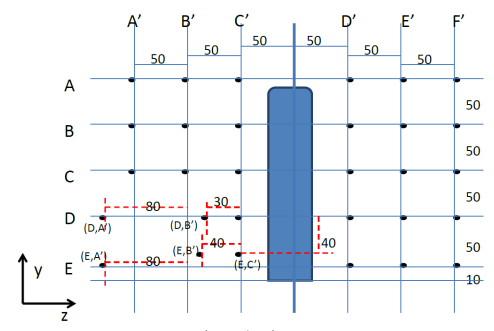


Figure 225: Scatter measurements (Y—Z plane)

Table 46: Scatter measurements (Y—Z plane) (μRad/100 mAs)

	A'	B'	C'	D'	Ε'	F'
А	300	676	852	591	870	437
В	218	539	2320	2940	1090	465
С	137	328	931	1120	461	233
D	27.8	1080	2790	2240	765	282
Е	8.96	844	441	85.3	538	345

#### Dose profile/Geometric Efficiency

A graphical presentation of the **dose profile** along a line -Z perpendicular to the **tomographic plane** and centered at the **isocenter**, determined in free air for one **Axial** scan, in the center location of the head-dosimetry phantom, and the center location of the body-dosimetry phantom - is given in the accompanying documents for each selectable value of N  $\times$  T. When more than three different values of N  $\times$  T are available, the information is provided for at least the minimum, maximum and one mid-range value. The **dose profile** is presented on the same graph and to the same scale as the corresponding **sensitivity profile** required by 203.111.

Dose profile was measured for 32 rows by taking a stationary scan with the radio-chromic film, centered on top of the detector array. The scan protocol was as follows: 120 kV, 200 mA, 5 sec. After the scan was taken, the radio-chromic film was scanned, and the profile extracted. The following figure shows the dose profile and detector array.

Geometric efficiency was calculated as the ratio of the detector array to FWHM of dose profile using the following formula:

$$Geom_{Efficiency} = \frac{N_{rows} \cdot w_z \cdot M_f}{FWHM}$$

where  $N_{rows}$  is the number of detector rows;  $w_z$  is the width of the detector in the z-direction;  $M_f$  is the magnification factor; FWHM is the full width at half maximum of the profile. Table 47 on page 251 lists the measured geometric efficiencies for the two existing collimations of the BodyTom Elite.

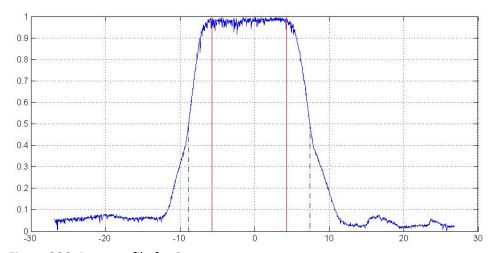


Figure 226: Dose profile for 8 rows

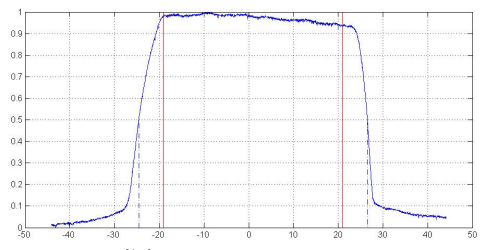


Figure 227: Dose profile for 32 rows

Table 47: The geometric efficiency of the two different collimations of the BodyTom Elite

Collimation	Geometric Efficiency
32 rows collimation	83 ± 5%
8 rows collimation	60 ± 5%

## **Chapter 8** Patient Registration

**Patient Registration** is the first step in the patient scan process. You can register a patient in the following ways:

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

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



Figure 228: Activated Patient Registration tab

#### **Navigating the Patient Registration screen**

Make sure the Patient Registration tab is selected.

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

Table 48: Patient Registration buttons

Patient Registration button	Action
Query	Searches the <b>HIS/RIS</b> server for scheduled patients. The population of patient information could take several minutes to appear, depending on the number of patient entries the query retrieves after clicking the <b>Query</b> button.
Cancel	Cancels the current query. Entries retrieved prior to cancellation appear in the <b>Query Results</b> list and Stored <b>Results</b> if they are moved there.
Register	Registers the selected patient and takes you to the <b>Acquisition</b> tab to select a protocol to be used for scanning.

Patient Registration button	Action
View	Shows selected patient details.
Search	Searches queried patient entries for specific information.
Store	Selects patient(s) from query results and moves them into the <b>Stored Results</b> list.
Delete	Removes patient(s) from the <b>Stored Results</b> list.
Manual	Manually enters a new patient and, when completed, takes you to the <b>Acquisition</b> tab to select a protocol to be used for scanning.

# Registering the patient

The following procedures show you how to register or enter patient information into the system before scanning a patient. Patients are registered manually or queried from the Hospital Information System/Radiology Information System (HIS/RIS). The system can be configured to add or create specific patient information when the patient is registered.

## Querying patient information

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



Figure 229: Patient Registration tab

2. Click the **Query** button at the bottom of the screen. The **Query Information** dialog box appears.

Figure 230: Query Information dialog box

3. Click the **HIS/RIS Server** dropdown and select the worksite to pull data from.

The default worksite appears at the top. If there is no list, see your site administrator to set it up.

4. Double-click any of the named Query Fields you would like to use to query for patients by entering the value in the Value column. A popup associated with the Query Field you selected appears. For example, if you double-click the Scheduled Start Date row, the Edit Value popup appears. Enter the desired start date. Another example would be to click the Patient Name value row. Again, the Edit Value popup appears; however, this time Patient Name text boxes are provided so you can type the patient's name to query. You can click any of the Value rows to fill in data to help query for the patient you are looking for. You can enter as much or as little information as needed. If no information is available, leave the value blank.



Figure 231: Edit Value popup for name

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

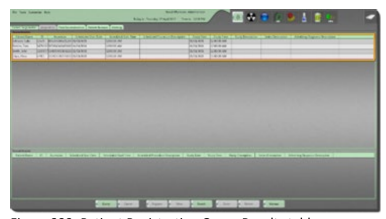


Figure 232: Patient Registration Query Results table

- 7. Select a patient and click the **Register** button to register the patient for the exam.
  - The system enables and opens the **Acquisition** tab. To perform the acquisition steps, see "Performing a scan" on page 264.

### Storing patients in the Stored Results list

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

- 1. If necessary, go to the **Patient Registration** tab to query the patients(s).
- 2. Perform steps 2 through 5 in "Querying patient information" on page 253.
- 3. Click 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, click anywhere in the patient's row.
  - To select more than one patient at a time, press and hold the Ctrl key and click patient entries until finished and release the Ctrl key.
  - To select all the patients, press and hold the **Shift** key, click the first patient in the list, then click the last patient to highlight all patients between the first patient selected and the last.
- 5. Click the **Store** button.

The patient information you selected appear in the **Stored Results** list at the bottom of **Patient Registration**.



Figure 233: Patient Registration Stored Results table

- 6. Click the patient you want to select from the **Stored Results** table.
- 7. Click the **Register** button to register the patient for the exam. The system enables and opens the **Acquisition** tab. To perform the acquisition steps, see "Performing a scan" on page 264.

### Manually registering a patient

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 **Patient Registration** tab.



Figure 234: Patient Registration tab

2. Click the **Manual** button at the bottom of **Patient Registration**. The **Exam Information** dialog box appears with the **Patient** tab open.

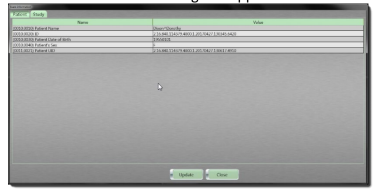


Figure 235: Exam Information dialog box

- 3. For **Patient Name**, double-click the **Patient Name** value. The **Edit Value** dialog box appears with patient name fields.
- 4. Enter patient name information in the fields provided and click one of the following buttons:
  - Click the **Update** button to save your entries and close the **Edit** Value dialog box.
  - Click the Close button to close the Edit Value dialog box without saving your work.

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.

Note The Patient ID is automatically generated by the system; you can replace this automated identifier with your patient's hospital ID number.



Figure 236: Patient ID field

- 5. For the **Patient Date of Birth**, perform the following:
  - Double-click the Patient Date of Birth field.
  - Enter the patient's birth date in the Patient Date of Birth field.
     Be sure to move the cursor to the far left to ensure two digits are included for the month and the day; four digits are required for the year.
  - Perform one of the following:
    - Click the **Update** button to save your work and close the **Edit** Value dialog box.
    - Click the Close button to close the Edit Value dialog box without saving your work.
- 6. For the **Patient's Sex**, perform the following:
  - Double-click the Patient's Sex field.
  - Enter the patient sex in the field by entering the appropriate letter:
    - o **F** for Female
    - o **M** for Male
    - o O for Other
  - Perform one of the following:
    - Click the **Update** button to save your work and close the **Edit Value** dialog box.

- Click the Close button to close the Edit Value dialog box without saving your work.
- 7. Perform one of the following:
  - Click the Register button to register your patient data.
  - Click the **Cancel** button to exit without entering your data.

When you click the **Register** button, the system enables and opens the **Acquisition** tab.

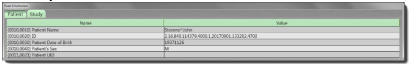


Figure 237: Patient data filled in

After your patient is registered, you can view the **Patient Exam Details** to ensure your data is correct.

If it is not correct, go to the next step to make the necessary changes.

8. Click the **Expand** link.

The **Exam Information** popup appears.

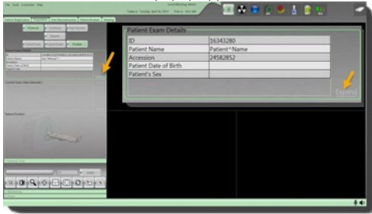


Figure 238: Expand link in context and close up

9. Make your changes in the **Exam Information** popup.

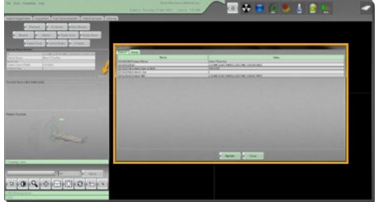


Figure 239: Exam Information popup

10. Click the **Update** button to save changes.

# Viewing patient information

This procedure lets you view, but not change, the patient information.

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



Figure 240: Patient Registration tab

- 2. Select a patient from the **Query Results** list or the **Stored Results** list.
- 3. Click the **View** button.
- Review the patient's information.
   This popup presents static information that you cannot change.
- 5. Click the **Close** button to exit the **View Entry Information** popup.

# Deleting patients from the Stored Result list

Patient information can be manually deleted from the **Stored Results** list, you cannot delete patients from the **Query Results** list.

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



Figure 241: Patient Registration tab

- 2. Select one or more patients from the **Stored Results** list to delete. Select patients in the following ways:
  - To select one patient, click anywhere in the patient's row.
  - To select more than one patient at a time, press and hold the Ctrl
    key and click patient entries until finished and release the Ctrl
    key.
  - To select all the patients, press and hold the **Shift** key, click the first patient in the list, then click the last patient to highlight all patients between the first patient selected and the last.
- Click the **Delete** button.The patients you selected are removed from **Stored Results** list.

#### Chapter 9 **Patient Scanning**

After you register the patient, the **Acquisition** tab automatically opens. The **Acquisition** tab lets you check that the selected patient information is accurate before you perform the scan. The Acquisition tab is also where you select protocols for the scan before you scan the patient. A protocol determines the parameters used to acquire patient images.



Figure 242: Active Acquisition tab

After the protocol is selected, you can scan the patient. See "Performing a scan" on page 264.

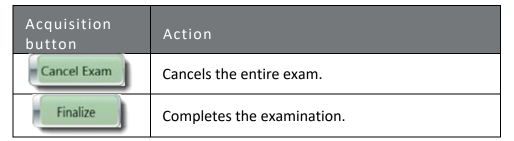
The following table provides information on the buttons on the **Acquisition** tab and what they are used for. Later you will learn how to set protocols for the scan.



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

Table 49: Acquisition buttons

Tuble 43. Acquisition buttons		
Acquisition button	Action	
Protocol	Allows you to modify the protocol selected or choose a new protocol.	
Continue	Authorizes the scanner to move to the next step if applicable.	
Start Recons	Begins any post-reconstructions that were defined during the protocol setup.	
Repeat	Allows you to repeat a portion or all the scan.	
Extend	Allows you to extend the currently active scan.	
Pause Scan	Allows you to pause the scan acquisition.	
Pause Exam	Allows you to pause the entire multi-step protocol acquisition.	
Cancel Scan	Cancels the current scan within a protocol.	



The following shows what appears in **Acquisition**:



Figure 243: What appears on Acquisition

### **Identifying Scan Types**

Scan types identify how images are acquired during a scan. The following Scan types are available.

#### **Axial**

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

#### Helical

The **Helical** scan type acquires data continuously as the x-ray tube rotates around the patient and the scanner translates over the patient in the Z axis.

### **Dynamic**

The **Dynamic** scan type acquires data at multiple time points over the same anatomic location while the scanner remains stationary; x-ray exposure can be continuous or intermittent.

### Reference

The **Reference** scan type acquires 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** and **Dynamic** scanning during a **CT Angiography** (**CTA**) or **Perfusion** protocol.

#### Scout

The **Scout** scan type acquires data continuously as the x-ray tube remains stationary at a designated angle and the scanner translates over the patient in the Z axis. The resulting **2D** projection is used during scan planning.

### Performing a 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, centipede movement, and gantry rotation immediately.

For a controlled stop, press the Cancel Scan button.

Note Be sure the scanner is calibrated for the room you will scan in. See "Selecting a room for the BodyTom Elite" on page 188 and "Performing a daily (air) calibration" on page 222.

- 1. From the workstation, go to the **Patient 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.
     See "Chapter 8 Patient Registration" on page 252.

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

- 2. From the **Acquisition** tab, click the **Protocol** button to open the **Exam Planner** dialog box.
- 3. Move scanner and align patient as needed.

See "Positioning the scanner before a scan" on page 96.

4. On the pendant, press the **Laser** button to turn on the laser and use it to align the patient to the scanner.

See "Positioning the patient using the laser lights" on page 97.

5. On the workstation screen, click the appropriate option:

Adult	To create and/or scan with adult scan protocols, which
	are stored by anatomical location.
Pediatric	To create and/or scan with pediatric scan protocols,
	which are stored by anatomical location.
Trauma	The <b>Trauma</b> orb can be used to store protocols
	commonly used for emergency scans.
Adult     Pediatric	By selecting either an <b>Adult</b> or <b>Pediatric</b> patient, the corresponding list of saved protocols becomes available.

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.

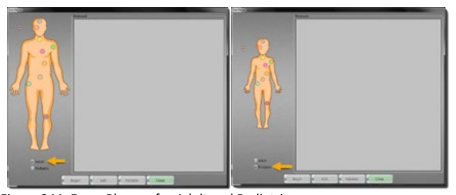


Figure 244: Exam Planner for Adult and Pediatric

6. Click the colored orb corresponding to the appropriate body part you will scan.

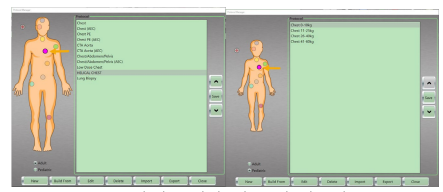


Figure 245: Anatomical orbs, with the Chest orb selected

- 7. Click the appropriate protocol from the list.
- 8. Click the **Edit** button to review the selected protocol.

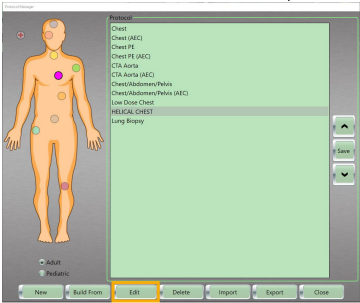


Figure 246: Protocol selected and Edit button active

The **Edit Protocol** dialog box appears.

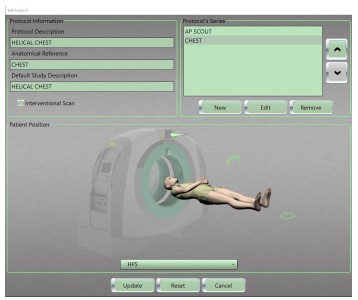


Figure 247: Edit Protocol dialog box

The **Protocol Information** tab displayed on the left and the Protocol's **Series** boxes displayed on the right show the series that are already created. The **Patient Position** appears identical whether it is for an adult, pediatric, or trauma patient.

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 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.

- 9. To edit an existing protocol, perform the following:
  - In the **Edit Protocol** dialog box, go to the **Protocol's Series** list and select the series to modify.
  - Click the Edit button.
     The Edit Series dialog box appears

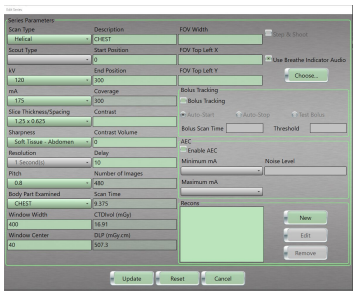


Figure 248: Edit Series dialog box

- Make desired changes. Select the **Update** button in the **Edit** Series dialog box.
- Alternatively, click the **Reset** button to remove any changes and return to the previous settings or click the **Cancel** button to return to the previous dialog box.
- 10. Click the **Update** button on the **Edit Protocol** dialog box.

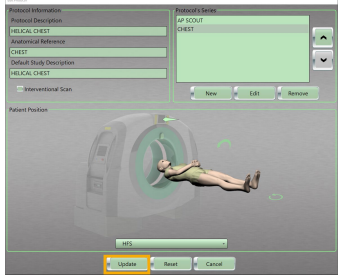


Figure 249: Update button

- 11. Click the **Begin** button from the **Exam Planner** dialog box.
- 12. When the **Is Scanner Properly Positioned?** popup appears, click the **Continue** button.



Figure 250: Is Scanner Properly Positioned? popup

Note If zero reference is not selected when starting your scan, the scanner considers the last known zero reference point to be the origin and start-point for the next scan. *Always* make sure to zero reference the scanner, when you set up a scan.

The system state orb will change color from yellow to green. The **System Ready to Scan** popup appears.



Figure 251: System Ready to Scan



**WARNING Do not** stand in either the forward or reverse paths of the scanner during the scan.

**Note** The scanner's side panels permit a low radiation exposure rate of <0.01mR/sec/100mAs – when x-ray is emitted.

The **START** button on the scanner control panel turns green when it is enabled.



Figure 252: Scanner control panel – START button

13. Press the **START** button on the scanner control panel to acquire your scan.

The pre-set scan delay countdown begins. The green light turns off when the **START** button is pressed.



Figure 253: Countdown popup

You can press **CANCEL** on the scanner to end the current scan operation. If pressed when lit, the system cancels the current scanning operation. If pressed during scanning, 1 current scan rotation, or 1 second, completes and then the scan is terminated. Alternatively, you can press the **Cancel Scan** button on the screen to cancel the entire scan or **Cancel Exam** button to cancel the entire exam.



Figure 254: Scanner control panel – CANCEL button

**Note** During the scan, observe the following:

Yellow lights on top of the scanner, and an audible beep identify that radiation is being emitted.

The patient's scan results appear; approximately one image per second.

When scanning begins, the **Continue**, **Repeat**, **Extend**, **Pause Scan**, **Pause Exam**, and **Cancel** buttons are enabled.

When you click the **CANCEL** button, the message "Scan is terminated" appears on both scanner and workstation.

14. If applicable, set your parameters and **Field Of View (FOV)** on your scouts.

**Note FOV** can only be adjusted when two scouts are acquired.



Figure 255: Scouts and FOV button

Scan coverage can be modified by selecting the drag boxes and adjusting the lines and can be centered by clicking on the small green circle and dragging the plan box.

15. Click **Continue** to proceed with your planned scan.



Figure 256: Continue button

The **Pending Scanning Movement** popup appears.



Figure 257: Pending Scanner Movement popup message

- 16. Click the **Continue** button to scan.

  Click the **Cancel** button to cancel the scan.
- 17. The **System Ready to Scan** popup appears.

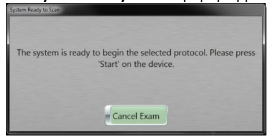


Figure 258: System Ready to Scan popup message – System is ready to begin scan

18. The **START** button on the scanner control panel turns green when it is enabled.



Figure 259: Scanner control panel – START button

19. Press the **START** button on the scanner control panel to acquire your scan.

The pre-set scan delay countdown begins. The green light turns off when the **START** button is pressed.

- 20. If the **Perform Reconstructions** popup appears, do one of the following:
  - Click the Yes button to perform post reconstructions now.
  - Click the **No** button to pause the reconstructions until a later time. When ready, click the Start Recons button.

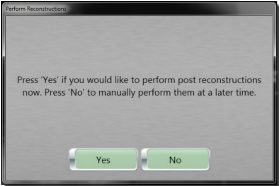


Figure 260: Perform Reconstructions popup message – To perform post reconstructions

- 21. Use the **Viewing** tools to review the scan.

  See "Examining the scanned image with tools" on page 299.
- 22. Click the **Finalize** button when finished.

  The dose report if **Show Dose Report** in **System Configuration** is enabled appears. In addition, the examination details are saved.

**Note** You must press the **Finalize** button before you can send the patient's data to **PACS**.



Figure 261: Dose report

## Repeating an image

The **Repeat** function can be used to repeat a scan if necessary. The entire scan 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, click the **Repeat** button. The **Protocol Viewer** dialog box appears.



Figure 262: Protocol Viewer – Start Position and End Position

- 2. Review the protocol parameters.
- 3. Click the **Repeat** button from the **Protocol Viewer** dialog box.
- 4. The **Repeat Protocol** popup appears.



Figure 263: Repeat Protocol popup

**Note** You can change the start and end position or use what appears.

- Click the Repeat button on the Repeat Protocol popup.
   Scout lines appear in blue, which indicates the second scan and modifications of the start and end points if made.
   The scanner will move to the start position.
- 6. Press the **START** button on the scanner's control panel to begin the repeat scan.

# **Extending** a scan

The **Extend** function can be used to add additional scan length to **Axial** scans only.

**Note** If scouts have been acquired, **Extend** can only be used within the boundaries of the Scout. If scouts are not acquired, there are no limits to extending the scan.

1. While the **Acquisition** tab remains active, click the **Extend** button.



#### The **Protocol Viewer** dialog box appears.

Figure 264: Protocol Viewer dialog box

- 2. Review the protocol parameters
- Click the Extend button.The Extend Protocol popup appears.



Figure 265: Extend Protocol popup

Number of Millimeters

20

Extend Cancel

4. Enter the length of the extension in the **Number of Millimeters** text box. Length must be in 10mm increment.

Figure 266: Extend Protocol popup

- 5. Click the Extend button.
- 6. Press the **Start** button on the scanner's control panel to perform the **Extended** scan.

### Scanning with special features

The following features are available for use in protocols.

# Using the step-and-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. Click the **Step & Shoot** option in the **Edit Series** dialog box.

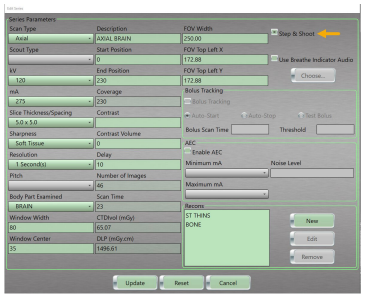


Figure 267: Step & Shoot option in the Edit Series dialog box

- 3. Click the **Update** button in the **Edit Series** dialog box.
- 4. Click the **Update** button in the **Edit Protocol** dialog box.
- 5. Click the **Begin** button in the **Exam Planner** dialog box to begin the scan.

The system state orb will change color from yellow to green. The **System Ready to Scan** popup appears.



Figure 268: System Ready to Scan popup

6. To continue the scan, go to the scanner and press the **Scan** button on the screen.

The first set of images are acquired at this position.

The **Step & Shoot** popup appears for you to control the next acquisition.



Figure 269: Step & Shoot popup

- Click the **Shoot** button to start the scan.To cancel the scan, click the **Cancel** button.
- 8. Continue for the length of the scan.
- 9. Click the **Finalize** button when finished.

### Performing a scan with Automatic Exposure Control

**Note** Depending on system's configuration, not all functions may be available to perform this procedure.

**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 adaption of mA based on user specified image quality and attenuation characteristics of the scanned body region.

Scout scans provide a graph of mA values based on object density and desired noise level. Axial or Helical scans 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<sub>vol</sub> and DLP values. It tends to increase CTDI<sub>vol</sub> for large patients and decrease it for small patients relative to a reference patient size.

**Note** Ensure patient is accurately centered in the 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 following Sharpness options:

Helical Soft Tissue - Head Helical Soft Tissue - Abdomen Axial Soft Tissue Axial Pos. Fossa/Vessel Axial Sharp

1. Under AEC, click the Enable AEC option.

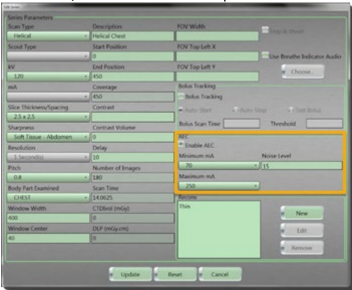


Figure 270: Edit Series dialog box with AEC options selected

- 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 range is 30 to 300.
- 4. Select the **Noise Level** to set the standard deviation of noise value for the acquired images. The noise range is 1-200.

**Note** Consult with the site's physicist for guidance specific to the department.

- 5. Click the **Update** button in the **Edit Series** dialog box.
- 6. Click the **Update** button in the **Edit Protocol** dialog box.
- 7. Make sure your patient and scanner are properly positioned.
- 8. Click the **Begin** button to begin the scan.
- 9. Press the **START** button on the scanner control panel.
- 10. After the scouts are acquired and the scan region is set, click the **AEC** tab.
- 11. Click the **Toggle Graph** button to view the graph on the scout.

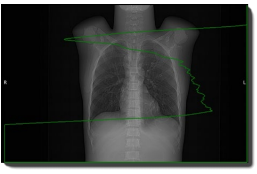


Figure 271: Toggle Graph button

The graphs will now appear on the scout(s). Review the mA modulation to ensure it meets your clinical needs.



Figure 272: Graphs on the scout(s)



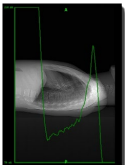


Figure 273: AEC modulation graph

- 12. To return to the scout parameter view, click the **Toggle Graph** button, again.
- 13. If changes to the mA or Noise levels are required, you can modify the Minimum, and Maximum mA and noise as needed on the image.



Figure 274: Minimum mA and maximum mA; noise level

- 14. When the desired level is achieved according to your department policy, click the **Continue** button to start the scan.
- 15. Press the **START** button from the scanner.

**Note** While reviewing the scan you will see mA modulation as per your graphs.

16. Click the Finalize button.

# 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.

Table 50: Bolus tracking parameters and tools

Option	Description
Bolus Scan Time	The amount of time allowed to monitor the bolus.
Auto-Start	Begins the scan after the specified bolus scan time if no bolus is detected.
O Auto-Stop	Stops the scan after the specified bolus scan time if no contrast is detected.

Option	Description
O Test Bolus	A small amount of contrast is injected, and a timing graph is displayed after specified bolus scan time.
Threshold	Hounsfield Unit detection at the area being monitored – ROI.
Use Calculated	When performing the test bolus, click the <b>Use Calculated</b> option to use the bolus timing calculated from the test bolus scan.
O Use Manual	Allows you to select a manual timing of bolus after the test bolus has calculated a timing.

- 1. Perform steps 1 through 8 in "Performing a scan" on page 2644.
- 2. Click the **Bolus Tracking** option and set parameters, such as **Auto-Start**, **Auto-Stop**, **Bolus Scan Time**, and **Threshold**.

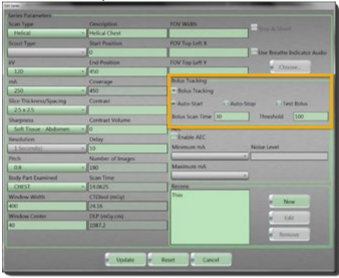


Figure 275: Bolus Tracking option

- 3. Click the **Update** button in the **Edit Series** dialog box.
- 4. Click the **Update** button in the **Edit Protocol** dialog box.
- 5. Click the **Begin** button to acquire the scout(s).
- 6. Acquire the scout(s).
- 7. To move the scout or reference line go to the **Viewing** tools and select **Active Scan Region**.

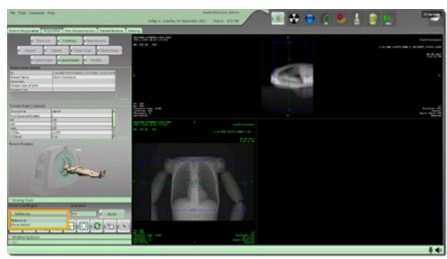


Figure 276: Active Scan Region – Bolus Reference or Helical CTA

Note If the protocol contains one or more scouts, the system places the scan region for each series on the scout based on start and end positions that were entered for each series in the protocol. The Active Scan Region dropdown populates the data with the ID that corresponds to each region. You can adjust the scan region prior to continuing the exam, as described below.

8. On the scout, use the Blue lines to plan the helical scan.

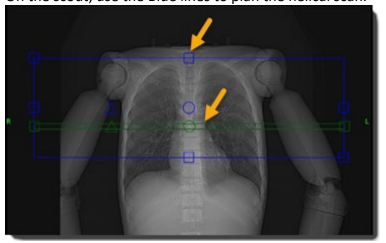


Figure 277: Scout line (blue) and Reference line (green)

Note The distance between the Reference and Bolus Tracking scans cannot exceed 100mm.

- 9. Position the green **Reference** line at the desired anatomical location.
- 10. Click the **Continue** button.

  The scanner will move to the reference line noted on the scout.
- 11. Go to the scanner and press the **START** button on the scanner control panel.
- 12. Click the **ROI** tool and draw the **ROI** on the **Reference** image.

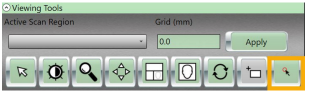


Figure 278: Bolus ROI tool

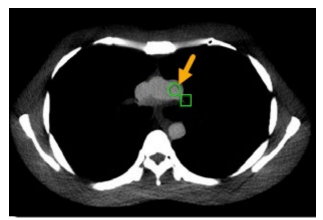


Figure 279: ROI on the Reference scan

- 13. Click Continue.
- 14. Load the injector and set your desired flow and rate and arm the injector.
- 15. When the scanner is ready, press **START** on the scanner and manually start the injector at the same time.
- 16. The Helical scan will trigger when the threshold value set for the ROI is exceeded.

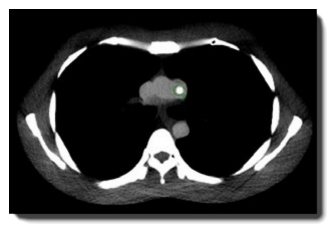


Figure 280: Scan triggers when bolus enters reference point/ROI

17. Review the completed scan.

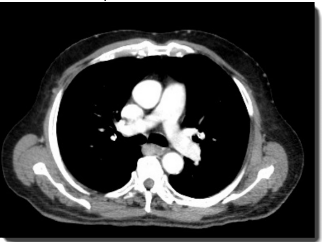


Figure 281: Scan at peak enhancement

18. Press the **Finalize** button when complete.

# **Performing Test Bolus**

**Test Bolus** tests the timing of the injected contrast.

A small amount of contrast is injected, and a timing graph is displayed after the specified bolus scan time. When the contrast is detected, the system stops scanning and a report on the recommended delay-time for CTA protocols appear.

- 1. Perform steps 1 through 8 in "Performing a scan" on page 2644.
- 2. Click the **Bolus Tracking** option, click the **Test Bolus** option and set parameters.

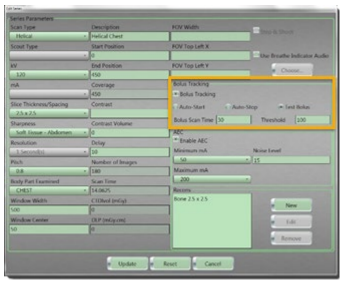


Figure 282: Test Bolus option

- 3. Click the **Update** button in the **Edit Series** dialog box.
- 4. Click the **Update** button in the **Edit Protocol** dialog box.
- 5. Click the **Begin** button to acquire the scout(s).
- 6. Acquire the scouts(s).
- 7. To move the scout or reference line go to the **Viewing** tools and select **Active Scan Region**.



Figure 283: Active scan region

8. On the scout, use the blue lines to plan the helical scan.

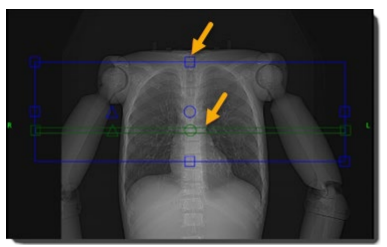


Figure 284: Scan planning lines

- 9. Position the green **Reference** line at the desired anatomical location.
- 10. Click the **Continue** button.

  The scanner will move to the reference line noted on the scout.
- 11. Go to the scanner and press the **START** button on the scanner control panel.

The reference image will be scanned and displayed.

12. Click the ROI tool and draw the ROI on the Reference image.

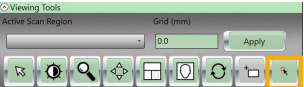


Figure 285: Bolus ROI

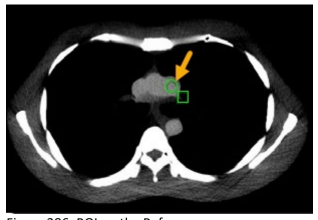


Figure 286: ROI on the Reference scan

13. Click Continue button.

- 14. Load the injector and set your desired flow and rate and arm the injector.
- 15. When the scanner is ready, press **START** on the scanner and manually start the injector simultaneously.
- 16. The **Bolus Timing graph** appears and shows the calculated, bolustracking time.

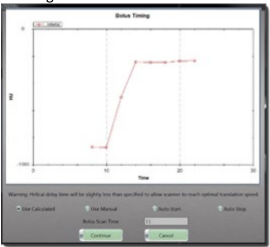


Figure 287: Bolus timing graph

17. Select from one of the following:

Use	Uses the bolus timing calculated from the <b>Test</b>
Calculated	Bolus scan.
Use Manual	Allows you to manually set the delay time prior to
	the start of the helical scan.
Auto Start	Begins the scan after the specified bolus scan time
	if no bolus is detected.
Ata Ctan	Stops the scan after the specified bolus scan time
Auto Stop	if no contrast is detected.

- 18. Click the **Continue** button. Review completed scan.
- 19. Press the **Finalize** button when complete.

## **Performing a CT Perfusion Scan**

CT perfusion (CTP) is a technique used to evaluate cerebral perfusion of the level of blood flow in the brain, by monitoring the initial phase of iodinated contrast media through the vasculature of the brain.

1. Perform steps 1 through 8 in "Performing a scan" on page 2644.

Scan Type FOV Width Description PERFUSION Dynamic FOV Top Left X Scout Type Start Position Use Breathe Indicator Audio FOV Top Left Y Choose... 40 80 Bolus Tracking mA Coverage 40 Bolus Tracking 50 Slice Thickness/Spacing Contrast 10.0 x 10.0 Bolus Scan Time Threshold Contrast Volume Sharpness Soft Tissue Resolution Delay Minimum mA Number of Images Pitch Maximum mA 240 Body Part Examined Scan Time HEAD 60 Window Width CTDIvol (mGy) New 135 DLP (mGy.cm) Edit Update Reset

2. After selecting CTP protocol, review the Dynamic CTP options including Scan Time and make your selections.

Figure 288: Edit Series CTP Scan Time

- 3. Click the **Update** button in the **Edit Series** dialog box.
- 4. Click the **Update** button in the **Edit Protocol** dialog box.
- 5. Click the **Begin** button to acquire the scout(s).
- 6. Perform scout and set Dynamic CTP scan location.
- 7. To move the Dynamic CTP location use the green circle inside the reference line.
- Click the Continue button.
   The scanner will move to the reference line noted on the scout.
- 9. Load the injector and set your desired flow rate. When the scanner is ready, press **Start** on the scanner and manually start the injector at the same time.
- 10. Review completed scan.

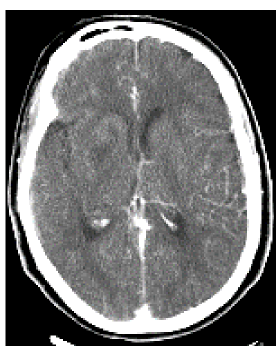


Figure 289: Brain Perfusion Image

#### 11. Select Finalize.

## Calculating and creating perfusion maps

- 1. Select a perfusion patient from **Patient Browser** and select the series to view.
- 2. To open the image, click the **View Images** button or double-click the series.

The **Viewing** tab is enabled, and the **CTP** tab automatically opens.

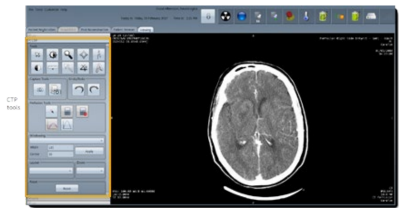


Figure 290: CTP tools

Table 51 CTP Tools

CTP Viewer tools		
9	Perfusion Artery/Vein Selection	Select to place the arterial and venous ROIs on the images.
	Calculate CBF, CBV, MTT Map	Select to calculate the CT Perfusion maps.
	Clear Perfusion Map	Cancels the calculations and returns to <b>Calculation</b> mode.
	Show Artery/Vein Flow Graph	Displays the Arterial Venous Flow graph.
	Peak Image	Displays the image that has the highest HU value based on the arterial ROI placement.

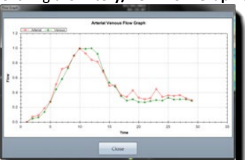
- 3. Click the **Perfusion Artery/Vein Selection** tool.
- 4. Place an Arterial ROI.
- 5. Place a Venous ROI.
- 6. Click the **Calculate CBF, CBV, MTT Map** tool.
- 7. The Perfusion Maps are calculated and displayed:



Figure 291: Perfusion maps

The calculations produce three maps and the perfusion image:

Perfusion Images	Top, left corner
Cerebral Blood Flow (CBF)	Top, right corner
Cerebral Blood Volume (CBV)	Bottom, left corner
Mean Transit Time (MTT)	Bottom, right corner



8. Clicking the **Artery/Vein Flow Graph** displays the following:

Figure 292: Arterial Venous Flow

This graph displays arterial and venous flow rates with respect to time.

- Click the Peak Image tool to display the peak image.
   The peak image displays the image that has the highest HU value based on the arterial ROI placement.
- 10. Click the **Clear Perfusion Maps** tool to cancel calculations and return to **Calculation** mode.
- 11. Click the **Reset** button to reset images back to the original setting(s). You cannot undo this action.

#### Using the Interventional Package

The Interventional Package is designed to make Interventional procedures quick and easy for the technologist and physician. When activated the gantry will spin continuously to make the transition to a scan as fast as possible.

When enabled the **Interventional Tab** will be displayed as seen below.



Figure 293: Interventional Tab

When activated the **Interventional Tab** provides a streamlined workflow specifically for interventional cases.

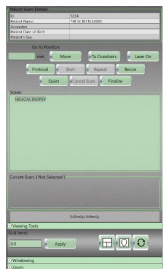


Figure 294: Interventional Tab - Patient exam details

The Interventional Tab includes the following options:

Option	Description Description
Go To Position:  mm Move	Allows a typed in value, when Move is clicked the scanner goes to that location
To Crosshairs	Moves the scanner to the Laser location
Laser On	Toggles the laser lights on or off
Protocol	Allows protocol parameters for the Interventional procedure to be modified
Start	Initiates X-Ray when the user is ready
Repeat	Repeats the last scanned protocol
Recon	Allows a selected raw data set to be Post Reconstructed without leaving the Interventional workflow
Quiet	Stops the gantry rotation
Cancel Scan	Cancels the currently planned scan
Finalize	Ends the current examination

The tab also includes a **Scan Tree, Current Scan,** and an updated **Dose Gauge.** 

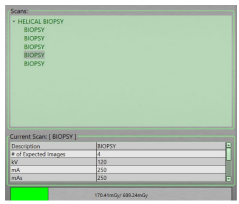


Figure 295: Scan Tree, Current Scan, and Updated Dose Gauge

The **Scan Tree** allows access to acquired scans for quick image loading or repeat scanning using those parameters.

- When a series in the Scan Tree is selected, the images associated with it will be displayed, allowing them to be reviewed to ensure the proper start and end locations as well as the thickness and spacing are selected for repeat scans.
- Clicking any of the scans in the Scan Tree, then clicking 'Repeat', moves the scanner to the Start location from the selected series and automatically initiates scanning.

The **Current Scan** box shows the protocol parameters used for the most recent active scan.

The **Dose Gauge** displays the accumulated CTDI and DLP in mGy for the current procedure and is updated each time a new scan is initiated.

**Viewing Tools, Windowing, and Zoom** options are available by clicking the appropriate line below the Dose Gauge.

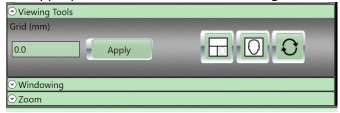


Figure 296: Viewing Tools, Windowing, and Zoom options

A right click menu on the active image, also allows access to a full array of viewing and measuring tools.



Figure 297: Interventional Workflow

#### **Interventional Workflow**

Any protocol can be used as an Interventional protocol.

- 1. Register a patient
- 2. Click 'Protocol'
- 3. Select the desired protocol
- 4. Ensure the 'Interventional Scan' box is selected.

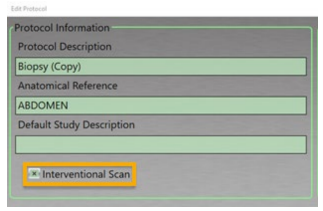


Figure 298: Interventional Workflow - Protocol Information dialogue box

- 5. Click 'Begin' to accept the protocol and perform the scout scan(s).
- 6. Plan your scan over the region of the procedure and acquire that scan.
- 7. Once the scan has been acquired the system will automatically move to the Interventional tab and show the following:



Figure 299: Interventional Workflow - Scan acquired

8. Review the acquired images from the previous scan to determine the start and end locations of the Interventional scan.



Figure 300: Modify protocol parameters

- 9. Click **'Protocol'** to modify the protocol parameters for the Interventional procedure.
- 10. Once parameters are selected click 'Begin'
  The scanner will move to the selected Start position.
- 11. Click the 'Start' button to initiate the scans set in the protocol.

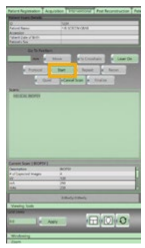


Figure 301: Initiate Scans - Interventional protocol

- 12. When satisfied the anatomy of interest is included in the images, you can enter a location in the 'Go To Position' box and click 'Move'.
  - This moves the scanner out of the way to allow the physician access to the patient.



Figure 302: Move the Scanner

- 13. When ready to perform another scan, click 'Repeat'
  - The scanner will move back to the previously defined start location and perform additional scans as defined in the protocol.

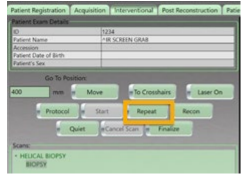


Figure 303: Repeat Scans - Interventional protocol

- 14. You can now use the 'Move' and 'Repeat' options to move the scanner back and forth to allow the physician access to the patient and perform repeat imaging to confirm devise position in the patient.
- 15. If changes to the protocol are necessary, click 'Protocol'.
- 16. Modify the parameters to the desired values and click 'Begin'.
- 17. You must then click 'Start' to initiate the new scan.

#### **Examining the scanned image with tools**

The image tools can only be used when the **Acquisition** tab is enabled and an image is displayed, or when images are loaded to the viewer from the Patient Browser tab.

From the **Acquisition tab**, you can zoom, pan, modify window width, and 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 55: 2D, MPR, 3D, and CTP image tools.

Table 52: Image tools

Image tool	Tool name	Action
<b>⊠</b>	Clear Tool	Resets the tool to the default pointer.
<b>(A)</b>	Window Width/Center	Adjusts window width and center of image.
Q	Zoom	Magnifies the image.
	Pan	Adjusts image on X or Y axis
	Toggle Scouts	Display's or removes scouts from Acquisition.

Image tool	Tool name	Action
	Toggle Layout	Changes the layout to 2x2. Repeat process to return to 1x1.
+0	Scan Region Re-Draw	If scout lines and the scan region is deactivated, allows you to reactivate.

# **Chapter 10** Patient Browser

The **Patient Browser** lets you view patient information and images associated with the patient information after the patient's scan.



Figure 304: Active Patient Browser tab

The following table identifies the buttons found on the **Patient Browser**.

Table 53: Command buttons

Button	Action
Archive	Selects the archive destination for selected information.
Import	Imports exam information from <b>PACS</b> or <b>Media</b> .
Delete	Deletes the selected exam information from the <b>Patient Browser</b> tab.
Register	Reregisters a patient who is already in the system.
Build Dose	Generates a Dose Report and Dose SR for the selected patient.
Merge	Combines two different image sets.
Show Info	Shows patient, study, series, and image information.
View Images	Displays selected patient images in Viewing.
Compare	Allows you to compare two series.

## **Navigating the Patient Browser**

The **Patient Browser** lets you perform tasks on existing series, for example archiving and previewing the series. This section will introduce you to **Patient Browser** and identify the symbols, areas, and buttons you can use.

The **Patient Browser** can be broken down into the following sections:

- Exam table
- Series table
- Selected protocol table
- Patient Browser's active buttons
- · Preview window



Figure 305: Patient Browser sections

# **Identifying symbols on Patient Browser**

Symbols may appear next to series in the exam and/or series tables. These symbols are more vivid when active; they are identified below as active symbols.

Locked 6	Indicates the series is locked and cannot be deleted.
	Indicates the series is marked to be read or read by the physician.
PACS 🗸	Indicates the series has been sent to <b>PACS.</b>
	Indicates the series has been sent to <b>PACS</b> ; the archived series appear below the initial table
and the second s	Indicates the series has been sent to a media device, for example USB.

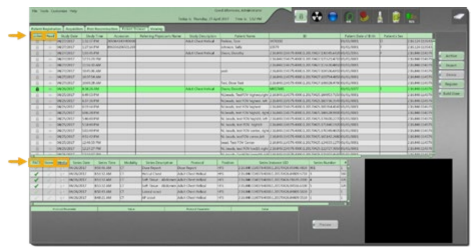


Figure 306: Patient Browser locked, read, PACS and Stored (archived), and media symbols

# Using the vertical and horizontal scroll bars on Patient Browser

Navigation scroll bars let you move the lists in the Patient Browser sections up, down, and horizontally to view all available exam information.

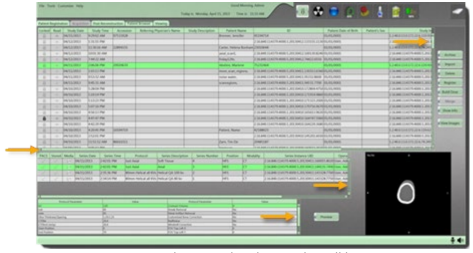


Figure 307: Patient Browser horizontal and vertical scroll bars

# Locking a study

- 1. Click the Patient Browser tab.
- 2. Select the study to lock.

- 3. Right-click the mouse button.
- 4. Click **Lock** on the floating menu.



Figure 308: Floating menu - Lock

The **Lock** symbol appears for any selected series.



Figure 309: A locked series

A series cannot be deleted while in the **Lock** mode.

5. To unlock right-click and click **Unlock**.

**Note** All **QA** series are locked to prevent deletion. The **QA** series can only be unlocked by your field service representative.

## Marking a series to read

- 1. Click the **Patient Browser** tab.
- 2. Select the study to mark.
- 3. Right-click the mouse button.

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4. Click Mark on the floating menu.

Figure 310: Floating menu - Mark

The **Mark** symbol appears for any selected series.

5. To unmark any series, right-click **Unmark**.

# Using the preview window

- 1. Click a patient in the exam table.
- 2. Click a series in the series table.
- 3. Click the Preview button to the right of the Series Protocol Table.

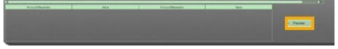


Figure 311: Preview Button

4. The selected series will appear in the **Preview** window.



Figure 312: The series appears in the preview window

#### **Archiving patient series**

You can archive patients and studies (or series) to **PACS**, media (USB or CD), or surgical navigation devices.

#### **Archiving to PACS**

- 1. Click the Patient Browser tab.
- 2. Select the patient study for **PACS** in the following way:
  - To select one patient and all associated series, click the patient, and click the **Archive** button.
  - To select specific series for a patient, press and hold the Ctrl key, then click each individual series from the Series table, and click the Archive button.

The Archive Destination popup appears.



Figure 313: Archive Destination popup

- Click the PACS button.The Archive to Server popup appears.
- 4. Click the **Select Archive Location** dropdown and select a site.



Figure 314: Archive To Server popup

- 5. Review the **Image Range** items to make sure all those items you selected in step 2 are captured.
- 6. Click the **Archive** button to begin the archive process.

If enabled the **Store/Print Queue** dialog box will appear to show the status of your image transfer. You can also activate the **Store/Print Queue** dialog box by clicking **Tools > Store/Print Queue** from the main menu.



Figure 315: Store/Print Queue dialog box

- 7. Watch the status of each series:
  - **Pending** informs you that the series is paused because you clicked the **Pause** button.
  - **Connecting** informs you that the series is in process of archiving to its targeted location.
  - Each series will move from the top portion of the popup to the bottom portion of the Store/Print Queue popup when it has been processed.
- 8. While the archiving is in process, you can perform one of the following from the buttons in the **Store Print Queue** dialog box.

Table 54: Store and Print Queue buttons

Store and Print Queue button	Action
Pause	When you select one or more series, temporarily stops the series from being stored. This is a toggle button with the <b>Resume</b> button.

Store and Print Queue button	Action
Delete	When you select one or more series, deletes either a series to be stored, or a series that failed to store.
Cancel	Stops the archive to USB or a drive.
Retry	When you select one of more series, tries to archive the selections.
Details	When you select one of more series, displays an explanation of why a series failed to store.
Close	Closes the <b>Store/Print Queue</b> popup.

- 9. 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.
- 10. If there are failed archived series; click the **Retry** button to attempt to archive the series you selected.

Note Any Storing Failure status appears in the bottom of the popup to inform you why the failure occurred. If an archive job fails, it will be sent to the Failed Store/Print Jobs list.

11. When the archiving is complete, click the **Close** button to exit the **Store/Print Queue**.

You can also click the **Close** button and the archiving process will continue as you do other tasks.

## **Archiving to Media**

- 1. Click the Patient Browser tab.
- 2. Select the patient study to archive following way:
  - To select one patient and all associated series, click the patient and click the **Archive** button.

 To select specific series for a patient, press and hold the Ctrl key, then click each individual series from the Series table, and click the Archive button.

The **Archive Destination** popup appears.



Figure 316: Archive Destination popup

- 3. If you are archiving a USB device, insert the USB drive into the USB port.
- 4. Click the Media button.

The Archive to Media popup appears.

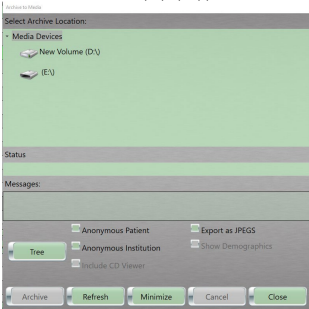


Figure 317: Archive to Media popup

- 5. Click the targeted drive and path destination.
- 6. The **Archive** button is active.



Figure 318: Archive Button active

7. Click the appropriate check boxes for your archive process:

Anonymous Patient	Makes the patient's information anonymous for HIPAA standards.
Anonymous Institution	Makes institutional information anonymous for HIPAA standards.
Include CD Viewer (requires CD viewer software installed)	Includes a CD viewer application to view images from the media.
Export as JPEGS	Exports image files in .JPG format.
Show Demographics	Includes the demographic information in archive if you clicked the Export as JPEGS check box.

- 8. Click the **Archive** button to begin the archive process.
  - The **Cancel** button is active after clicking the Archive button; click the **Cancel** button to stop the archive.
- 9. The **Archive to Media** dialog will update the status when archiving is complete.

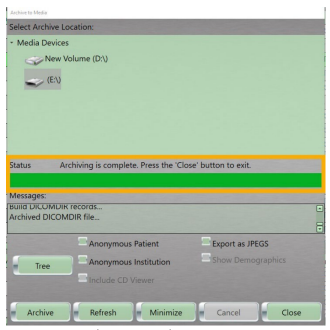


Figure 319: Archiving complete

- Click the Refresh button to remove any messages that appear in the Message box.
- Click the **Minimize** button to continue working in other areas while the archiving process runs.
  - A disk appears at the bottom; click it to maximize the Archive to Media popup.
- Click the Close button to exit the Archive to Media popup after the archive process is complete.

## **Archiving to Navigation**

- 1. Click the Patient Browser tab.
- 2. Select the patient study or series.
- 3. Click the **Archive** button.
- 4. Click the **Navigation** button.
- 5. Click the **Select Archive Location** dropdown and select the location.



Figure 320: Archive to Server popup

6. Under **Image Range**, select the image(s) you want to send to navigation.

To return to the default selections, click the **Reset** button.

- 7. Perform one of the following:
  - Click the **Archive** button to send the image to **Navigation**.
  - Click the Cancel button to return to Patient Browser.

#### **Import**

**Import** allows you to add patient images to the Patient Browser.

## **Importing from PACS**

- 1. Click the Patient Browser tab.
- 2. Click the **Import** button to import data. The **Import Location** popup appears.

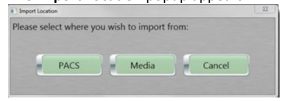


Figure 321: Import Location popup

Click the PACS button.The Import from PACS dialog box appears.

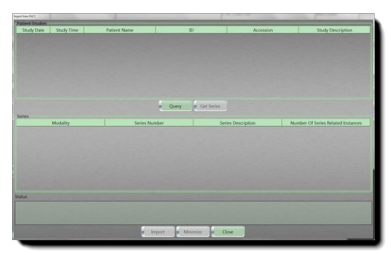


Figure 322: Import from PACS dialog box

4. Click the Query button.

The **Query Information** dialog box appears.



Figure 323: PACS Query Information dialog box

- 5. Perform the following:
  - Select a **HIS/RIS** server from the dropdown.
  - Set the values to search in your query.
  - Click one of the following buttons:
    - Click Query to save the search results
    - o Click **Reset** to clear the query information.
    - Click **Cancel** to exit the **Query Information** popup.
- 6. From the **Queried** results, select a patient and click the **Get Series** button.

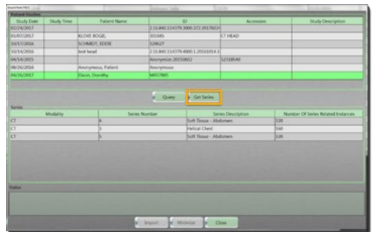


Figure 324: Import PACS dialog box with active Get Series button

- 7. Click the **Import** button.
- 8. Click the **Close** button to exit the **Import from PACS** dialog box.

## Importing from media

- 1. Click the **Patient Browser** tab.
- 2. Click the **Import** button on **Patient Browser**.

The **Import Location** popup appears.



Figure 325: Import Location popup

3. Click the **Media** button.
The **Import from Media** popup appears.

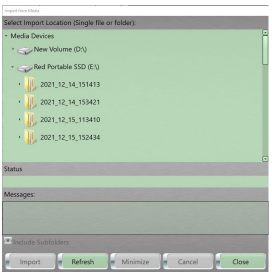


Figure 326: Import from Media popup

4. Click the drive and path where images were previously stored. The **Import** button is active.



Figure 327: Active Import button

- If necessary, click **Subfolders** to see the entire path.
- 5. Click the **Import** button.

The imported images appear in **Patient Browser**.

#### **Delete**

- 1. Click the Patient Browser tab.
- 2. Select the study or the series to delete.

3. Click the **Delete** button.

The **Confirm Deletion** popup appears.



Figure 328: Confirm Deletion popup

4. Click the **Delete** button on the **Confirm Deletion** popup. The patient data will be deleted from the **Patient Browser**.

#### Registering a patient from Patient Browser

If additional scans must be performed on a patient that is listed in the **Patient Browser**, you can register them by performing the following:

- 1. Click the Patient Browser tab.
- 2. Select the patient to register.
- 3. Click the **Register** button.



Figure 329: Patient browser register button

4. The Create New Study popup appears.

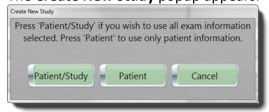


Figure 330: Create New Study popup

5. Perform one of the following:

- Click the Patient/Study button to use all exam information selected, including the accession number.
- Click the **Patient** button to use only patient information.
- Click the Cancel button to exit the Create New Study popup.

## **Building dose from Patient Browser**

The **Build Dose** button in the Patient Browser, allows you to manually create a Dose Report and Dose SR image which will appear in the Series table when completed.

- 1. Click the Patient Browser tab.
- 2. Select the patient to use for the **Build Dose**.
- 3. Click the Build Dose button.



Figure 331: Build dose button

4. The Build Dose Please Wait popup appears.

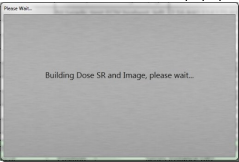


Figure 332: Please Wait popup

- 5. The Dose Report and Dose SR image will be saved to the Series Table.
- 6. If the **Dose Build Failed** popup appears, click the OK button, make the necessary changes, and try again.



Figure 333: Dose Build Failed popup

Note Dose Structured Reports (Dose SR) cannot be viewed in the BodyTom Elite system; Dose SR can be viewed in PACS with the appropriate viewer.

### Merge

The merge function is used to combine two different series for review.

Note Only series with same patient name, ID, slice thickness, slice spacing, kernel, and pixel spacing can be merged.

- 1. Click the **Patient Browser** tab.
- Select the study to merge.The study must include several scans.
- 3. Select two series to merge by pressing and holding the **Ctrl** key and highlighting the two series.

The **Merge** button appears and is active.

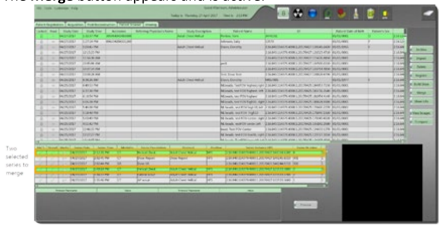


Figure 334: Two series selected to merge

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4. Click the Merge button to merge the series.

Figure 335: Active Merge button

- 5. The **Please Wait** popup appears while the application loads the series to be merged.
- 6. When the **Merge** window appears, the selected series are displayed side by side in separate windows.
- 7. The bottom 'filmstrip' is empty when the window first displays.



Figure 336: Image viewer without image(s) selected

- 8. Images can be drug from the series displays on the sides of the **Merge** window to the Image Viewers.
  - These images will be outlined in green in the series list.
    - Multiple images in the series lists can be selected by clicking each image.
    - You can also right click on the series lists and choose Select All.
  - To Deselect a single image, click the image, the green outline will change to white.
    - You can also right click on the series lists and choose **Deselect** All.

9. Drag the images you wish to merge from the series lists to the bottom 'filmstrip'.



Figure 337: Selecting images to move to image viewer or bottom filmstrip

10. To reorder the filmstrip, drag one or more images to the new location.

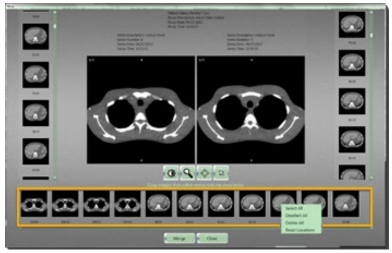


Figure 338: Finished filmstrip

11. When finished with the bottom strip, click Merge.

The application generates an additional series with the selected images. The **Series Save** popup appears indicating if the merge operation was successful.



Figure 339: Merge Series Saved popup

- 12. Click the **OK** button.
- 13. Click Close.

The merged series will display in the **Patient Browser** and will be marked as a merged series under the Series Description column.

## Using Show Info to view, update, and move a series

- 1. Click the Patient Browser tab.
- 2. Select the patient.
- Select the series.The **Show Info** button becomes active.
- 4. Click the **Show Info** button.

The View/Update Information dialog box appears.

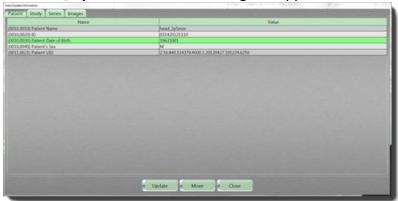


Figure 340: View/Update Information dialog box

Patient	Data about the patient, such as patient name,
	date of birth, and sex of patient.
Study	Data about the study, such as date, time, and
	referring physician.
Series	Data about the series, such as the position
	reference indicator, model, pixel-padding
	value, and series date and time.
Images	Data about the image, such as the instance
	number (sequential), exposure time.

- 5. Click the tab(s) to review and update the necessary information.
- 6. Double-click any editable field and make your change(s).
- 7. Click one of the following buttons
  - Click the **Update** button to save your changes.

**Note** If information is invalid, you are prompted to correct the information and click **Update** again.

If the field cannot be edited, a prompt appears tell you the selected field is not editable.

Click the Move button to show the Move Series popup.

Note The Move Series function is used when a scan has been acquired under the wrong patient file or to move a patient that was registered manually to the Patient Registration tab. Moving the patient to the Patient Registration tab will capture all the patient's information.

- Click the Registration button to move the patient into an existing patient or by manually creating a new patient using the Patient Registration process.
- Click the **Cancel** button to exit the **Move Series** popup.
- Click Browser to go to the Patient Browser tab and move the series.
- Click the **Move** button to confirm moving the series.
- Click the Cancel button to exit the Patient Browser tab and return to the View/Update Information popup.
- 8. Click the **Close** button to exit the **View/Update Information** popup.

Note An audit log of both old and new patient series, including the date and time of change and who performed it, is generated.

## Modifying a series scanned under the wrong patient

If a patient has been scanned under the wrong identification, the series can be corrected.

- 1. Click the Patient Browser tab.
- 2. Select the series that was scanned with incorrect patient identification to modify the data.
- Click the Show Info button.
   The View/Update Information dialog box appears.



Figure 341: View/Update Information dialog box

4. Click the **Move** button.

The **Move Series** popup appears, denoting where to retrieve patient information from.

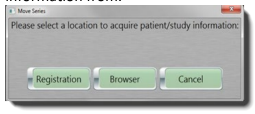


Figure 342: Move Series popup

The following defines what each batton performs.	
Registration	If patient information is stored within hospital's
	HIS/RIS server, click the Registration button,
	which will open the <b>Register Patient</b> tab to let you
	choose patient/study information.
Browser	If patient information is stored within system's
	browser, click <b>Browser</b> button, which show the
	Patient Browser tab information to let you select a
	series with correct patient information.
Cancel	Returns you to the previous dialog box.

The following defines what each button performs:

- 5. Perform one of the following:
  - If you clicked the **Registration** button in the previous step, go to the next step.
  - If you clicked the **Browser** button in the previous step, go to step 11.
- 6. Click the Manual button.

The **Exam Information** dialog box appears.

- 7. Enter the corrected data in any of the fields. See "Registering the patient" on page 253.
- 8. Click the **Update** button to save the change(s).
- 9. Click the **Move** button.

A prompt appears to review changes made to the patient and/or series information for changes to take effect.

- Click the **Ok** button and then the **Update** button.
   The corrected patient and moved data will appear in the **Patient** Browser.
- 11. If you selected the **Browser** button, the **Patient Browser** tab is showing; select the correct patient and series.
- 12. Click the Move button.

A prompt appears to review changes made to the patient and/or series information for changes to take effect.

- 13. Click the Ok button.
- 14. Review the patient to ensure it is the proper one.
- 15. Click the **Update** button.

16. Click the Cancel button to return to the Patient Browser.

#### Loading a series into view

- 1. Click the Patient Browser tab.
- 2. Select the patient.
- 3. Select the series.
- Click the View Images button or double-click the selected series.
   The Viewing tab opens, and the series appears for viewing and manipulating.

#### Appending a series

Note Regardless of how many series are appended, the series are listed chronologically. This tool can be used to put all images from a patient together on a CD or to PACS.

- 1. Click the **Patient Browser** tab.
- 2. Select the study to append.
- 3. Select the first series.
- 4. Right-click the mouse to select the second series.
  The **Append Images** appears on the floating menu.

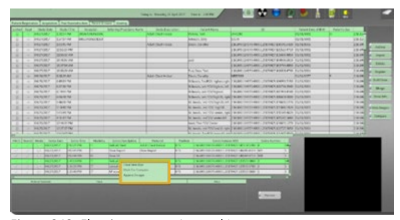


Figure 343: Floating menu - Append Images

5. Click Append Images.

The Please Wait popup appears.



Figure 344: Please Wait popup

A new series is created with (Appended) at the end of the description.



Figure 345: (Appended) series created

## **Chapter 11 Viewing Images**

**Viewing** lets you see already-scanned images from previous examinations. To view images, select the patient in **Patient Browser** and then select the series to view. To open the image, click the **View Images** button or double-click the series.



Figure 346: Active Viewing tab

The following tables identify the tools in the **Viewing** tabs that let you manipulate images. Some image tools appear on specific viewing tabs, only. The view tabs are **2D**, **MPR**, **3D**, and **CTP** (if enabled).

Table 55: 2D, MPR, 3D, and CTP image tools

Table 55: 2D, MPR, 3D, and CTP image tools			
Image tools	Tool name	Action	
Common tools			
B	Clear Tool	Resets the tool to the default pointer device.	
<b>Q</b>	Window Width/Center	Adjusts window width and center of image.	
Q	Zoom	Magnifies the image.	
\$	Pan	Adjusts image on X or Y axis.	
0	Invert	Inverts black to white and white to black.	
	Capture	Saves a screen capture of a selected viewport.	
	Capture All	Saves screen captures of all visible viewports.	
Reset	Reset	Reverts all images back to their original mode.	
2D and CTP tools			

Image tools	Tool name	Action	
A	Region of Interest (ROI)	Defines a circular <b>ROI</b> and displays the <b>ROI</b> information.	
R	Arrow	Draws an arrow on the image.	
2D, CTP, and	MPR tools		
*	Line	Draws a line on the image and is used for measurement.	
**	Angle	Draws an angle on the image and displays the angle information.	
2D only tools			
	Add Annotation	Create text box for annotation.	
Rotate	Rotate	Rotates images.	
	Reverse Image Stack	Reverses the order in which images display.	
1-1	Flip Vertically	Flips images up or down.	
<b>←</b>  →	Flip Horizontally	Flips images right or left.	
<b>«</b>	Cine Reverse	Cine backward through the image.	
<b>»</b>	Cine Forward	Cine forward through the image.	
	Stop	Stop the cine loop.	
MPR only too	ls		
Ø	Tilt	When selected a White 'steering' wheel allows you to correct a rotated image.	
3D only tool			

Image tools	Tool name	Action	
Color Preset  CT Angio 1	Color Preset	Dropdown menu allows you to select from multiple color options.	
Render Mode Color	Render Mode display images in Colo		
Orientation Superior	Orientation	Dropdown menu that allows you to select from multiple orientation options.	
	Rotate	Rotates the 3D image.	
1	Undo	Reverses the most recent action taken.	
C	Redo	Restores the most recent Undo action taken.	
CTP only tools			
•	Perfusion Artery/Vein Selection	Select to place the arterial and venous ROIs on the images.	
	Calculate CBF, CBV, MTT Map	Select to calculate the CT Perfusion maps.	
	Clear Perfusion Map	Cancels the calculations and returns to <b>Calculation</b> mode.	
	Show Artery/Vein Flow Graph	Displays the Arterial Venous Flow graph.	
	Peak Image	Displays the image that has the highest HU value based on the arterial ROI placement.	

## **Using keyboard shortcuts**

Keys are a quick way to navigate around. The table below provides keyboard shortcuts you can use to manipulate images in the **Viewing** tab.

Table 56: Arrow key navigation

Arrow keys	Action	
	To scroll through images.	
< >	To adjust the window center.	
PgUp	To quickly scroll through images.	

#### Setting window width and center

**Note** Any modifications you make are not saved to the image.

- 1. Select a patient from **Patient Browser**, select the series to view.
- 2. To open the image, click the **View Images** button or double-click the series.

The Viewing tab is enabled and the 2D viewer opens.

- 3. The following options allow you to adjust the window width and center of the image:
  - Click the Window Width/Center icon in the Tools menu, then while holding the left mouse button down drag up/down to modify Window Center and right/left to modify Window Width.
  - To adjust with a preset, click the Windowing dropdown and select a preset.



Figure 347: Windowing preset dropdown list

 Type values in the Width and Center text boxes and click the Apply button.



Figure 348: Windowing Width and Center text boxes, and the Apply button

Right click over an image and use the Activate Window Tool
option then while holding the left mouse button down drag
up/down to modify Window Center and right/left to modify
Window Width.



Figure 349: Right click menu

## Viewing images in 2D

**2D** lets you view scanned images in a **2-Dimensional** space. Standard **2D** mode is used when *only* one dataset is loaded. The default layout is a  $2 \times 2$  grid.

The **Viewing** tab and **2D** viewer opens when you select a dataset from the **Patient Browser**.

- 1. Select a patient from **Patient Browser** and select the series to view.
- 2. To open the image, click the **View Images** button or double-click the series.



The **Viewing** tab is enabled and the **2D** tab is opened.

Figure 350: 2D tools

- 3. Use any of the image tools to manipulate your images.
- 4. Click the **Reset** button to reset images back to the original setting(s). You cannot undo this action.

## **Comparing images**

You can compare images in two different ways:

**Note** You can compare two series from the same patient or two series from different patients.

## Using the floating menu to compare images

- 1. Select the patient in **Patient Browser**.
- 2. Select the first series from the series window.
- 3. Right-click and click Mark for Compare from the menu.

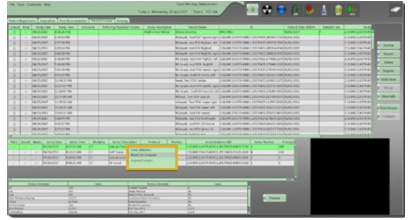


Figure 351: Floating menu - Mark For Compare

- 4. Select the second series or a series from a different patient.
- 5. Right-click and then click **Compare With Selected Series** from the floating menu.

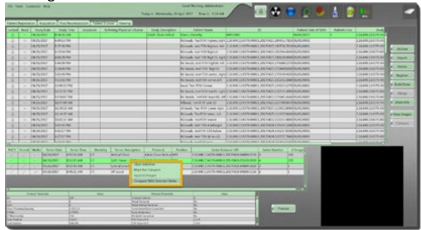


Figure 352: Floating menu - Compare with Selected Series

Both series are loaded into **Viewing** to compare.

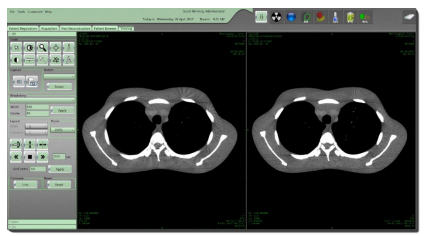


Figure 353: Compared series

6. Click the **Link** button to link both images together to view.



Figure 354: Link button

The **Unlink** button replaces the **Link** button.

7. Click the **Reset** button to reset images back to the original settings.

## **Using the Compare button to compare two images**

1. Select the patient in Patient Browser.

- 2. Select the first series.
- 3. Press and hold the **Ctrl** key.
- 4. Select the second series. Both series are highlighted.
- 5. Click the **Compare** button.



Figure 355: Using the Compare button

## Comparing a scout and a scan

- 1. Select a patient from **Patient Browser**.
- 2. Select a scout from the series window.
- 3. Press and hold the Ctrl key on the keyboard.



Figure 356: Scout and scan selected to compare

- 4. Select the scan from the series window. Both images are highlighted.
- Click the Compare button.
   The scout and the scan will appear on screen at the same time. A
  green localizer line appears on the scout.

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6. Compare the scout to the scan.

Figure 357: Comparing a scout (1) and a scan (2)

7. To remove the scout, right-click on the scout and click **Hide Scout Viewpoint**.

To return the scout to view, right-click in the viewing window and click **Show Scout Viewport**.

#### Using the ROI

- 1. Select a patient from **Patient Browser** and select the series to view.
- 2. To open the image, click the **View Images** button or double-click the series.

The **Viewing** tab is enabled and the **2D** tab is opened.

- 3. Click the ROI tool.
- Move the mouse pointer to the image where you want the ROI located.
- 5. Click the left-mouse button and drag the **ROI** diameter to the required size. To lock the **ROI** in place, click the left-mouse a second time.
  - To change the location of the ROI or the details of the ROI, click the ROI or measurements you wish to move. The ROI and its measurements will turn yellow, and the pointer becomes a hand. Click and hold the mouse button on either the ROI or it's measurements and drag to a new location. Click anywhere outside the ROI to freeze it in the new location.
    - When you move the ROI to a different location the measurements of the ROI are automatically updated based on the new location.

6. To remove the **ROI**, left click anywhere on the **ROI**, right click to see the floating menu, and click **Delete Annotation**, or click on the **ROI** and when it turns yellow, press **Delete** on the keyboard.

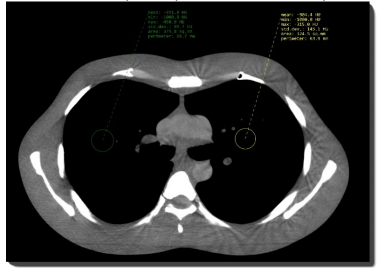


Figure 358: ROI

#### Using Layout and Rotate in 2D view

**Layout** lets you alter the number of images presented on the **Viewing** tab. **Rotate** lets you turn the images. Select a patient from **Patient Browser**, select the series to view.

- 1. Select a patient from **Patient Browser** and select the series to view.
- 2. To open the image, click the **View Images** button or double-click the series.

The **Viewing** tab is enabled and the **2D** tab is opened.

3. To adjust the layout of the viewing area, click the **Rows** and/or **Columns** dropdowns to select the number of rows or columns you want to show.



Figure 359: Layout (viewing tools)

4. To rotate the image, click the **Rotate** dropdown and select the number of degrees to rotate the images.

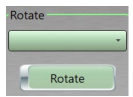


Figure 360: Rotate dropdown

5. Click the **Rotate** button to see the images turn to the new angle.

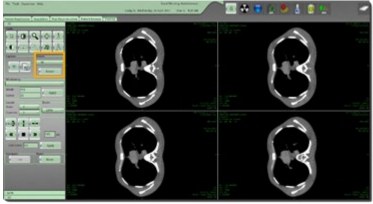


Figure 361: Rotate (viewing tools)

## Applying a grid to your images in 2D

- 1. Select a patient from **Patient Browser** and select the series to view.
- 2. To open the images, click the **View Images** button or double-click the series.

The **Viewing** tab is enabled and the **2D** tab is opened.

3. Change the size of the grid in the **Grid (mm)** text box.

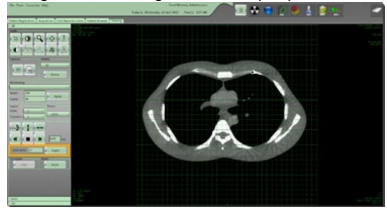


Figure 362: Grid (mm)

- 4. Click the **Apply** button to apply a grid over the image.
- 5. Perform one of the following to remove the grid:

- Click the Reset button.
- Enter 0 in the **Grid (mm)** text box and click the **Apply** button.

## Viewing images in MPR

Multi-Planar Reformation (MPR) allows images to be created from the original Axial plane into Coronal, Sagittal or Transverse (Axial) planes. MPR is fast, uses all the attenuation values in the dataset, and can be easily performed on the workstation. MPR however, provides on a two-dimensional (2D) display of the image data.

Viewer layout is 2 x 2 as seen below.

- 1. Select a patient from **Patient Browser** and select the series to view.
- 2. To open the images, click the **View Images** button or double-click on the series.

The **Viewing** tab is enabled and the **2D** tab is opened.

3. Click the **MPR** tab.

The MPR screen appears.

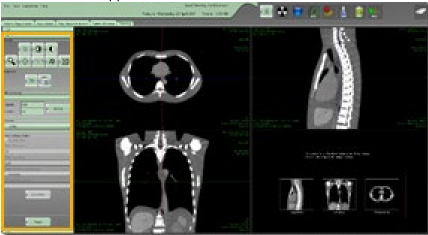


Figure 363: MPR tools

4. Select the image reformat at the bottom of the screen.

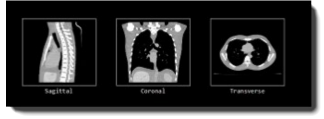


Figure 364: Image reformat selections

- 5. Use any of the image tools to manipulate the images.
- 6. The tilt tool can be used to modify the rotation of the images.
- 7. Adjust the image angle by moving the circle.

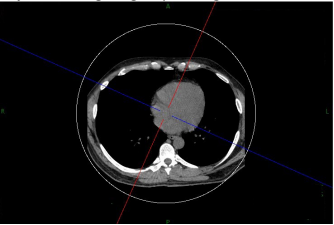


Figure 365: Tilt tool

8. Click the reset button to reset the images back to the original settings.

You cannot undo this action.

## Understanding and using slab

Through the reformation process, axial images are stacked creating a volume, or slab, which can be assessed in different planes. The thickness and spacing of each slab can be varied to meet the needs of the viewer. The reformations can be displayed in an average, maximum or minimum projection.

**MPR's** should be created using 1.25mm slices with a spacing of 0.625mm whenever possible.

## Creating the slab

- 1. Select a patient from **Patient Browser** and select the series to view.
- 2. To open the images, click the **View Images** button or double-click the series.

The Viewing tab is enabled and the 2D tab is opened.

- Click the MPR tab.The MPR screen appears.
- 4. Click the **Sagittal**, **Coronal**, or **Transverse** plane to create your slab.



Figure 366: Image formats

- 5. The **Secondary Series** option is enabled.
- 6. Select Enable Slab.



Figure 367: Enable Slab option

The **Enable Slab** option is inactive if no **MPR** view is selected.



Figure 368: Enable Slab option under Secondary Series

7. Set the **Cyan** lines to determine the beginning and end of the slab.

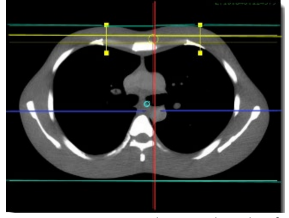


Figure 369: Cyan Line and cyan circle to drag for FOV

- 8. Use the **Cyan** circle to drag the planned slab if required.
- 9. Define the **Slab Thickness** and **Slab Spacing** in the text boxes.



Figure 370: Slab Thickness and Slab Spacing text boxes

You can also use the **yellow** squares found on the slab thickness display to modify the **Slab Thickness**.



Figure 371: Small yellow boxes to manually drag for desired slab thickness

10. Click the **Slab Rendering Options** dropdown to select the appropriate option.



Figure 372: Slab Rendering Options dropdown

The following options are available in MPR Slab mode:

Slab Thickness	The thickness of the MPR slab.	
Clab Chasing	The space between the start of one slab and	
Slab Spacing	the next.	
	Where you define the pixel values that will be	
Slab Rendering	displayed in each slab: options include,	
Options	Average, Maximum Intensity and Minimum	
	Intensity.	

Maximum	The highest pixel values for all slices within		
Intensity	the slab is displayed.		
Minimum Intensity	The lowest pixel values for all slices within the		
William intensity	slab is displayed.		
	The pixel values of all slices within the slab		
Average	are combined and the average value for each		
	pixel is displayed.		
Series Description	Text field for naming the series of images		
Series Description	created when clicking the <b>Generate</b> button.		
	Define the slab thickness. The boxes on the		
Yellow lines	lines allow you to adjust the thickness using		
	the mouse.		
	Define the slab <b>FOV</b> and dictate the range of		
	the new series to be generated. The cyan		
Cyan lines	lines are adjustable by clicking and dragging		
Cyan inies	on the lines themselves; both lines are moved		
	by clicking and dragging the central circle		
	marker.		
Red, blue, and	Define the cross sections of the anatomy		
green lines	being viewed.		
	Generates a new series with the name given		
Generate	in the <b>Series Description</b> field, based on the		
	selected MPR view pane.		

11. Select the **Tilt Tool** to correct any rotation on the image.



Figure 373: Tilt tool

12. Use the mouse pointer to move the white **Tilt** circle.

Note The circle does not represent the Field of View



Figure 374: Tilt white circle

- 13. Enter the slab name in the **Series Description** text box.
- 14. The slab can be previewed in the bottom right viewport.

Note Make sure Zoom is at 100% or below.

Ensure all expected anatomy is included when previewing the created MPR.

15. When you are ready to save, click the **Generate** button. A **Saving Series** popup appears.

When the series is complete, the **Capture Complete** pop-up appears.



Figure 375: Capture Complete popup message – Series saved

- 16. Click **OK** to close the **Capture Complete** popup message.
- 17. The new **MPR** images appear in the Patient Browser with the description in the Series Description text box.

CT	50cm Helical	Adult Abdomen Helical	
CT	Coronal Abdomen	Adult Abdomen Helical	
CT	Lateral Scout	Adult Abdomen Helical	

Figure 376: MPR images in Patient Browser

18. To create additional MPR's. 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 3D-volume. The initial display shows the **3D** volume and a box appears around it. **MPR** planes also appear.

- 1. Select a patient from **Patient Browser** and select the series to view.
- 2. To open the images, click the **View Images** button or double-click the series.

The **Viewing** tab is enabled and the 2D tab is opened.





Figure 377: 3D tools

- 4. To rotate the image up to 360°, click **Rotate** and move the image with the mouse pointer to the rotation of choice.
- 5. You can change the **Color Presets** from the dropdown menu:

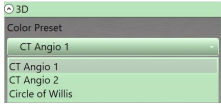


Figure 378: 3D Color Presets

6. You can change the **Render Mode** from the dropdown menu:



Figure 379: 3D Render modes

7. Click the **Orientation** drop-down box to assign an orientation:



Figure 380: 3D Orientation options

8. Click the **Reset** button to reset images back to the original settings. You cannot undo this action.

## **Chapter 12** Post Reconstruction

The system stores multiple patient series of raw data to allow post reconstruction of images. **Post Reconstruction** allows reconstructing of the acquired data using different algorithms, slice thicknesses, or use of image enhancement algorithms, such as **Metal Artifact Reduction**, **Noise Reduction** and **Windmill Correction**.

#### **Reconstruction Overview**

#### Metal artifact reduction

Streak artifacts are often seen around metal leads, prostheses, applicators, bone, or metal screws. Numerous factors can contribute to these streaks including under-sampling, photon starvation, patient motion, beam hardening, and scatter. You can use **Metal Artifact Reduction (MAR)** to reduce these streaks. **MAR** removes the metal from the image to reconstruct the soft tissue only; then it adds it back, to reduce the artifacts. This is currently used only on **Axial** scans.

#### Noise reduction

Noise appears as grain on the image and is caused by a low signal to noise ratio. This occurs more commonly when a thin-slice thickness is used. It can also occur when the radiation dose is insufficient to penetrate the anatomy being scanned.

Note Noise reduction applies to post-processing filters that reduce the amount of noise in the images. In clinical practice, using noise reduction may allow for a reduction in CT patient-dose depending on the clinical task, patient size, anatomical location, and clinical practice. Consult with the site's radiologist and physicist to determine the appropriate dose to obtain diagnostic image quality for a particular clinical task.

#### Windmill Correction

The Windmill Correction reduces artifacts that are common in **Helical** scans due to the nature of the cone-beam reconstruction.



Figure 381: Active Post Reconstruction tab

The tools available to **Post Reconstruction** are identified in the table below.

Table 57: Reconstruction tools

Image tools	Tool name	Action	
Load	Load Images	Loads images from selected series into viewing.	
Stop	Stop	Cancels the current, post- reconstruction request. All images are generated until you click the <b>Stop</b> button.	
FOV	FOV	Adjusts the <b>FOV</b> prior to reconstruction.	
₩ Z	Clear Tool	Resets tool to default pointer device.	
<b>O</b>	Window Width/Center	Adjusts the width and level of selected image.	
Q	Zoom	Magnifies the image.	
	Pan	Adjusts the image on X or Y axis.	
0	Reset	Resets the display to default viewer settings.	
Start Recons	Start	Begins your <b>Post Reconstruction</b> .	
Resend	Resend	Sends the last acquired scan from the recon workstation to the <b>Patient Browser</b> .	

## **Performing Post Reconstruction**

The following figure identifies parts of **Post Reconstruction**:

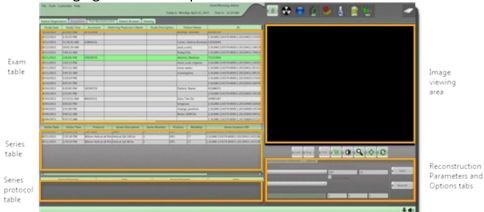


Figure 382: Post Reconstruction areas

- 1. Click the **Post Reconstruction** tab.
- Select a study in the Exam Table.
   When you select a study, all the scanned series for that study appear in the Series Table.
- 3. Select the series to reconstruct.

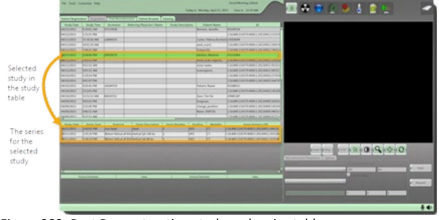


Figure 383: Post Reconstruction study and series tables

4. Click the **Load** button.

The scan or series will load into the viewer. The series protocol table and the **Reconstruction Parameters** and **Options** tabs are active. The scan will appear in the **Image Viewing Area**.

5. View the study in the Image Viewing Area.



Figure 384: Post Reconstruction viewing image area

- 6. To modify the **FOV**, perform the following:
  - Click the FOV tool, click and drag the mouse to form a square on the image in the Image Viewing Area.
    - The size of the square appears in the Width box in the Reconstruction Parameters and Options tab.
  - Click the circle in the middle of the FOV square and drag to move the FOV.
  - Click one of the two drag boxes on the corners to adjust the size of box.
    - The Width dimension, and X/Y coordinates are adjusted as the size changes.
    - The **FOV** size cannot exceed the range of 50 600mm square.



Figure 385: FOV resizing boxes

**Note** You can also enter a number in the **Width** or X and Y box to define a specific **FOV**.

Alternatively, click the **Use FOV Max** option to use the maximum **FOV**.

7. Click the **Sharpness** dropdown to select a reconstruction algorithm from the **Reconstruction Parameters** tab.

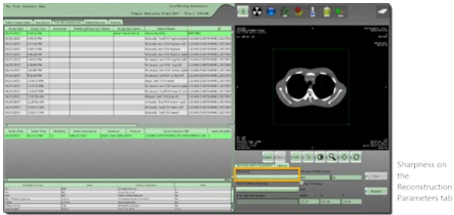


Figure 386: Sharpness on the Reconstruction Parameters tab



Figure 387: Reconstruction Parameters Sharpness dropdown

8. Click the Slice Thickness/Spacing dropdown to select.



Figure 388: Reconstruction Parameters Slice Thickness/Spacing dropdown

The slice thickness and spacing options available are determined by the type of scan that was acquired (Axial vs. Helical).

The # of Expected Images text box shows the calculated number of images that will be reconstructed based on the parameters used for the reconstruction.



Figure 389: # of Expected Images

- 10. Click the **Options** tab. The following are only available during **Post Reconstruction.**
- 11. Perform the appropriate action:
  - If desired, select Noise Reduction for an Axial or Helical scan.



Figure 390: Noise Reduction on the Options tab for a Helical scan

• If desired, select **Perform Windmill Correction** for a **Helical** scan.



Figure 391: Perform Windmill Correction and/or Noise Reduction on the Options tab for a Helical scan

If desired, select Metal Artifact Removal for an Axial scan.



Figure 392: Metal artifact removal

12. Click the **Start** button to generate a new dataset.

When you click the **Start** button, the reconstructed images appear in the viewing pane.



Figure 393: Please wait while the system performs data reconstruction message

13. When the reconstruction is complete, the images appear in **Patient Browser.** 

## Resending images from the scanner to the workstation

Pressing the Resend button lets you send the last acquired scan from the recon computer to the **Patient Browser**. This may be necessary when you have wireless interruptions and/or workstation shuts down unexpectedly.



Figure 394: Resend button

#### Chapter 13 **Accessories and Options**

In this chapter you will learn how to convert a bed, stretcher, or any type of adjustable surface into a scanning platform using the Universal Transfer Board.

To request the catalog(s) to reference product descriptions/details and part numbers for the available accessories/options that are used with the BodyTom Elite, see "Contact information" on page 25:

When using a fixed scanner, the table moves from one portion of anatomy to another while the gantry remains stationary. With the BodyTom Elite, 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 can be used for most beds or stretchers. It is placed under the patient and secured to the bed or stretcher with straps.



**WARNING** 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 the Universal Transfer Board

The universal transfer board is a carbon-fiber, radiolucent board that is designed to work with any ICU bed or stretcher. The carbon-fiber board comes with a 0.5-inch-thick headboard and 2-inch x 5-foot straps to strap the board to the ICU bed or stretcher.

You can use the universal transfer board on any bed, table, or stretcher. 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 BodyTom Elite is moved into position and the scan is performed.

The universal transfer board is always used with mattress stiffeners.

The mattress stiffeners provide a solid 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 BodyTom Elite for easy transport.

The universal transfer board is used for adults.

**Note** The universal transfer board is an optional accessory that does not come with the system.

#### Table 58: Universal Transfer Board weight-bearing restrictions

The weight limit of the Universal Transfer board is equal to the weight limit of the patient bed. The weight limit on the portion of the Universal Transfer board that supports the patients head is 7.5 kg / 17lbs. The universal transfer board is used to support and scan the patients head, only.

See also "Parts that potentially come into contact with the patient" on page 88.



**WARNING** The weight limit for the superior portion of the Universal Transfer Board is 7.5kg or 17lbs.

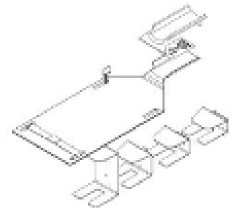


Figure 395: Universal transfer board and stiffeners

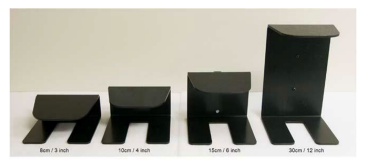


Figure 396: Four types of mattress stiffeners

**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 397: Mattress stiffener in place

- 3. The universal transfer board requires mattress stiffeners that provide a solid 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 398: 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 399: 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.

# Chapter 14 Cleaning and Storing the System and Workstation Specifications

Be familiar with this section before using the cleaning or storing the system.

#### Cleaning the scanner and workstation

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

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



WARNING Do not use flammable or potentially explosive disinfecting

sprays, since resultant vapor could ignite, causing personal

injury and/or damage to the equipment.

• w

**WARNING** To prevent short-circuiting or possible electrical shock, do not

spray cleaning agents or spill liquid cleaning agents directly

onto the machine.

 $\triangle$ 

**WARNING** Always electrically isolate this equipment from the main

electrical supply before cleaning and disinfecting it to prevent

short-circuiting or possible electrical shock.

<u>^</u>

**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.

<u>^</u>

**WARNING** Before cleaning the workstation (drive system), be sure to

disconnect the workstation from the wall outlet (power source). Failure to do so could result in electrical shock and cause severe injury to you and/or damage to electrical

components.

<u>^</u>

**CAUTION** Do not allow electrical components to become wet. For eye

and hand protection, it is important to wear safety glasses

and rubber gloves, respectively.



#### CAUTION

Do not spray cleaning agents directly on the system. Spray a clean cloth with the solution and then wipe down the scanner and workstation.

#### Cleaning the outside of the scanner and workstation

- Prepare detergent/disinfectant (regulated by EPA as hospital disinfectant) solution according to instructions on label for correct usage.
  - Use a basin or spray bottle (with product label).
  - Use a pump (usually on detergent/disinfectant containers) to dispense the concentrate in the basin or spray bottle, then fill with correct amount of tap water.
  - If using a spray bottle, empty and rinse out after use.

Note The stability of the solution is unknown after 24 hours; therefore, a fresh preparation of cleaning solution *must* be prepared for each day of cleaning.

- 2. Use general purpose germicidal cleaner on the external covers and rails. Do not use cleaners on the screens.
- 3. Use swabs moistened with cleaning solution, clean and remove any dust, soil, dried contrast media, or foreign matter; allow all components to air dry.
- 4. Wipe down and clean the frame of device and allow to air dry; return to its storage area.

**Note** Wash (at 25°C) with neutral detergents, *only*; softening agents *are not allowed*.

The following recommended products are registered by the EPA as hospital disinfectant; these solutions are quaternary ammonium compounds and are used in environmental sanitation of non-critical surfaces:

- TB Quat™ is a cleaning solution manufactured by ABC Compounding Co.
- Wex-cide™ is a disinfectant manufactured by Wexford Labs, Inc., product number Wexcide128.

#### Maintenance of the workstation



**WARNING** Maintenance checks and all service must be performed by service personnel trained by NeuroLogica Corp. See "Contact information" on page 25.

#### Storing the system

#### Storing the scanner and workstation

Store the scanner in a dry, well ventilated, climate-controlled area. You can use the key to lock the scanner when not in use. See "Identifying operator control panel buttons" on page 80 to locate the lock.



**CAUTION** When the scanner is not in use and stored, it must be plugged in a 120V or 250V outlet to charge the batteries.



**CAUTION** When the workstation is not in use and stored, it must be plugged in a 120V (or other compatible) outlet to charge the batteries.

Store the scanner on its centipedes or castor wheels (feet).

**Note** If the floor surface is soft (spongy) store the system on its centipedes to disperse the weight of the system evenly.

 After transporting the scanner to an acceptable storage location, you can either store the system in **Transport** mode (on its caster wheels) or Scan mode (on its centipedes).



Figure 400: BodyTom Elite castor wheels

2. Turn off the scanner and workstation. The system is now ready to be stored. See "Powering the workstation" on page 104.

Note It is recommended practice that the scanner is plugged in and turned on even when it is not in use.

## Storing the QA phantom

Store the phantom in a secure location with easy access for the daily QA procedure.

## **Workstation specifications**

Table 59: Workstation specifications

Phase	Single	
Voltage Range	100-240VAC ± 10%	
Factory Outlet Recommendations	NEMA 5-20R	NEMA 6-30R
Frequency	50 or 60Hz	
Battery Capacity	Fully Charged/ 12 hrs. (Typical)	
Typical Usage	110-120 VAC 60 Hz	230-240VAC 50 Hz
Wiring	125V, 2 Pole, 3 Wire Grounding	250V, 2 Pole, 3 Wire Grounding
Battery Operating Voltage	51.8VDC	
Overall width	41in. (104cm)	
Overall height	79in. (199cm)	
Overall length	101in. (256.5cm)	
Weight EST	3510 lbs. (1592kg)	
Battery power (2) 12 VDC (lithium polymer)	800W	
Max programmed speed fwd.	1.6 MPH	
Max recharge time ~ 8hrs.		
Max continuous operation		
Locking and unlocking cycles		
Hrs transport over floors	2hrs.	
Hrs system locked no external power	2hrs.	
Max slope holding angle with scanner	7º C (44.6º F)	
Max doorway threshold	1in. (2.54cm)	

Max elevator threshold	¾in. (1.905cm)
Height to locking adapter	8.59in 8.69in.
Min/Max storage temperature	-25º C to 70º C (-13º F to 158º F)
Min/Max operating relative humidity	
Min/Max storage relative humidity	20% to 85% (non-condensing)
Min/Max ambient operating temperature	15º C to 35º C (59º F/95º F)

## Understanding the symbols and product-marking plate

Table 60: Symbols and product-marking plate

Symbol	Description
$\sim$	Indicates alternating current.
	Indicates protective earth (ground).
<u> </u>	Indicates a caution: consult accompanying documents.
4	Indicates a caution: risk of electrical shock.
	Indicates electrostatic sensitive devices.
<u> </u>	Indicates a warning: high temperature.
-3 ·c -3 ·c (-3 ·r)	Indicates temperature limits.
ر آر	Indicates mechanical deactivation device.
(( <u>(</u> ))	Indicates a radiation precaution; may be affected by radiation from other sources; may produce interference that affects other equipment.
	Indicates manufactured by.
	Indicates toe/foot crush hazard; positioned in the front of the unit adjacent to the safety bumper and is a warning for population when lowering the unit.
$\bigcirc$	Indicates a coil power cord.
	Indicates a chain hazard could cause severe personal injury.

Symbol	Description
	Indicates to keep away from rain for packaging.
XXX eH	Indicates a humidity limit for packaging.
	Indicates a warning: battery charging.
	Indicates fuse usage.
	Indicates accompanying operating instructions in the user manual must be followed to safely
	operate equipment.

Note Disregarding information on safety is considered abnormal use.

## Locating the product-marking plate on the workstation



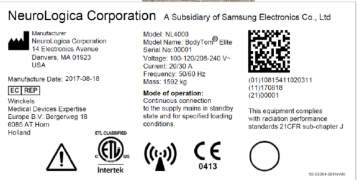


Figure 401: Product-marking plate on side of the workstation

## Listing of replacement parts for workstation

To ensure proper compliance requirements of replacement parts, (for example, cables and accessories), parts must be purchased through NeuroLogica Corp.



WARNING

Using other manufacturer cables and accessories may affect EMC performance. Unauthorized use of these items will void warranty and may cause harm to patient, others and/or equipment.

## **Product Safety and Electromagnetic Comparability**

Tested by: Intertek Testing Services NA, Inc., 70 Codman Hill Road,

Boxborough, MA 01719

# Appendix A Glossary

А

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 Control	Software used to adjust or modulate the
(AEC)	mA 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. Also
	referred to as step and shoot.

В

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 Axial, Helical or Dynamic scan when a threshold level of contrast enhancement is reached at a specified
	region of interest.

(

Collimation	Restricts x-ray to only the selected
	anatomy, minimizing dose to patient and
	reducing scatter.

Computed Tomography Angiography (CTA)	A test that uses x-rays to provide detailed pictures of 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 <sub>vol</sub> )	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. It is not patient dose. The CT dose index volume is noted as CTDI <sub>vol</sub> . It is reported in units of mGy. The CTDI <sub>vol</sub> is based on measurements made by the manufacturer in a factory setting. The CTDI <sub>vol</sub> is calculated differently for both the Axial and the Helical mode:
	For Axial scan mode: $CTDI_{vol} = [(N \times T)/I] \times CTDI_w$ .
	For Helical scan mode: $CTDI_{vol} = 1/pitch x$ $CTDI_{w}$ .
	See also dose and patient dose.
Computed Tomography Dose Index (CTDI <sub>w</sub> ) 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 <sub>w</sub> .
Computed Tomography (CT) number	Relative value assigned to each pixel to quantify the attenuation occurring in each voxel 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 Communication in Medicine (DICOM)	Digital Imaging and Communications in Medicine, or DICOM, is a standard that helps people doing work in 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	The generic term that refers to the CTDI <sub>vol</sub> , the standardized parameter to measure scanner radiation output – or the amount of 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 Protocol (DHCP)	A standardized network protocol used on Internet 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 (multiple detector widths)	Data acquisition at multiple time points over the same anatomic location(s).

E

Electromagnetic Compatibility (EMC)	The branch of electrical sciences that studies the unintentional generation, propagation, and reception of electromagnetic energy with reference to the unwanted effects (Electromagnetic interference (EMI)) that such energy may induce.
Electromagnetic Interference (EMI)	A disturbance generated by an external source that 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.

F

Field of View (FOV)	The diameter of the acquired attenuation
	data displayed across the image matrix.

Н

Helical scan mode	A CT acquisition whereby an x-ray
	acquisition whereby 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	A Radiology Information System (RIS) is the
System/Radiology	core system for the electronic
Information Systems	management of imaging departments. The
(HIS/RIS)	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.

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.
Iterative Bone Correction (IBC)	A feature build into the reconstruction software, which performs a correction on every single 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.

K

	· · · · · · · · · · · · · · · · · · ·
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.

M

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 (2D) grid numbers
	arranged in rows and columns.
Maximum Intensity	The multiplanar reformation technique
Projection (MIP)	that displays only the maximum pixel value
	along a ray traced through the object to
	the viewers assumed perspective in front
	of the scanner display screen.
Mean Transit Time (MTT)	A common measurement during CTP
	studies of the brain. Refers to the average
	transit time, in seconds, needed for blood
	to pass through a given region of brain
	tissue.

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
Procedure Step (MPPS)	information about the imaging performed
	back to the HIS/RIS or PACS.
Modality worklist 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.
Multi-Planar Reformation	The process of displaying CT images in a
(MPR)	different 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.

Ν

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.

P

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 Browser, local	Where the already-scanned patient list is
database	stored.
Patient coordinates	References are as follows:
	X left to right.
	<ul> <li>Y anterior to posterior.</li> </ul>
	Z head to feet.
Patient dose	The absorbed dose to a patient.
	See also CTDI <sub>vol</sub> .

Peak kiloVoltage (kV)	The penetrating power of the photons
	coming from the x-ray tube.
Picture Archive and	Stores medical information, including 2D
Communications Systems	images, and 3D medical images. All modern
(PACS)	PACS setups will work 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	The person within an organization
(RSO)	responsible for the safe use of radiation and
	radioactive materials as well as regulatory
	compliance.
Radio Frequency	Also called Electromagnetic Interference
Interference (RFI)	(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
	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

Casa dalan	
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. Like 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.
Size Specific Dose Estimate	Not dose to any specific organ but rather
(SSDE)	the mean dose in the center of the scanned
	volume. That is, SSDE is not the exact
	patient dose, as factors such as scan length
	and patient composition can differ from the
	assumptions used to calculate SSDE, for
	example conversion factors based on
	patient size provided to estimated patient
	dose for a patient of a particular size.
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.

Τ

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
(TAC)	versus time. 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

Volume Rendering (VR) image or object	A 3D modeling technique that utilizes the entire 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 CT 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 Listing of All Buttons, Tools, and Icons

## Status bar icons

Table 61: Status bar icons

Status bar icon	Status bar icon name	Status description	
	X-ray status	Identifies x-ray as on or off. The icon changes from a gray/black icon when x-ray is off to a rotating yellow/black icon when x-ray is on.	
	System state  System state  The orb changes color depending on the state the system is in.  See Table 25 on page 118 for a list of the different orb colors and system states they identification.		
	Scanner position	Identifies the system's current position relative to its zero reference.	
8	System E-STOP status	Identifies when <b>E-STOP</b> is engaged. The icon will flash when <b>E-STOP</b> is pressed.	
	System tube heat status	Indicates the current X-Ray tube heat status. The values are color coded as follows:  Blue 0% - 25%  Yellow 26% - 50%  Orange 51% - 75%  Red 76% - 100%	
	Scanner 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%	

Status bar icon	Status bar icon name	Status description
	System air freshness status	Indicates the air freshness status; it is recommended that an air calibration be performed:  Every eight (8) 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%  After calibration it returns to 100%.
	Workstation battery capacity status	Indicates the remaining workstation battery capacity 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 workstation into an outlet to charge if the battery capacity is low; a scan cannot complete when the battery capacity is 10% or lower.  When the workstation reaches the red capacity range, the system will shut down. A message informs you that the system will shut down due to low battery.  The lightning bolt icon signifies that the workstation is currently charging and goes away when

Status bar icon	Status bar icon name	Status de	scription
	Image storage space status	for image s	torage. The available es are color coded as  100% - 51% 50% - 20% 19% - 0%

## System state orbs

Table 62: System state orbs

Orb	Color	State
	Dark gray	The system is in an unknown state.
	Light gray	The system is powering up or down.
	Dark purple	The system is busy.
	Purple	The system is completing air calibration.
	Light purple	The system is archiving.
	Blue	The system is idle.
	Green	The system is ready to perform a scan.
	Light yellow	The system is planning.
	Dark yellow	The system is preparing.
	Light orange	The system is reconstructing.
	Dark orange	The system is scanning.

Orb	Color	State
	Pink	The system is not ready.
	Red	The system is in fault.

## **Workstation buttons**

Table 63: BodyTom Elite workstation buttons

	e workstation buttons
Workstation	Action
button	
Add	In <b>Protocol Manager</b> – adds a new protocol from the list.
Archive	In <b>Patient Browser</b> – selects the archive destination for selected information.
Begin	In <b>Patient Browser</b> – used to begin a protocol.
Build From	In <b>Protocol Manager</b> – used to create a new protocol from a previously saved protocol.
Build Dose	In <b>Patient Browser</b> – generates the dose for the selected patient.
Cancel	In <b>Patient Registration</b> – cancels the current query. In <b>Patient Browser</b> – cancels any series being imported.
Cancel Exam	In <b>Acquisition</b> – cancels the entire exam being performed.
Cancel Scan	In <b>Acquisition</b> – cancels the current scan within a protocol.
Clear	In <b>System Configuration</b> – clears information in fields.

Workstation button	Action
Close	In Protocol Manager – closes popup. In Store/Print Queue – closes the Store/Print Queue popup. In System Configuration – closes System or User Configuration dialog boxes.
Compare	In <b>Patient Browser</b> – allows you to select multiple series of patient images to compare in <b>Viewing.</b>
Continue	In <b>Acquisition</b> – authorizes the scanner to move to the next step.
Delete	In Patient Registration – deletes patient(s) from the Stored Results list.  In Patient Browser – deletes selected exam information from Patient Browser.  In Store/Print Queue – deletes a series to be stored or a series that failed to store.  In Protocol Manager – deletes a saved protocol.  In System Configuration – clears a saved dose setting to remove the restriction.
Delete All	In <b>System Configuration</b> – deletes saved dose settings to remove all restrictions.
Details	In <b>Store/Print Queue</b> – when you select one or more series, displays an explanation of why a series failed to store.
Dwn	Move selected item down the list.
Edit	In <b>Patient Registration</b> – used to modify protocols.
Echo	In <b>System Configuration</b> under <b>DICOM Servers</b> – echoes the selected server to test the connection.
Export	In <b>Protocol Manager</b> – exports protocols to a media device.
Extend	In <b>Acquisition</b> – provides extension of current protocol.

Workstation button	Action
Finalize	In <b>Acquisition</b> – completes the examination. Completes all protocols, builds Dose SR and images, and directs user to <b>Patient Browser</b> .
Generate	In Viewing - generates a new series with the Series  Description field information – based on the selected MPR.
Import	In <b>Patient Browser</b> – imports the exam information from <b>PACS</b> or <b>Media</b> .  In <b>Protocol Manager</b> – imports previously exported protocols to the workstation.
Manual	In Patient Registration – manually enters a new patient and, when completed, takes you to the Acquisition tab to acquire the data for a scan.
Media	In <b>Patient Browser</b> – used to select the destination for patient data to media.
Merge	In <b>Patient Browser</b> – combines two different image sets.
Minimize	In <b>Patient Browser</b> – minimizes the <b>Import for Media</b> popup.
New	In <b>Protocol Manager</b> – used to create a new protocol.
Ok	To accept selections you make.
PACS	In <b>Patient Browser</b> – used to select the destination for patient data to <b>PACS</b> .
Patient	In <b>Patient Browser</b> – used to select only patient information for a patient when using <b>Register</b> feature.
Patient/Study	In <b>Patient Browser</b> – used to select all exam information for a patient when using <b>Register</b> feature.
Pause	In <b>Store/Print Queue</b> – when you select one or more series, temporarily stops the series from being stored. This is a toggle button with the <b>Resume</b> button.

Workstation button	Action
Pause Exam	In <b>Acquisition</b> – pauses entire protocol.
Pause Scan	In <b>Acquisition</b> – pauses current scan within a protocol.
Play	In System Configuration under Audio Configuration – used to play audio files.
Prepare	For <b>Quality Assurance</b> – used to prepare workstation to run a <b>Quality Assurance</b> test.
Protocol	In <b>Acquisition</b> – selects an existing protocol for the current study.
Query	In Patient Registration – searches the HIS/RIS server for scheduled patients.
Register	In Patient Registration – registers the selected patient and then takes you to the Acquisition tab to acquire the data for an examination (scan).  In Patient Browser – opens the Create New Study dialog box and moves selected patient to Acquisition.
Registration	In Patient Registration – used to take patient information stored in HIS/RIS server to Patient Registration to choose the patient and study.
Repeat	In <b>Acquisition</b> – repeats the last scan that was performed.
Resend	In <b>Post Reconstruction</b> , sends the last acquired scan from the recon workstation to the Patient Browser.
Reset	In Viewing – reverts all images back to original state. In System Configuration – resets information back to default or clears information in fields.
Resume	In <b>Store/Print Queue</b> – when you select one or more series, continues to store previously paused series. This is a toggle button with the <b>Pause</b> button.

Workstation button	Action
Retry	In <b>Store/Print Queue</b> – when you select one of more series, tries to archive the selections.
Save	In <b>System Configuration</b> – saves updated information.
Search	In <b>Patient Registration</b> – searches queried patient entries for specific information.
Show Info	In <b>Patient Browser</b> – shows patient, study, series, and image information; used to modify series scanned under a wrong patient.
Start	For <b>Daily Calibration</b> – begins the daily (air) calibration.
Start Recons	In <b>Acquisition</b> – begins any post-reconstructions that were defined during the protocol setup. In <b>Post Recons</b> – begins a manual reconstruction
Stop	In <b>System Configuration</b> under <b>Audio Configuration</b> – stops audio files from playing.
Store	In Patient Registration – selects patient(s) from query results and moves them into the Stored Results list.
Unlock	In <b>System Configuration</b> under <b>User Accounts</b> – used by administrators to unlock a user's account.
Up	Move selected item up the list.
Update	In <b>Protocol Manager</b> – updates information on an existing protocol.  In <b>System Configuration</b> – updates information.
Validate	In <b>Acquisition</b> – prompts the application to send the selected protocol to the scanner and verify that the scanner has tube and battery capacity to perform the protocol.
View	In <b>Patient Registration</b> – shows selected patient details. In <b>System Configuration</b> – shows information.

Workstation button	Action
View Images	In <b>Viewing</b> – to load and views images.

# Viewing tools

Table 64: Viewing tools

Tool	Tool name	Action		
Y <sub>0</sub>		2D, CTP, and Viewing tool – draws		
1 4º 1	Angle	an angle on the image and		
		displays the angle information.		
53		2D, CTP, and Viewing tool – draws		
	Arrow	an arrow on the image, which can		
)		be repositioned.		
		CTP only tool – calculates the		
	Calculate CBF,	Cerebral Blood Flow (CBF),		
	CBV, MTT Map	Cerebral Blood Volume (CBV) and		
		Mean Transit Time (MTT) maps.		
	Clear	CTP only tool – cancels the		
	Perfusion Map	calculations and returns to		
	· cirasion map	Calculation mode.		
· [6]	_	Common tool - saves a screen		
	Capture	capture of selected viewport.		
	Capture all	Common tool - saves screen		
	Viewports	captures of all visible viewports.		
<b>«</b>	Cine Backward	<b>2D only tool</b> – cines backward		
	Cilie Backward	through the image.		
<b>&gt;&gt;&gt;</b>	Cine Forward	<b>2D only tool</b> – cines forward		
	- CC. C. T. T. C.	through the image.		
		Common tool (Acquisition, Post		
13	Clear Active	Reconstruction, Viewing) - resets		
10	Clear Active	the tool to the default pointer		
		device.		

Tool	Tool name Action					
	Change the Window Width Level	Common tool (Acquisition, Post Reconstruction, Viewing) — click and move pointer over image. Left click and hold down the mouse button and drag in chosen direction to adjust image width and level. Width and level values appear in the Width/Level status display. A pre-defined width level setting can also be selected. Select the preset from the dropdown list below the WL Preset button. Width and level presets can also be saved or deleted.				
FOV	Field Of View	Post Reconstruction tool – adjusts the Field Of View (FOV) prior to reconstruction.				
<b>+</b>	Flip Horizontal	<b>2D only and Viewing tool</b> – flips images right or left.				
-1-	Flip Vertical	2D only and Viewing tool – flips images up or down.				
	Invert	Common tool (Viewing) - inverts black to white and white to black.				
Load	Load	Viewing tool – loads images from selected series into viewing.				
	Measure (Line)	<b>2D, CTP, and Viewing tool</b> – draws a line on the image and displays length information.				
	Pan	Common tool (Acquisition, Post Reconstruction, Viewing) - click and move pointer over image. Left click and hold down the mouse button and drag the image in the chosen direction. Release mouse button to position image in new location.				

Tool	Tool name	Action		
	1001 Hame	CTP only tool – displays the image		
	Peak Image	that has the most visible contrast		
	i can image	(based on arterial <b>ROI</b> placement).		
	Perfusion	CTP only tool – selects the artery		
· Q	Artery/Vein	and vein to be used for performing		
	Selection	perfusion calculations.		
	Selection	Axial and Helical Viewing tool –		
		restores the last text editing or		
		resizing and positioning of controls		
( - 1	Redo	- if no other action occurred since		
		last time the <b>Undo</b> button was		
		clicked.		
		Viewing tool – performs the most		
Redo 1	Redo	recent action, again. The button is		
		disabled if the application cannot		
		redo the application.		
	Danian of	2D, CTP, and Viewing tool – defines		
" A 1	Region of	a circular <b>ROI</b> and displays the <b>ROI</b>		
	Interest (ROI)	information (5mm diameter by		
		default).		
. 0	Reset	Post Reconstruction and		
		Acquisition tool – resets the		
		display to default viewer settings.		
5.1	Reverse Image	2D only and Viewing tool –		
	Stack	reverses the order in which		
		images display.		
.50	Datata	MPR only and Viewing tool –		
	Rotate	rotates the image.		
		Acquisition tool if secut lines		
* +	Scan Region	Acquisition tool – if scout lines		
	Re-Draw	and the scan region is deactivated,		
_		allows you to reactivate.		
		CTP only tool – displays the Arterial Venous Flow graph.		
	Show	Antenal venous riow graph.		
		9		
	Artery/Vein	1"		
	Flow Graph			
		Owe		
		Doct Deconstruction tool consols		
		Post Reconstruction tool – cancels		
Stop	Stop	the current, post-reconstruction		
		request. All images are generated		
		until you click the <b>Stop</b> button.		

Tool	Tool name	Action		
	Stop Cine	<b>2D only tool</b> – stops the cine forward and backward.		
A	Text (Annotation)	2D only and Viewing tool – creates text box for annotation.		
	Tilt	MPR only tool – corrects a rotated image.		
	Toggle Layout	Acquisition tool – changes the layout to 2x2. Repeat process to return to 1x1.		
	Toggle Scouts	Acquisition tool – removes scouts from Acquisition.		
1	Undo	Axial and Helical Viewing tool – reverses the most recent action taken (a successful copy, cut, delete, undo or paste action).		
Undo	Undo	Viewing tool – removes the most recent action performed on image. The workstation remembers the last five adjustments made. The tool is disabled if the workstation canno redo the adjustment.		
<b>O</b>	Windowing	Common tool - adjusts the width and level of the selected image.		
Q	Zoom	Common tool (Acquisition, Post Reconstruction, Viewing) - click and move the pointer over the image. Left-click the mouse and hold down the left-mouse button and move in upward direction to zoom in (enlarge) and downward to zoom out (shrink).		

Table 65: Pendant buttons

Pendant butto	Button	Description	Action
	POWER	POWER	Illuminates when power is supplied to pendant.
	*	LASER	Turns on all three positional lasers. While the lasers are on, the scanner spins for the internal laser to be seen within the scanner opening.
POWER  GO TO  LASER SCAN PLANE  ZERO  REFERENCE		GO TO SCAN PLANE	Moves the scanner forward approximately 30cm. This is the distance between the internal and external lasers.
GANTRY—		ZERO REFERENCE	Sets the scanner to zero before starting a scout or a scan.
	<b>(</b>	MOVE BACKWARD (slow)	Pressing and holding moves the scanner backward 10mm per second.
SET MEMORY	<b>&gt;</b>	MOVE FORWARD (slow)	Pressing and holding moves the scanner forward 10mm per second.
SCAN REST POSITION POSITION  Neura Logica	<b>®</b>	MOVE BACKWARD (fast)	Pressing and holding moves the scanner backward 60mm per second.
	<b>③</b>	MOVE FORWARD (fast)	Pressing and holding moves the scanner forward 60mm per second.
	<b>(2)</b>	SET MEMORY	Allows the user to program <b>Scan</b> and <b>Rest</b> positions for the scanner.

		SCAN POSITION	Moves the scanner to the <b>Scan Position</b> saved using the Set Memory feature.
		REST POSITION	Moves the scanner to the <b>Rest Position</b> saved using the Set Memory feature.

# Appendix C Sample of Reference Protocols Provided

Table 66: Sample of BodyTom Elite adult protocols and important estimates

Protocol Name	Туре	kV	m A	Slice Thickness /Spacing	Sharpness	Resolution	Coverage	CTDI <sub>vol</sub> (mGy)	DLP <sup>5</sup> (mGy.cm)
Adult Head Axial	Axial	120	200	5.0 x 5.0	Soft Tissue	1 Sec.	250	47.32	1183
C-Spine Helical	Helical	120	250	1.25 x 1.25	Soft Tissue - Abdomen	Pitch = 0.8	400	24.16	966.4
Adult Chest Helical	Helical	120	150	1.25 x 1.25	Bone	Pitch =0.8	450	14.49	652.05
Adult Abdomen Helical	Helical	120	250	2.5 x 2.5	Soft Tissue - Abdomen	Pitch = 0.8	500	24.16	1208

Table 67: Sample of BodyTom Elite pediatric protocols and important estimates

Protocol Name	Type	kV	m A	Slice Thickness /Spacing	Sharpness	Resolution	Coverage	CTDI <sub>vol</sub> (mGy)	DLP <sup>6</sup> (mGy.cm)
Pediatric Head Axial	Axial	100	175	5.0 x 5.0	Soft Tissue	1 Sec.	200	32.02	640.4

<sup>&</sup>lt;sup>5</sup> DLP is based on length from coverage column

<sup>&</sup>lt;sup>6</sup> DLP is based on length from coverage column

## Appendix D Automatic Exposure Control

#### 1 Introduction:

Automatic Exposure Control (AEC) is a feature which allows the exposure to automatically be modified based on the attenuation of the scanned object. The main objective of AEC is to optimize the x-ray current based on prior knowledge of the scanned objects profile. AEC is used to optimize patient exposure while attempting to maintain acceptable diagnostic quality of the reconstructed images.

AEC uses image noise to optimize the scan current. The image noise on CT scanners can be traced to two sources: **Electronic Noise** and **Quantum Noise**. Electronic Noise is generated by the electronic components of the Data Acquisition System (DAS). **Quantum Noise** is related to x-ray generation. Currently **Quantum Noise** is the major component of noise on CT images, the contribution of **Electronic Noise** has become less significant since the early days of CT scanners.

## 2 Image Noise:

#### 2.1 Electronic Noise:

The DAS is composed of the crystals, the photodiodes, the Analog to Digital Convertors (ADC) and other electronic components known as "converter cards" since they convert x-rays into a quantifiable current. Thermal Noise is the most common source of electronic noise in the CT system. As the scanners internal temperature increases thermal noise becomes the dominant component of the DAS's electronic noise. Imperfections in the semiconductor chips used in the DAS also contribute to the Electronic Noise. However, with the advance in semiconductor crystals this has become less relevant. Currently Electronic Noise has no significant impact on image quality when using proper scan parameters, i.e., scan voltage, current and exposure.

#### 2.2 Quantum Noise:

The generation of x-ray photons can be described by a Poisson random process. Poisson random processes are used to describe event generation over a fixed time interval. A Poisson random process is used to describe Queues in general. In a queue the number of new arrivals to the queue over a fixed time interval follow a Poisson distribution.

The **Quantum Noise** is related to the standard deviation of the Poisson distribution which inversely proportional to the square roots of the number of events:

$$\sigma_I \propto \frac{1}{\sqrt{N}}$$

Where  $\sigma$  is the image noise and N is the number of detected photons. The above equation can help relate the image noise to the scan current, I, the scan time, t, and the slice thickness, Sw, since the number of photons is proportional to either one:

$$N \propto It \ and \ N \propto Sw$$

The image noise of a given scanner can be written as a function of the scan parameters:

$$\sigma = \frac{K}{\sqrt{I \cdot t \cdot Sw}}$$

To reduce the image noise, we can either increase the scan current, the scan time or the slice width, or any combination thereof. K is constant that is dependent on the image reconstruction process. It follows, for two different scan currents  $I_1$  and  $I_2$  over the scan times  $t_1$  and  $t_2$  and using the same slice thickness, the image noises for the two scans can be related using the following equation:

$$\frac{\sigma_2}{\sigma_1} = \sqrt{\frac{I_1 \cdot t_1}{I_2 \cdot t_2}} \quad or \quad \sigma_2 = \sqrt{\frac{I_1 \cdot t_1}{I_2 \cdot t_2}} \quad \sigma_1$$

The same relation exists between the image noise and the slice thickness

$$\frac{\sigma_2}{\sigma_1} = \sqrt{\frac{Sw_1}{Sw_2}} \quad or \quad \sigma_2 = \sqrt{\frac{Sw_1}{Sw_2}} \quad \sigma_1$$

As a result, doubling the slice thickness can reduce the image noise by almost 40%. Figure 1 shows the Noise as a function of the scan current (left) and the inverse of the scan current (right). The scan done for the same slice thickness of 2.5 mm and same scan time of 1 second.

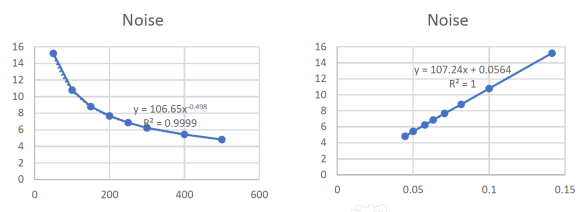


Figure 1: The Noise in a 20 cm water phantom as a function of the scan current I (left) and the inverse of square root of the current  $1.0/\sqrt{I}$ .

## 3 AEC working instructions:

#### 3.1 AEC input parameters:

AEC requires three different input parameters: Minimum mA, Maximum mA, and the Noise Level.

**Minimum mA:** The Minimum mA is set to prevent an unacceptable amount of noise in the reconstructed images.

**Maximum mA:** The Maximum mA value is used to prevent over-exposure of the patient being scanned. This is typically set to reduce the chances of over exposure which may lead to radiation sickness. However, in the case of the BodyTom Elite, the maximum scan current is set to protect the x-ray tube.

**The noise level:** The first step of AEC is to build an equivalent stack of cylindrical water phantoms or Water Equivalent Diameter (WED). The WED is created based on the measured attenuation from the scout. The noise level represents the noise in each section of the stack of water equivalent phantoms. The selected noise level should be within an acceptable range, and it should be dictated by the scan protocol.

One of the key features of the scanners AEC is the ability the user has to re-adjust the noise level based on the estimated mA until the desired mA profile is attained. The feature works as follows:

- 1. A protocol is created with the AEC feature enabled, and a Minimum mA, Maximum mA, and Noise Level are defined.
- 2. The above-mentioned protocol is used to create a Scout

image.

- 3. The user can toggle the AEC graph to view the scan current profile. The profile will be overlayed on top of the scout.
- 4. If the scan current profile is acceptable, then the user will initiate the diagnostic scan as desired.
- 5. If the scan profile is not acceptable, the user can **adjust** the noise level and **recalculate** the scan current.
- 6. The process can be repeated as many times as needed until the user is satisfied with the current profile. The AEC tool will allow the user to view the scan current before initiating the actual scan.

#### 3.2 The scan parameters:

The scan protocol parameters are not needed for AEC however they do affect the current estimation:

- 1. **kV**: the scan kV is used to select the appropriate noise table used for estimating the scan mA.
- 2. **Slice Thickness:** The noise is measured at a slice thickness of 5.0 mm; however, the scan protocol slice thickness can be any of the allowable thickness values. The selected slice thickness is then used to adjust the noise table using the equation in section 2.2. The entire noise table will be multiplied by the square root of the slice thickness ratio. The multiplication factor is:

$$\alpha = \sqrt{\frac{5.0}{Selected Sw in mm}}$$

 The reconstruction kernel: The noise image depends on the reconstruction kernel. AEC is limited to SoftTissue and PostFossa Kernels. AEC will be disabled if the user selects a different reconstruction kernel.

#### 3.3 Notes

When AEC is selected the user should be aware of:

1. **Patient Positioning:** The patient should be properly positioned as close as possible to the scanner iso-center. Failure to do so can lead to an over-estimate of the scan current leading to an increase in patient dose.

- 2. **Presence of metal implants:** AEC should not be used if the patient has metal implants in the region to be scanned.
- 3. **The measured noise:** the final noise in the image depends on the size of the scanned patient. AEC assumes that the patient is cylindrical, as such the measured noise level could be different then the selected noise level.
- 4. Anatomical features: AEC should be used when the region to be scanned includes significant differences in attenuation, such as the chest and abdomen. Anatomical regions with slight differences in attenuation like the head, will not benefit from AEC use.

#### 3.4 Sample protocols:

Below are some suggested protocols. The noise levels depend on the size and weight of the patient. The noise levels in the table below are for illustration purposes. The site physicist and CT manager should dictate the final noise levels.

	Noise Level	Minimum mA	Maximum MA	Slice thickness
Chest scanning	15	50	250	1.25
Chest/Abdomen	13	50	280	2.5

## 4 AEC algorithm description:

AEC uses the measured attenuation of the scanned object and the selected noise level to estimate the scan current at each planned scan location during the scan. The mA is typically estimated using different water phantom diameters.

The BodyTom Elite uses Z-modulated AEC where each planned scan location is modeled using a cylindrical water phantom, or WED. The WED is calculated using the scout profile. Once the WED is estimated a specific mA value is assigned to each planned scan location. The Flowchart (Figure 2) below describes the basic steps for using AEC:

- Select the appropriate AEC parameters to be used, those values are:
  - a. Minimum mA
  - b. Maximum mA
  - c. Noise Level

- 2. Acquire a Scout using the same kV that will be used for the Axial or Helical acquisition.
  - a. An AP or Lateral Scout can be used; however, AP scouts are preferred.
- 3. For each planned slice, the system calculates the WED and assigns a specific mA value to that location. Figure 3 shows the WED of the equivalent water phantoms as calculated based on the scout image (left). The estimated diameters (right) shows that the water portion of the phantom match the true diameter of the phantom.
- 4. Using the measured noise in different diameters water phantoms at different mA levels, find the mA that generates the selected noise level.
- 5. Adjust the mA based on the scan mode. Figure 4 shows the estimated mA as well as the adjusted mA for the axial scan mode.

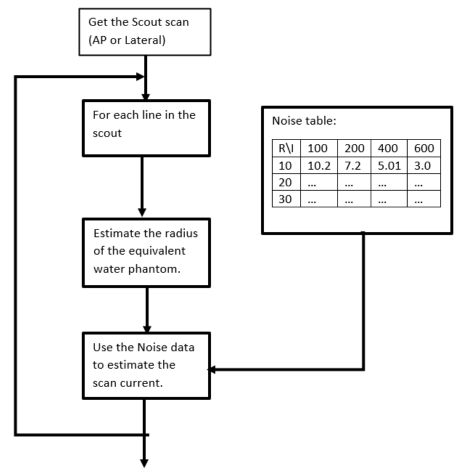


Figure 2: The AEC flowchart

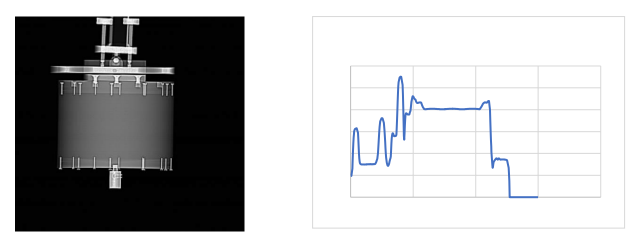


Figure 3: the AP scout of a 20 cm water phantom and the estimated radius based on the scout.

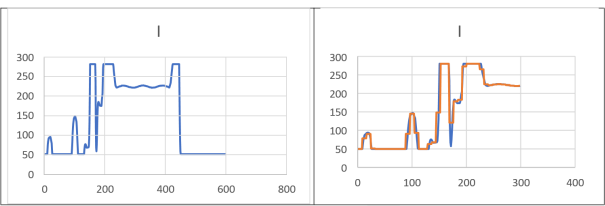


Figure 4: The estimated current using the equivalent diameter (left) and the scan current for axial scan mode (right).

#### 5 The Noise measurements:

The noise tables used for predicting the scan current is measured for each scan voltage. For each scan voltage the noise is measured using a set of predefined scan currents of 50, 100, 150, 200, 250 and 300 mA. The noise is measured using different water phantoms. In our case we have used 150, 200 and 300-mm water phantoms. Table 68 shows a sample of the noise table at 120 kV.

Phantom Diameter	Scan Currents (mA)					
(mm)	50	100	150	200	250	300
100	7.5868	5.4078	4.4009	3.8519	3.4769	3.1679
150	10.76	7.6690	6.2493	5.4768	4.8741	4.4806
200	17.41	12.3721	10.1349	8.8315	7.8699	7.1964
250	28.400	20.2000	16.4000	14.3000	12.7000	11.6000
300	49.952	35.6299	28.7523	24.9660	22.3726	20.2553

For example, if we desired to determine scan current in a 175mm water phantom that has 10 HU noise at 120 kV using the above table. The Noise is modeled using a 2-dimensional function of the current and the scan current. The scan current is then extracted from the 2D model. The noise is measured using a nominal slice width, typically 5.0 mm. The noise table will then be normalized based on the selected slice thickness of the scan protocol.

#### **Appendix E Rotating Anode X-Ray Tube**



GS-3073

Rotating Anode X-Ray Tube Tubes Radiogénes à Anode Tournante Drehanoden - Röntgenröhre Tubos de Rayos-X con Ánodo Giratorio



Note: Document originally drafted in the English language. Note: Document à l'origine rédigé dans l'anglais. Anmerkung: Dokument ursprünglich gezeichnet in der englischen Sprache. Nota: Documento elaborado originalmente en la lengua Inglesa.

Product Description
The GS-3073 is a 5.5" (140 mm)
150 kV, 2.5 MJ (3.5 MHU) maximum anode heat content, rotating anode insert. This insert is specifically designed for CT Scanners. The insert features a 7° tungstenrhenium facing on molybdenum with a graphite backed target and is available with the following nominal focal spot:

> 0.7 x 0.8 1.2 x 1.4 IEC 60336

Loading Factor for silt focal: Small - 120 kV. 100 mA Large - 120 kV, 200 mA

Maximum Anode Cooling Rate: 8,750 W (12,250 HU/sec)

mum continuous anode heat dissipation: 3,400 W (4,760 HU/sec)

Nominal Anode Input Power: Small - 23 kW IEC 60613 Large - 42 kW IEC 60613

Reference Axis:

This insert is intended for use in a Varex Imaging B-240H housing.

Le tube GS-3073, est une tube à anode tournante de plateau 140 mm. (5.5 pouces), 150 kV, d'une capacité therimque de 2,5 MJ (3,5 MUC). Il est à spécialement concu pour une utilisation avec les scanners CT. Le pente de l'anode en molybdéne traitée, tungsténe, rhènium, recourte de graphite, est de 7°. La dimension des foyers est

0,7 x 0,8 1,2 x 1,4 CEI 60336

Facteur de charge pour foyer à fente: Petit - 120 kV, 100 mA Grand - 120 kV, 200 mA

Toux maximum de efroidissement de l'anode: 8,750 W (12,250 UC/sec)

Description calorifique maximim de l'anode (en continu): 3,400 W (4,760 UC/sec)

Pulssance Nominale de l'anode: Petit - 23 kW CEI 60613 Grand - 42 kW CEI 60613

Perpendiculaire à la face de sortie

Ce tube est essentiellement destiné à être employé dans les gaines Varex Imaging des séries B-240H.

Produktbeschreibung
Die GS-3073 ist eine 140 mm
(5.5") Doppelfokus Drehanoden-Röntgenröhre, mit einer Anoden Wärmespeicherkapazität von 2.5 MJ (3.5 MHU) und einer max. Spannungsfestigkeit von 150 kV. Die Röntgenröhre wurde für den Einsatz an CT Scanners entwickelt. Der rückseitig graphitbeschichtete Wolfram Rhenium-Molybdan Anodenteller besitzt einen Winkel von 7°. Folgende Brennfleckkombination ist lieferbar

> 0.7 x 0.8 1.2 x 1.4 IEC 60336

Ladefaktor: Klein - 120 kV, 100 mA Gross - 120 kV, 200 mA

Maximale kontinuleriiche Wärmeableitung des Anodentellers: 3,400 W (4,760 HU/sek)

Nominale Anoden

Gross - 42 kW IEC 60613

Referenz Achsen:

Die Röntgenröhre ist für den Einbau n die Varex Imaging Strahlerhaube B-240H vorgesehen.

Descrincion del Producto
El GS-3073 es un tubo de ánodo
giratorio de 140 mm (5.5"), 150 kV,
2.5 MJ(3.5 kUC), la cual es el maximo almacenaje termal del anodo, es diseñado especificamente para uso en CT scannners. El blanco emisor es una combinación de tungsteno, renio y molibdeno con grafito en la parte posterior con un rayo central de 7 grados. Disponible con las siguientes combinaciones de marcas focales:

> 0.7 x 0.8 1.2 x 1.4 IEC 60336

Carga Electrica Para la Abertura Focal: Pequeño - 120 kV, 100 mA Grande - 120 kV, 200 mA

Medida Maxima dei Enfriamiento dei Anodo: 8,750 W (12,250 HU/seg)

Maxima disipación termal

continuo del Anodo: 3,400 W (4,760 HU/seg)

El Poder de Penetración para el Anodo Nominal: Pequeño - 23 kW IEC 60613 Grande - 42 kW IEC 60613

Referencia de axes: Perpendicular a la abertura facial

Este tubo es diseñado, para uso en los encajes Varex Imaging de la serie B-240H.

133595-000 Rev A 01/17

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GS-3073

Volumetric / Helical Scan Ratings IEC 60613 Tableaux des Caractéristiques Nominales de Balayage Volumétrique/Hélicoïdal CEI 60613 Volumen-/Spiralbelichtungs-Leistungdlagramme IEC 60613 Volumétrico/Clasificación Grafica del Escán/Helicoideo IEC 60613

#### 3Ø 60 Hz ■

0.7 x 0.8 Focal Spot 7 Degrees 0,7 x 0,8 Dimension Focale 7 Degrés 0.7 x 0.8 Brennfieck 7 Grad 0.7 x 0.8 De Marcas Focales 7 Grados

Volume Scan Time		AS A FUNCT			ALLOWED TUBE VING STARTING F			BE VOLTAGES	
(Seconds)	120 kV	Starting H.S. = 1 130 kV	6% 140 kV	120 kV	Starting H.S. = 3 130 kV	140 kV	120 kV	arting H.S. = 5 130 kV	50% 140 kV
1 2 4 10 20	125 125 125 125 125 125	100 100 100 100 100	100 100 100 100 100	125 125 125 125 125 125	100 100 100 100 100	100 100 100 100 100	125 125 125 125 125 125	100 100 100 100 100	100 100 100 100 100
30 40 50 60 70	125 125 125 125 125	100 100 100 100 100	100 100 100 100 100	125 125 125 125 125 125	100 100 100 100 100	100 100 100 100 100	125 125 125 125 125 100 (a)	100 100 100 100 100 100 (a)	100 100 100 100 100 (a)

#### 3Ø 180 Hz

0.7 x 0.8 Focal Spot 7 Degrees 0,7 x 0,8 Dimension Focale 7 Degrés 0.7 x 0.8 Brennfleck 7 Grad 0.7 x 0.8 De Marcas Focales 7 Grados

Volume Scan Time	AS A FUNC		ALLOWED TUBE ING STARTING F			BE VOLTAGES			
(Seconds)	120 kV	Starting H.S. = 1 130 kV	16% 140 kV	120 kV	Starting H.S. = 3 130 kV	3% 140 kV	120 kV	arting H.S. = 5 130 kV	50% 140 kV
1 2 4 10 20	175 175 175 175 175	175 175 175 175 175	150 150 150 150	175 175 175 175 175 175	175 175 175 175 175	150 150 150 150 150	175 175 175 175 175 175	175 175 175 175 175	150 150 150 150 150
30 40 50 60 70	175 175 175 175 175	175 175 175 175 175	150 150 150 150	175 175 175 175 175 150 (a)	175 175 175 175 150 (a)	150 150 150 150 125 (a)	175 175 150 (a) 125 (a) 100 (a)	175 150 150 (a) 125 (a) 100 (a)	150 150 125 (a) 100 (a) 100 (a)

#### 3Ø 60 Hz

1.2 x 1.4 Focal Spot 7 Degrees 1,2 x 1,4 Dimension Focale 7 Degrés 1.2 x 1.4 Brennfieck 7 Grad 1.2 x 1.4 De Marcas Focales 7 Grados

Volume Scan Time									
(Seconds)	120 kV	tarting H.S. = 10 130 kV	5% 140 kV	120 kV	tarting H.S. = 3 130 kV	3% 140 kV	120 kV St	arting H.S. = 5 130 kV	50% 140 kV
1 2 4 10 20	250 250 250 250 250 250	250 250 250 250 250 250	225 225 225 225 225 225 225	250 250 250 250 250 250	250 250 250 250 250 250	225 225 225 225 225 225 225	250 250 250 250 250 250	250 250 250 250 250 225	225 225 225 225 225 225 225
30 40 50 60 70	250 250 (b) 225 225 200 (a)	250 225 (b) 225 200 200 (a)	225 200 (b) 200 200 175 (a)	250 (b) 250 (b) 225 (a) 175 (a) 150 (a)	250 225 (b) 200 (a) 175 (a) 150 (a)	225 200 (b) 175 (a) 150 (a) 125 (a)	225 175 (a) 150 (a) 125 (a) 100 (a)	225 175 (a) 150 (a) 125 (a) 100 (a)	200 150 (a) 125 (a) 100 (a) 100 (a)

#### 3Ø 180 Hz

1.2 x 1.4 Focal Spot 7 Degrees 1,2 x 1,4 Dimension Focale 7 Degrés 1.2 x 1.4 Brennfleck 7 Grad 1.2 x 1.4 De Marcas Focales 7 Grados

Volume Scan Time		AS A FUNCTI	ON OF THE	AXIMUM A FOLLOWIN	LLOWED TUBE NG STARTING H	CRRENT (M EAT STORA	a) GE AND TUB	E VOLTAGES	
(Seconds)	120 kV	Starting H.S. = 16 130 kV	% 140 kV	120 kV	itarting H.S. = 33 130 kV	140 kV	120 kV Sta	rting H.S. = 5 130 kV	0% 140 kV
1	375	350	325	375	350	325	375	350	325
2	375	350	325	375	350	325	375	350	325
4	375	350	325	375	350	325	375	350	325
10	350 (b)	300 (b)	300 (b)	350 (b)	300 (b)	300 (b)	350 (b)	300 (b)	300 (b)
20	350 (b)	300 (b)	300 (b)	350 (b)	300 (b)	300 (b)	325	300	275
30	300 (b)	275 (b)	250 (b)	300 (b)	275 (b)	250 (b)	250 (a)	225 (a)	200 (a)
40	250 (b)	225 (b)	200 (b)	250 (b)	225 (b)	200 (b)	175 (a)	175 (a)	150 (a)
50	250 (b)	225 (b)	200 (b)	225 (a)	200 (a)	175 (a)	150 (a)	150 (a)	125 (a)
60	250 (b)	225 (b)	200 (b)	175 (a)	175 (a)	150 (a)	125 (a)	125 (a)	100 (a)
70	250 (a)	200 (a)	175 (a)	150 (a)	150 (a)	125 (a)	100 (a)	100 (a)	100 (a)

# Note: 1. Limits are based on maximum track rating except for the following codes: a - Limited by wardalael heat storage. b - Limited by window heating. c - Limited by finanent emission. 2. H.S. = Heat Storage kV = Tube Voltage

Remarque.

Les limites sont fonction de l'indice maximal de surface de l'anode, sauf pour les codes sulvants.

a - Limite par le stockage thermique disponible.

b - Limite par le chauffage de la fenêtre.

c - Limite par le rayonnément des l'allaments.

C - Limite par le chauffage.

L'allaments.

L'allaments.

H = Tube Voltage.

Anmerkunger:

1. Grenwerte basieren auf der maximalen Anodenobertischenlielstung mit Ausnahme der folgenden Godes:
a – Durch vertiggder Warmekapazität begrenzt.
b – Durch öffnungserwärmung begrenzt.
c – Durch Glöhrädenemission begrenzt.
kV = Röhre Spännung

Nota:

1. La clasificación de la marca maxima son imitadas, eccepto por los siquientes codigos.

codigos.

disponible.

b - Limitado por el almacenaje de calor disponible.

b - Limitado por el calor de conducción de la venfanilla.

c - Limitado por la emisión del filamento.

kV = Tubo Voltáje

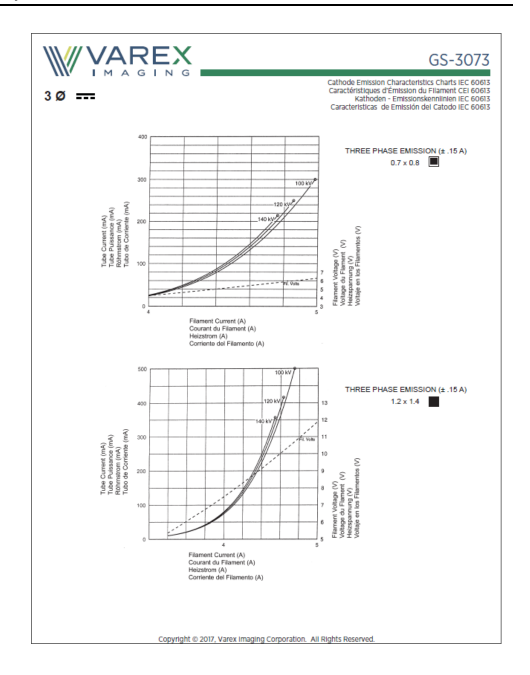
Note: Rating charts reflect maximim tube performance. Tube operation is ultimate limited by system software.

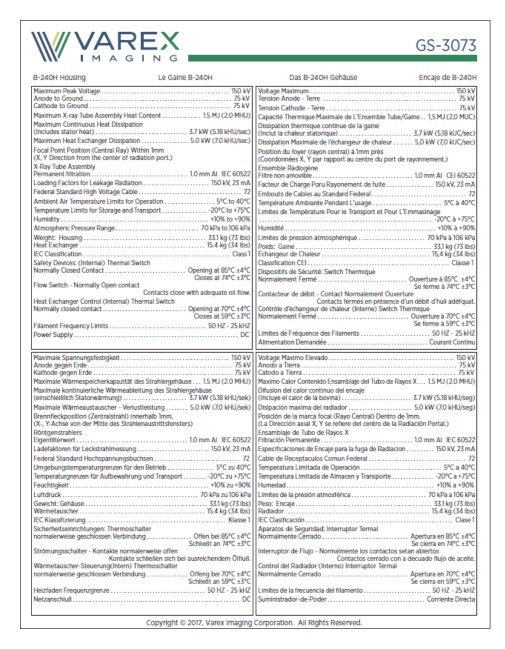
Remarque; Abaques de caractéristiques représentent des valeurs maximales. L'utilisation du tube est finalement limitée par le logiciel du système.

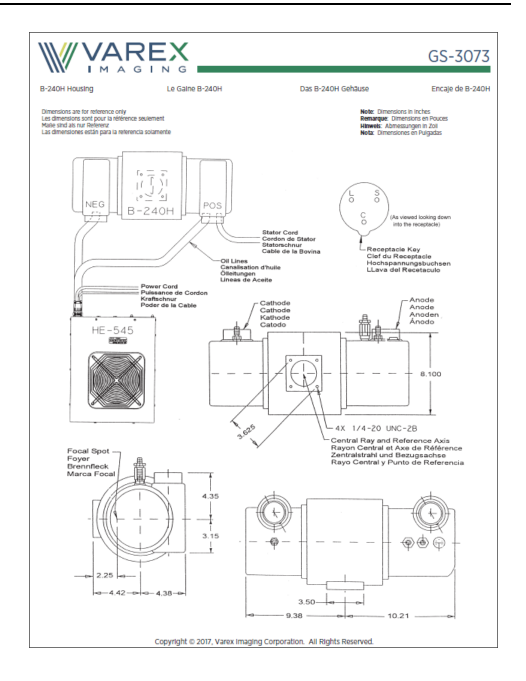
Anmerkungen:
Die leistungsdiagramme reflektieren die
maximale Röhrenleistung. Der Röhrenbetrieb ist ultimativ zu begrenzen durch die
sestamkontrolisorftware.

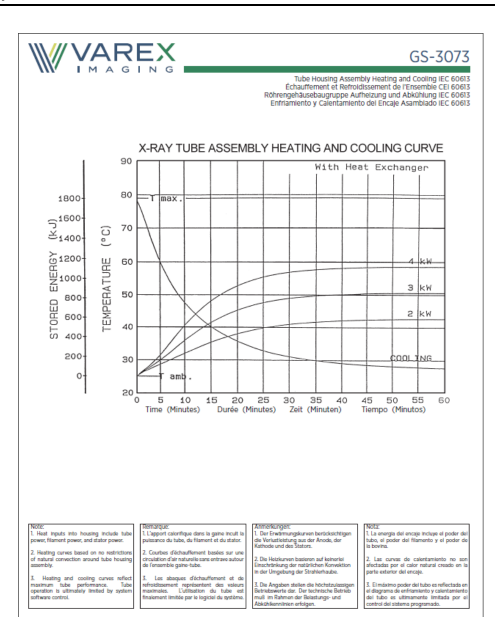
Nota; El máximo poder del tubo es reflectada en el clasificación diagrama. La operación del tubo es ultimamente limitada por el control del sistema programado.

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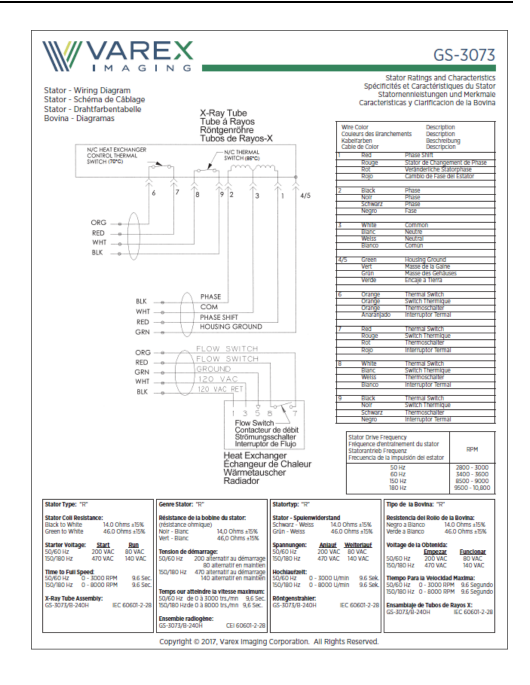


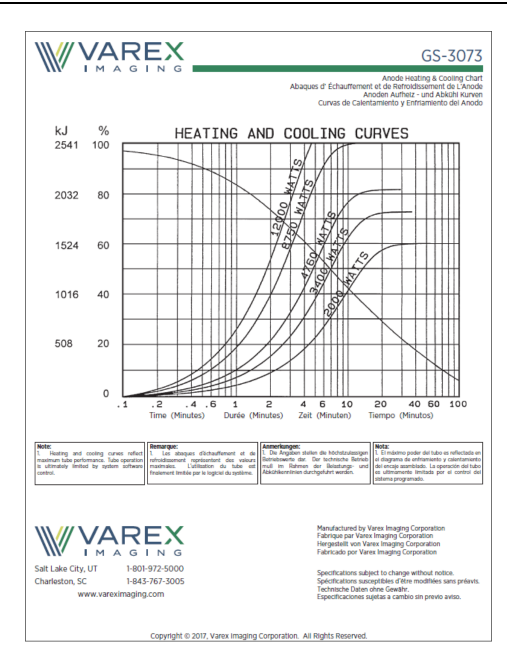






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## Appendix F Error Code

Table 69: Error code list

Error	Error code		
code	description	Popup description	Cause
0	ABORT_EXAM	Exam has been aborted. Please try again. If problem persists, contact customer service with code:	Generic message whenever a scan/exam has been terminated abnormally. User would not typically see this because a more specific error should be posted.
1	CAN_DEVICE_DISCO NNECT_CENT1	Communications fault. Please contact customer service with code:	Cannot communicate with centipede 1 device.
2	CAN_DEVICE_DISCO NNECT_CENT2	Communications fault. Please contact customer service with code:	Cannot communicate with centipede 2 device.
3	CAN_DEVICE_DISCO NNECT_ROTATE	Communications fault. Please contact customer service with code:	Cannot communicate with rotate device.
4	CAN_DEVICE_DISCO NNECT_BIB	Communications fault. Please contact customer service with code:	Cannot communicate with BIB device.
5	CAN_DEVICE_DISCO NNECT_OIB1	Communications fault. Please contact customer service with code:	Cannot communicate with OIB1 device. Note that this alone will not cause a Fault state but Start and Cancel buttons on one side of scanner will not operate.
6	CAN_DEVICE_DISCO NNECT_OIB2	Communications fault. Please contact customer service with code:	Cannot communicate with OIB2 device. Note that this alone will not cause a Fault state but Start and Cancel buttons on one side of scanner will not operate.

Error code	Error code description	Popup description	Cause
7	CAN_DEVICE_DISCO NNECT_POWER	Communications fault. Please contact customer service with code:	Cannot communicate with CCB device.
8	CAN_DEVICE_DISCO NNECT_TRANS	Communications fault. Please contact customer service with code:	Cannot communicate with Transport device.
9	CAN_DEVICE_DISCO NNECT_DCB	Communications fault. Please contact customer service with code:	Cannot communicate with DCB device.
10	CAN_DEVICE_DISCO NNECT_HVG	Communications fault. Please contact customer service with code:	Cannot communicate with HVG device.
11	HVG_LATCH_ERROR _ENABLE	High voltage fault. Please retry task. If problem persists, please contact customer service with code:	There was a problem with x-ray Enable signal.
12	HVG_LATCH_ERROR _INTERLOCK	High voltage fault. Please retry task. If problem persists, please contact customer service with code:	An interlock condition was asserted.
13	HVG_LATCH_ERROR _110_TIMER	High voltage fault. Please retry task. If problem persists, please contact customer service with code:	The 110% timer has expired and forced x-rays off.
14	HVG_LATCH_ERROR _XRT_THERM_SW	High voltage fault. Please retry task. If problem persists, please contact customer service with code:	X-ray tube thermal switch asserted.
15	HVG_LATCH_ERROR _HE_FLOW_SW	High voltage fault. Please retry task. If problem persists, please contact customer service with code:	Heat exchanger flow switch asserted.
16	HVG_LATCH_ERROR _WDT	High voltage fault. Please retry task. If problem persists, please contact customer service with code:	The watchdog timer has timed out and forced x-rays off (WD timer is controlled by the DCB firmware), this error would be unusual.

Error code	Error code description	Popup description	Cause
17	HVG_LATCH_ERROR _ARC_FAULT	High voltage fault. Please retry task. If problem persists, please contact customer service with code:	Arcs occurred.
18	HVG_LATCH_ERROR _HVG_FAULT	High voltage fault. Please retry task. If problem persists, please contact customer service with code:	A generic HVG fault condition occurred (look at HVG_ERROR_ code).
19	HVG_LATCH_ERROR _STARTER	High voltage fault. Please retry task. If problem persists, please contact customer service with code:	Anode (starter) device reported and error.
20	HVG_LATCH_ERROR _DAS_OVER_RANGE	High voltage fault. Please retry task. If problem persists, please contact customer service with code:	DAS (detector) data values are out of range.
21	HVG_ERROR_MA_R EGULATION	High voltage fault. Please retry task. If problem persists, please contact customer service with code:	There was a problem with mA regulation.
22	HVG_ERROR_KV_RE GULATION	High voltage fault. Please retry task. If problem persists, please contact customer service with code:	There was a problem with kV regulation.
23	HVG_ERROR_ANOD E_STARTER	High voltage fault. Please retry task. If problem persists, please contact customer service with code:	Anode (starter) device reported and error.
24	HVG_ERROR_INV_O VER_TEMP	High voltage fault. Please retry task. If problem persists, please contact customer service with code:	Inverter over temp condition.
25	HVG_ERROR_UNCO MMANDED_EXP	High voltage fault. Please retry task. If problem persists, please contact customer service with code:	An exposure happened without being commanded.
26	HVG_ERROR_ANOD E_OVER_VOLTAGE	High voltage fault. Please retry task. If problem persists, please contact customer service with code:	Anode over voltage condition.

Error code	Error code description	Popup description	Cause
27	HVG_ERROR_CATHO DE_OVER_VOLTAGE	High voltage fault. Please retry task. If problem persists, please contact customer service with code:	Cathode over voltage condition.
28	HVG_ERROR_ANOD E_OVER_CURRENT	High voltage fault. Please retry task. If problem persists, please contact customer service with code:	Anode over current condition.
29	HVG_ERROR_CATHO DE_OVER_CURRENT	High voltage fault. Please retry task. If problem persists, please contact customer service with code:	Cathode over current condition.
30	HVG_ERROR_FILAM ENT_OVER_CURREN T	High voltage fault. Please retry task. If problem persists, please contact customer service with code:	Filament over current condition.
31	HVG_ERROR_ARC_D ETECTED	High voltage fault. Please retry task. If problem persists, please contact customer service with code:	Arcs occurred.
32	HVG_ERROR_CURRE NT_RET_WIRE_DISC ON	High voltage fault. Please retry task. If problem persists, please contact customer service with code:	Current return wire disconnect.
33	HVG_ERROR_MA_O VER_PROG	High voltage fault. Please retry task. If problem persists, please contact customer service with code:	The actual mA was higher than programmed.
34	HVG_ERROR_KV_OV ER_PROG	High voltage fault. Please retry task. If problem persists, please contact customer service with code:	The actual kV was higher than programmed.
35	HVG_ERROR_FILAM ENT_REGULATION	High voltage failure, retry protocol, if problem persists contact customer service with code	There was a problem with filament regulation.
36	DCB_ERROR_BAD_D ETECTORS	Data acquisition fault. Please contact customer service with code:	DCB reported a bad detector condition.

Error code	Error code description	Popup description	Cause
40	RECON_PROTOCOL_ REJECTED_INVALID_ PROTOCOL_TYPE_O R_USAGE_PARAMET ER	Recon Protocol Rejected due to invalid Type or Usage parameter	Recon rejected protocol because of invalid parameters.
41	RECON_PROTOCOL_ REJECTED_INVALID_ PROTOCOL	Recon Protocol Rejected due to invalid Protocol	Recon rejected protocol because of invalid parameters.
42	RECON_PROTOCOL_ REJECTED_INVALID_ PROTOCOL_IMAGE_ COODINATES	Recon Protocol Rejected due to invalid Image Coordinates	Recon rejected protocol because of invalid parameters.
43	RECON_PROTOCOL_ REJECTED_INVALID_ PROTOCOL_ROI_CO ODINATES	Recon Protocol Rejected due to invalid ROI Coordinates	Recon rejected protocol because of invalid parameters.
44	RECON_PROTOCOL_ REJECTED_HELICAL_ QA_FAILED	Recon Protocol Rejected due to Helical QA Failure	Recon rejected protocol because of invalid parameters.
45	RECON_PROTOCOL_ REJECTED_RECON_B USY	Recon Protocol Rejected due to Recon Busy	Recon is in an invalid state to perform a protocol.
46	RECON_PROTOCOL_ REJECTED_SERIAL_LI NK_DISCONNECT_O CCURRED	Recon Protocol Rejected due to Serial Link Disconnect	Recon rejected protocol because the serial link is not connected.
47	RECON_PROTOCOL_ REJECTED_INSUFFICI ENT_MEMORY	Recon Protocol Rejected due to Insufficient Memory	Recon rejected protocol because of insufficient memory necessary to perform requested protocol.
48	RECON_PROTOCOL_ REJECTED_INVALID_ PROTOCOL_SLICE_C OODINATES	Recon Protocol Rejected due to invalid Slice Coordinates	Recon rejected protocol because of invalid parameters.
50	RECON_DATA_CORR UPTED	System has encountered an unexpected error. Please contact customer service with code:	The Recon app reported data corruption (view headers not correct).

Error code	Error code description	Popup description	Cause
51	RECON_OFFSET_CAL _FAILED	System failed to properly perform protocol calibration. Please try again. If problem persists, please contact customer service with code:	The Recon app reported Offset Cal failure.
52	RECON_AIR_CAL_FA ILED	System failed to properly calibration system. Please try calibration again. If problem persists, please contact customer service with code:	The Recon app reported Air Cal failure.
53	RECON_OFFSET_CAL _TIMEOUT	System failed to properly perform protocol calibration. Please try again. If problem persists, please contact customer service with code:	Scanner control app timed out waiting for an offset cal to complete.
54	RECON_OFFSET_PR OTOCOL_TIMEOUT	System failed to properly perform protocol calibration. Please try again. If problem persists, please contact customer service with code:	Scanner control app timed out waiting for Recon app to acknowledge a protocol request (offset or image).
55	RECON_PROTOCOL_ TIMEOUT	System is unable to perform protocol. Please try again. If problem persists, please contact customer service with code:	Scanner control app timed out waiting for a scan to complete.
56	RECON_PROTOCOL_ REJECTED	System is unable to perform protocol. Please try again. If problem persists, please contact customer service with code:	The Recon app rejected a scan protocol request.

Error code	Error code description	Popup description	Cause
57	RECON_POST_NO_S CAN_INFO	System is unable to perform post reconstruction. Please try again. If problem persists, please contact customer service with code:	The image data was not found for a Post Recon or Resend Images command (where image data could be scan_info file, .dcm files, .cor files, or entire directory).
58	RECON_POST_RECO N_FAILED	System is unable to perform post reconstruction. Please try again. If problem persists, please contact customer service with code:	A Post Recon or Resend Images command could not be run due to scanner state.
60	PREPARE_FAILED	The scanner encountered a fault preparing for scan. Please contact customer service with code:	A generic Prepare error occurred. This should be accompanied by a more specific error. See scanner log for detail.
61	DISK_PREPARE_ERR OR	Communications fault. Please contact customer service with code:	Disk subsystem reported an error during prepare.
62	CENTIPEDE_MOVE_ TIMEOUT	The scanner failed to move to the correct location. Please check for obstructions or debris on the floor that could impede the movement and try again. If problem persists, please contact customer service with code:	Scanner Control app timed out waiting for scanner to reach requested position; OR scanner did not reach required velocity for a scout or helical scan.
63	ANODE_STARTUP_TI MEOUT	X-ray power sequence fault. Please contact customer service with code:	Scanner Control app timed out waiting for anode device to report anode rotation is at speed.

Error code	Error code description	Popup description	Cause
64	DISK_NOT_IN_TICK_ MODE	Rotational speed fault. Please contact customer service with code:	Scanner Control app timed out waiting for disk to get to tick mode (during prepare for scans that specify disk rotation).
65	COLLIMATOR_POSITI ON_TIMEOUT	Failure to position collimator. Please contact customer service with code:	Scanner Control app timed out waiting for collimator to get to requested position.
66	DCB_READY_TIMEO UT	DCB communication fault. Please contact customer service with code:	Scanner Control app timed out waiting for DCB to report "ready" state.
67	HEAT_EXCHANGER_ ERROR	X-ray cooling fault. Please contact customer service with code:	Heat exchanger did not come on during prepare.
68	FILAMENT_ERROR	X-ray filament fault. Please contact customer service with code:	Scanner Control app timed out waiting for DCB to report "filament on"; OR for filament monitor to report > 10.
69	SERIAL_LINK_NOT_U P	Communications fault. Please contact customer service with code:	Scanner Control app timed out waiting for serial link to come up; OR serial link was lost before start Acq.
70	ROTATE_COMMAND _FAILED	Rotation communication error. Please contact customer service with code:	Unused error code.
71	TRANSLATE_COMM AND_FAILED	Translate fault. Please contact customer service with code:	Move command to centipede device failed.
72	COULD_NOT_START _SSP	Software initialization fault. Please contact customer service with code:	SSP software (Scanner State and Position GUI, the GUIs that are on the scanner displays) did not start up.

Error code	Error code description	Popup description	Cause
73	ROTATE_TO_ANGLE _FAILED	Rotational fault. Please contact customer service with code:	Scanner Control app timed out waiting for "Rotate to Angle" operation.
74	COULD_NOT_CLEAR _ROTATE_FAULT	Rotation fault. Please contact customer service with code:	The rotate device has reported a fault, and the fault cannot be cleared.
75	ENCODER_CONSISTE NCY_FAULT	Encoder consistency fault. Please contact customer service with code:	Unused error code.
76	INTERLOCK_TEST_FA ILED	Interlock Test Failed. Please ensure that e-stop is not engaged. If e-stop is not engaged, please contact customer service with code:	The interlock circuit either remained continuous after a Software Interlock was applied; OR the interlock circuit was broken when it was expected to be continuous.
80	DISK_SPACE_LOW	System disk space low, please contact customer service with code:	Unused error code.
81	DISK_SPACE_QUERY _FAILED	System disk space low, please contact customer service with code:	Unused error code.
82	NOT_ALL_BASE_DEV ICES_PRESENT	Base communications fault. Please contact customer service with code:	Not all devices were found on Base CAN bus (or a device has become disconnected). This should be accompanied by another error code describing which device(s) disconnected. You can look at Scanner Control GUI for a status of which devices are present.

Error code	Error code description	Popup description	Cause
83	NOT_ALL_DISK_DEVI CES_PRESENT	Disk communications fault. Please contact customer service with code:	Not all devices were found on Disk CAN bus (or a device has become disconnected). This should be accompanied by another error code describing which device(s) disconnected. You can look at Scanner Control GUI for a status of which devices are present.
90	CCB_BATTERY_OPER ATIONAL	Battery system fault. Please contact customer service with code:	CCB device reported a (not) Operational Alarm (scanning not possible).
91	CCB_BATTERY_INTE RLOCK	Battery system communication error. Please contact customer service with code:	CCB device reported an Interlock Alarm (scanning not possible).
92	CCB_BATTERY_MAI N_BREAKER	Circuit breaker has been tripped. Please reset and contact customer service with code:	Battery main breaker turned off. Scanner is running on wall power only. (Scanning not possible.)
93	CCB_LOW_BATTERY _ALARM	Low battery condition. Please charge system as soon as possible.	CCB device reported a Low Battery Alarm (scanning not possible).
94	CCB_DEAD_BATTERY _ALARM	Dead battery condition. The system is shutting down. Please charge system and report condition to customer service with code:	CCB device reported a Dead Battery Alarm (auto-shutdown of scanner is imminent) (scanning DEFINITELY not possible).
95	CCB_HIGH_BATTERY _ALARM	High battery condition has occurred. Battery charging has been disabled.	CCB device reported a High Battery Alarm

Error code	Error code description	Popup description	Cause
96	CCB_OVERCGARGED _BATTERY_ALARM	Please power down and unplug system and contact customer service immediately with code:	CCB device reported an Overcharged Battery Alarm. This is beyond a High Battery warning and is serious. The system should be turned off and unplugged immediately.
97	CCB_BATTERY_HIGH _TEMP_ALARM	Battery system fault. Please contact customer service with code:	CCB device reported a High Temp Alarm (scanning not possible).
98	CCB_BATTERY_OVER _TEMP_ALARM	Over temperature battery condition. Please power down and unplug system and contact customer service immediately with code:	CCB device reported an Over Temp Alarm. This is beyond a High Temp warning and is serious. The system should be turned off and unplugged immediately.
99	CCB_BATTERY_MEA SUREMENT_ERROR	Battery system fault. Please contact customer service with code:	CCB device reported a Measurement Error Alarm (scanning not possible)
100	CCB_BATTERY_IMBA LANCE_WARNING	Battery system fault. Please contact customer service with code:	One or more battery voltage levels are not the same as the others. No action required. System will try to correct itself.
110	RECON_PROTOCOL_ REJECTED_INVALID_ PROTOCOL_USAGE_ PARAMETER	Reconstruction fault. Please contact customer service with code:	Recon received invalid usage parameter in protocol.
111	RECON_PROTOCOL_ REJECTED_INVALID_ NUMBER_OF_VIEWS _PARAMETER	Reconstruction fault. Please contact customer service with code:	Recon received invalid number of views in protocol.
112	RECON_PROTOCOL_ REJECTED_RUN_DM A_SETUP	Reconstruction fault. Please contact customer service with code:	Recon failed to initialize DMA in preparation for scan.

Error	Error code	Danie danietiae	Commo	
code	description	Popup description	Cause	
	RECON_PROTOCOL_	Reconstruction fault.	Recon received	
113	REJECTED_UNDEFIN	Please contact customer	undefined usage	
	ED_USAGE	service with code:	parameter in protocol.	
	RECON_PROTOCOL_	Reconstruction fault.		
114	REJECTED_INVALID_	Please contact customer	Unused error code.	
	RAW_DATA_REPLAY	service with code:		
	RECON_PROTOCOL_	Reconstruction fault.		
115	REJECTED_FILES_RE	Please contact customer	Unused error code.	
	TRIEVE_FAILED	service with code:		
	RECON_PROTOCOL_	Reconstruction fault.	Recon received	
116	REJECTED_INVALID_	Please contact customer	incorrect structure	
110	PARAMETER_STRUC	service with code:	size.	
	TURE_SIZE	Service with code.	3120.	
	RECON_PROTOCOL_	Reconstruction fault.		
117	REJECTED_INVALID_	Please contact customer	Unused error code.	
117	PROTOCOL_FLASH_I	service with code:	onasca ciror coac.	
	O_CMD	Service with code.		
	RECON_PROTOCOL_	Reconstruction fault.		
118	REJECTED_PREPARE	Please contact customer	Unused error code.	
110	_AND_PRIME_POST	service with code:		
	_RECON_USAGE	Service with code.		
	RECON_PROTOCOL_	Reconstruction fault.	Recon received a	
119	REJECTED INVALID	Please contact customer service with code:	protocol while still	
	POST_RECON_STATE		processing previous	
		Service With code.	protocol/post recon.	
	RECON_PROTOCOL_	Reconstruction fault.	Recon received	
120	REJECTED_INVALID_	Please contact customer	incorrect structure	
	MESSAGE_BODY_LE	service with code:	size.	
	NGTH		-	
	RECON_PROTOCOL_	Reconstruction fault.		
121	REJECTED_INVALID_	Please contact customer	Unused error code.	
	RELOAD_PARAMETE	service with code:		
	R_FILES			
	RECON_PROTOCOL_	Reconstruction fault.	Recon was sent an	
122	REJECTED_INVALID_	Please contact customer	unsupported	
	UNSUPPORTED_CO	service with code:	command from	
	MMAND		scanner control.	
	RECON_PROTOCOL_	Barrada dia 6 h	Recon was sent an	
422	REJECTED_INVALID_	Reconstruction fault.	invalid helical filter	
123	HELICAL_FILTER_KER	Please contact customer	kernel from	
	NEL TYPE	service with code:	workstation/scanner	
	1455_1115		control.	

Error	Error code	Popup description	Cause
code	description		04450
124	RECON_PROTOCOL_ REJECTED_INVALID_ NUMBER_OF_HELIC AL_IMAGES_FOR_W INDMILL	Reconstruction fault. Please contact customer service with code:	Recon received an invalid number of images for windmill.
125	RECON_PROTOCOL_ REJECTED_GPU_FAIL ED_TO_START	Reconstruction fault. Please contact customer service with code:	Recon failed to initialize GPU during preparation for scan.
126	RECON_PROTOCOL_ REJECTED_RECON_B USY	Reconstruction fault. Please contact customer service with code:	Recon received a protocol while still processing previous protocol/post recon.
130	RECON_AIR_CAL_FA ILED_NON_AIR_IMA GE	Reconstruction fault. Please contact customer service with code:	Air image above threshold for air calibration.
131	RECON_AIR_CAL_FA ILED_SEND_EVENT	Reconstruction fault. Please contact customer service with code:	Air calibration failed to be performed.
132	RECON_AIR_CAL_IM AGE_EXCEEDS_THRE SHOLD	Reconstruction fault. Please contact customer service with code:	Air image above threshold for air calibration.
133	RECON_AIR_CAL_FA ILED_NO_VIEW_DAT A	Reconstruction fault. Please contact customer service with code:	No view data received during an air calibration.
134	RECON_AIR_CAL_FA ILED_CORRUPTED_V IEW_DATA	Reconstruction fault. Please contact customer service with code:	Corrupted views received during air calibration.
135	RECON_OFFSET_CAL _FAILED_SEND_EVE NT	Reconstruction fault. Please contact customer service with code:	Offset calibration failed to be performed.
136	RECON_OFFSET_CAL _FAILED_NO_VIEW_ DATA	Reconstruction fault. Please contact customer service with code:	No view data received during an offset calibration.
137	RECON_OFFSET_CAL _FAILED_CORRUPTE D_VIEW_DATA	Reconstruction fault. Please contact customer service with code:	Corrupted views received during offset calibration.
138	RECON_OFFSET_CAL _FAILED_BAD_REFE RENCE	Reconstruction fault. Please contact customer service with code:	Offset calibration failed due to bad reference.

Error code	Error code description	Popup description	Cause
139	RECON_OFFSET_CAL _80_PERCENT_BAD_ REFERENCE	Please contact Customer Service immediately and run a Quality Assurance (QA) Phantom test to verify image quality. Error code:	80% of reference detector values are above the acceptable threshold during an offset cal.
140	UPS_LOW_BATTERY _ALARM	Workstation low battery condition. Please charge system as soon as possible.	UPS device reported a Low Battery Alarm.
141	UPS_DEAD_BATTERY _ALARM	Workstation dead battery condition. The Workstation is shutting down. Please charge cart and report condition to customer service with code:	UPS device reported a Dead Battery Alarm.
142	UPS_HIGH_BATTERY _ALARM	Workstation high battery condition has occurred. Cart battery charging has been disabled.	UPS device reported a High Battery Alarm.
143	UPS_OVERCHARGED _BATTERY_ALARM	Please power down and unplug workstation cart and contact customer service immediately with code:	UPS device reported an Overcharged Battery Alarm. This is beyond a High Battery warning and is serious. The cart should be turned off and unplugged immediately.
144	UPS_BATTERY_HIGH _TEMP_ALARM	Workstation battery system fault. Please contact customer service with code:	UPS device reported a High Temp Alarm.
145	UPS_BATTERY_OVER _TEMP_ALARM	Workstation over temperature battery condition. Please power down workstation, unplug cart and contact customer service immediately with code:	UPS device reported an Over Temp Alarm. This is beyond a High Temp warning and is serious. The cart should be turned off and unplugged immediately.
146	UPS_BATTERY_MEA SUREMENT_ERROR	Workstation battery system fault. Please contact customer service with code:	UPS device reported a Measurement Error Alarm.

Error code	Error code description	Popup description	Cause
147	UPS_BATTERY_IMBA LANCE_WARNING	Workstation battery system fault. Please contact customer service with code:	One or more battery voltage levels are not the same as the others. No action required. UPS will try to correct itself.
148	UPS_CHARGER_FAU LT	Workstation battery system fault. Please contact customer service with code:	UPS device reported a Charger fault.

### Appendix G Revision History

Table 70: Workstation (only) revision history before combined with BodyTom Elite User Manual (below)

Revision	ECO number	Effective date	Author	Changes
00	ECO- 002019	None supplied	Keith Almeida	Releasing 1- 00132-060 User Manual, WS CART NL4000
01	ECO- 002266	None supplied	Keith Almeida	Added "Installing a UPS" instructions
02	ECO- 002735	None supplied	Rich DeSalvo	Removed "Cart Assembly" instructions Updated TOC, Figures, Tables Updated cart info
N/A	N/A	N/A See Table 71, below.	Cynthia Crow	Consolidated the BodyTom Workstation User Manual with the BodyTom User Manual. See Table 71, below.

Table 71: Revision history

Table 71: Revis	,			
Revision	ECO number	Effective date	Author	Changes
00	ECO-001673	2012/01/27	Christofer Krueger/Mike Limoli/Ibrahim Bechwati	New Release
01	ECO-001877	2012/11/25	Christofer Krueger	Correct minor edits uncovered during translations' review and pilot installations. Also needed to add 3 <sup>rd</sup> edition and HC requirements
02	ECO-002003	2013/12/19	Christofer Krueger	Update to meet IEC 60601-1 3 <sup>rd</sup> ed, 60825-1, and CFDA Requirements. Error codes, additional warnings, focal spot, CTDI <sub>100</sub> results, scatter values, product label location, etc.
03	ECO-001992	2014/04/03	Keith Almeida	Updated to software release 01.06
04	ECO-002326	2015/01/28	Keith Almeida	Releasing NL4000 Software Version 01.07
05	ECO-002441	2015/04/22	Keith Almeida	Releasing NL4000 Software Version 01.07.01
06	ECO-002869	2016/04/18	Keith Almeida	Update for consistent formatting and Samsung branding. Added 1.07 Error codes Reduced use of pronouns Inclusion of Information Bulletins CSB-00006 and CSB-00007.
07	ECO-002915	2016/07/13	Keith Almeida	Addition of Slab, Kernel, and Floor Flatness Certification corresponding with version 1.08 of the software.
08	ECO-002961	2016/10/08	Ross Caisse	Removing references to bolus tracking for software release 1.08.01.
09	ECO-003015	2016/11/18	Ross Caisse	Adding references to bolus tracking for software release 1.08.02.

Revision	ECO number	Effective date	Author	Changes
10	ECO-003600	2017/09/14	Cynthia Crow	Changed branding from BodyTom to BodyTom Elite. Applied new template/styles, step-by-step procedures, and workflow for improved audience experience. Consolidated the previous (Revision 02) BodyTom Workstation User Manual (1- 00132-060) with the BodyTom User Manual, Revision 10 (1- NL4000-060).
11	ECO-003712	2018/04/13	Derek Decoux	Updated the tube heat capacity icon description with correct ranges for different colors. Added Essential Performance factors to the preface. Added a Functional Earth Symbol (Table 55)
12	ECO-004280	2018/11/01	Christofer Krueger	Adding a note above Table 13 on EMC Characteristics. Modified Table 14 to detail the correct standard parameters. Changed title to include "NL4000 BodyTom". Added "BodyTom" to the "About this Manual" section. Table $4 \sim 0.25\Omega$ to $0.105\Omega$ .
13	ECO-004754	2020/03/03	Christofer Krueger	Added details to comply with IEC 60825-1, 3 <sup>rd</sup> ed. Table 2 and Laser Safety section of Chapter 1. EU MDR updates: Added Serious Injury, Electronic Copy and Clinical Benefit notes to the preface

Revision	ECO number	Effective date	Author	Changes
14	ECO-004919	2020/06/17	Christofer Krueger	Added Laser warning statement from the FDA.  Updated for new marking plate. Removed broken bookmarks. Corrected the workstation temp range. Modified an incomplete sentence in table 58. Updated glossary to remove the word "withring". Bug 5304. Added new statement and Figure 203 on Geometric efficiency for being less than 70%.
15	ECO-005032	2020/09/015	Stephen Dunn	Updated Geometric Efficiency wording, added ACR information
16	ECO-005301	2021/02/05	Gina Cunsolo	Updated to remove reference to BodyTom. Added symbols to Table 2 to reflect all symbols on the label.
17	ECO-005794	2022/02/23	Karen Reed/Keith Kaser/Stephen Dunn	Added statement to contact CS if there is an adverse event under Contact Info Sec.  Added reference for eIFU in "About this user manual" Section Updated Symbols Table for clerical purposes.  Updated Table 9 to reference both BodyTom and BodyTom Elite PNs.  Updated WS Cart weight in Table 11 for correct weight.  Removed old references of standards already mentioned under Compliance Statement.  Added Table 42 (p. 244) and 43 (p. 245).

Updated images to remove older logos. Removed Floor calibration. Remove Surgical clamp details. Added Interventional scanning information. Updated Cleaning instructions to remove references from Chapter One and move to Cleaning Chapter 14. Updated Pendant information. Modified multiple bulleted tables to make them actual tables. Modified Dose Alert information. Added detailed AEC information to Appendix D. Updated entire document Font to Calibri 12. Modified Appendix C to include only a sample of the reference protocols. Modified all line spacing to 1.0. Updated International Customer Service contacts to make them more user friendly. Made many changes to wording and grammar throughout document. Updated all images to remove red callouts and arrows and replaced with gold. Removed small filament information from Focal Spot section of Chapter One. Removed E-Stop instructions from Chapter 1 and moved to Chapter 3. Added scanner Battery Capacity figure to Chapter 1. Added X-Ray tube capacity information to Chapter 1. Removed Floor flatness certification information from Chapter 1. Removed multiple icons/orbs from text of document. Removed redundant 'Saving a Preset' under Applying Windowing Presets in Chapter 5. Removed AEC information from 'Creating a new protocol' section

Revision	ECO number	Effective date	Author	Changes
				in Chapter 6 since it is covered in detail in 'Scanning with special features' of Chapter 9. Added 'Editing and existing protocol' to Chapter 6. Updated 'Starting Quality Assurance' section of Chapter 7 to include better instructions for this process. Modified all contents of Chapter 10 to match appearance of items on the GUI. Updated all sections of Chapter 11 'Viewing Images' to make them clearer. Removed redundant 'Applying kernels in Post Reconstruction' section of Chapter 12. Updated 'Using Universal Transfer Board' in Chapter 13 to include more information. Updated Footers of pages 2-24. Added information to Site Specifications related to floor supporting product weight. Updated workstation weight. Updated Figure 5. Updated Figure 2, 12, 33, 38, 40 and 399 to remove BodyTom reference.
18	ECO-006198		Keith A. Kaser	Corrected multiple spacing issues. Added Brain CT Perfusion scanning instructions under the "Scanning with Special Features" section and moved Viewing Images in the CTP Viewer from the Viewing Images section to the new CT Perfusion section mentioned above.
19	ECO-006502		Keith A. Kaser	Updated 'Intended use of system' section. Replaced Figure 232 to change highlight section from red to gold. Replaced Figure 345 to change arrow from red to gold.

Revision	ECO number	Effective date	Author	Changes
				Updated Laser Section – changed numbers from Red to Green and Blue to match Figure 5 colors. Replaced Figure 343 to change highlight section from red to gold. Replaced Figure 392 to change highlight section from red to gold. Replaced Figure 398 to change arrows from red to gold.
				Updated Table 21 and 65 to correct direction of travel for Pendant buttons.

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