

User Manual

Trauma DR plus

Rev.: 05-11-2025



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List of revisions

Equipment	Trauma DR plus
Document	User Manual
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01	01-01-2017	Added C-arm illustrations and prepacked protocols
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04	05-11-2025	Changes regarding art.22 (MDR). Removed Fax nr Updated detector section to new detector type incl. BiAA and iNR

Head Office, The Netherlands

System Packer



Oldelft Benelux BV

Wiltonstraat 41
3905 KW VEENENDAAL
Tel. (31) 318 584 400
E-mail: info@delftdi.com
Internet: <http://www.delftdi.com>

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Preface

Identification Manual

Manuals are identified by a part number.

ODB-100-5040ENG-04



The part number is printed at the bottom of the title page. Numbers printed on other pages are for internal revision control and may differ.

CE Classification

Manuals made for products that have been CE (Conformité Européenne) marked have the CE mark on the front page. The number behind the CE Mark defines the responsible notified body. Oldelft Benelux B.V. systems are classified according to IEC 60601-1 as Class II, type B.

Oldelft Benelux B.V. systems meet the relevant provisions of the European MDR (Medical Device Regulation); this is based on conformity of the products and the quality system according to ISO 13485:2016.

The system meets the relevant provisions of the European Medical Device Regulation (EU) 2017/745 (MDR).

The Easy DR designed under the MDR art.22 can still have MDD certificates for the components due the transition period.

Symbols

Symbol	Meaning	Remarks
Location: Near or on the type identification plate. 	Symbol for separated collection of electrical and electronic equipment per Directive 2002/96 of the European Parliament and the Council of the European Union (Directive on Waste of Electrical and Electronic Equipment - WEEE). If applicable to the type of device, it indicates legally imposed obligations within EU member states, Iceland, Norway and Switzerland when the equipment is disposed of, at the end of its life-time.	Owners of marked equipment should contact the organization that imported the equipment into their country, when they want to dispose of the equipment, at the end of its lifetime. The Directive prioritizes re-use of equipment over re-use of components over re-use of materials over disposal as waste. Article 5 part 2d allows producers to decline the return of any used equipment that is or may be biologically or radiologically contaminated.
	Date of manufacture	
	SN: Serial number UDI: Unique Device Identifier	
	This mark indicated that this is CLASS I Equipment according to EN60601-1.	
	This mark indicated that this is a Type B Applied Part according to EN60601-1.	
	Consulting the accompanying documents is a mandatory action.	
	General warning sign	Placed together with a supplementary symbol or text. The text associated shall be an affirmative statement (i.e., a safety notice) describing the principal risk(s) foreseen (e.g. "Causes burns", "Risk of explosion", etc.).
	The CE Mark is a declaration by the manufacturer that the product complies with the requirements of the applicable European Union (EU) Medical Device Regulation (MDR) and that the product has been subject to conformity assessment procedures as provided in that regulation.	

Conventions

The Trauma DR plus has been designed to meet all safety requirements applicable to medical equipment. However, anyone using the Trauma DR plus must be fully aware of potential safety hazards. The information contained in this chapter is provided to help users to operate the Trauma DR plus safely.

Throughout this manual certain conventions are used. These are Warnings, Cautions and Notes. They provide a means of prioritizing information to be brought to the attention of the user. They are given as follows:

 WARNING:	Indication of an important warning that is to be obeyed to eliminate the risk of personal injury or an incorrect clinical diagnosis.
CAUTION:	Indication of important information to avoid serious system equipment damage.
Note:	Additional information for reader understanding or simplification of a task.

Precautions

The user is expected to use the product (soft and hardware) in accordance with the instructions given in this manual, which must be read before the system is used. Any unauthorized deviation from the procedures laid down in this manual can affect the contractual obligations between purchaser and vendor.

 WARNING:	This product was designed and manufactured to ensure maximum safety of use. It should be installed and serviced in strict compliance with safety instructions described in this Document.
 WARNING:	Never attempt a shortcut to procedures which may degrade the safety considerations of the system.
 WARNING:	Class 1 MD: To avoid the risk of electric shock, this equipment must only be connected to a supply mains with protective earth.
 WARNING:	The user shall take care that potential electromagnetic or other interferences between the unit and other equipment is avoided. If necessary certain actions must be taken.
 WARNING:	Only trained and qualified personal are allowed to operate the Trauma DR plus system.

 WARNING:	Do not modify any part of Oldelft Benelux supplied hardware, software or software configuration. Any modification, installation or running an application that has not explicitly been approved by Oldelft Benelux may compromise patient safety and will result in termination of all warranty and service contracts. a. If this equipment is modified, repaired or maintained, appropriate inspection and testing must be conducted to ensure continued safe use of the equipment. b. Do not leave problems unattended that may affect the safety of the product. In case you have doubts concerning the Trauma DR plus's save operation, call the service number of your vender for further instructions. c. The Trauma DR plus should be cleaned regular to prevent the accumulation of dust. The contact surface should be cleaned after each patient for hygienic reasons.
 WARNING:	The Oldelft Benelux products are designed for use and maintenance using only parts available from Oldelft Benelux. No liability for failure can be accepted by Oldelft Benelux through the use of parts obtained from other suppliers unless written permission is obtained from Oldelft Benelux. Copyright All rights reserved. No part of this manual may be reproduced or transmitted in any form whatsoever, including photocopying and recording without written permission of the copyright holder: Oldelft Benelux B.V., Wiltonstraat 41, 3961 KW VEENENDAAL, The Netherlands. Concerning this Publication Information contained in this publication is subject to change without notice. Whenever the equipment is delivered with a documentation medium such as a CD or DVD, the user shall always check the medium for latest information. This information is part of Release Notes and Document Upgrade files. Neither Oldelft Benelux B.V. and its subsidiaries nor anyone else who has been involved with the development, production or delivery of the documentation be liable for any special, incidental or consequential damages, whether based upon breach of warranty, breach of contract, negligence, strict liability in tort or any other legal theory. This manual was originally drafted in the English language.

 WARNING:	Do not use the Trauma DR plus if the system is damaged or does not function properly. Do not leave problems unattended that may affect the safety of the product. In case you have doubts concerning the Trauma DR plus save operation, call the service number of your vender for further instructions.
Note:	Any serious incident with the user and/or patient that has occurred in relation to the device should be reported to the manufacturer and the competent authority of the Member State in which the user and/or patient is established.

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1 Introduction

1.1 About this manual

The Trauma DR plus is a multifunctional X-ray unit. In this manual it is assumed that the operator is a professional and trained X-ray user, who has gained sufficient knowledge to operate the Trauma DR plus safely.

This user manual is a guide for the operator of the Trauma DR plus multifunctional X-ray unit. In this manual it is assumed that the operator is a professional and trained X-ray user, who has gained sufficient knowledge to operate Trauma DR plus system safely.

In particular this document describes the daily operation and interpretation of the user interface for the Trauma DR plus.

1.2 Classification

Medical Devices classes (also known as **risks classes**) specify the risk class depending on the vulnerability of the human body and the potential risk associated with the medical device.

Trauma DR plus is classified according to Class II type B (bear a medium to high risk).

Appliance classes (also known as **protection classes**) specify measures to prevent dangerous contact voltages on unenergized parts, of an electronic device.

	This mark indicated that this is CLASS I Equipment according to EN60601-1.	Equipment with two levels of protection against electrical shock. - Basic insulation of live parts and touchable conductive casing. - Earth connection of touchable conductive casing.
	This mark indicated that this is a Type B Applied Part according to EN60601-1.	No Electrical contact with patient and the patient can be immediately released from the Wall Stand or Patient Table.

The system meets the relevant provisions of the European Medical Device Regulation (EU) 2017/745 (MDR).

1.3 Reference manuals

Note:	This document complements the reference manuals provided with this system and cannot replace these manuals. The user should therefore take note of the reference manuals, especially the safety rules before using the system. When this document and the reference manual differ, the reference manuals are considered as leading.
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- GMM: User Manual Trauma DR Ceiling Stand
- Canon: Operation Manual CXDI Control Software NE
- Canon: User's Manual CXDI-720C Wireless
- Canon: User's Manual CXDI-420C Wireless
- Canon: User's Manual CXDI-820C Wireless
- CPI: Operation Manual CPI "MINI-CONSOLE" CONTROL UNIT
- CPI: Operation Manual for Canon Single Console CXDI NE

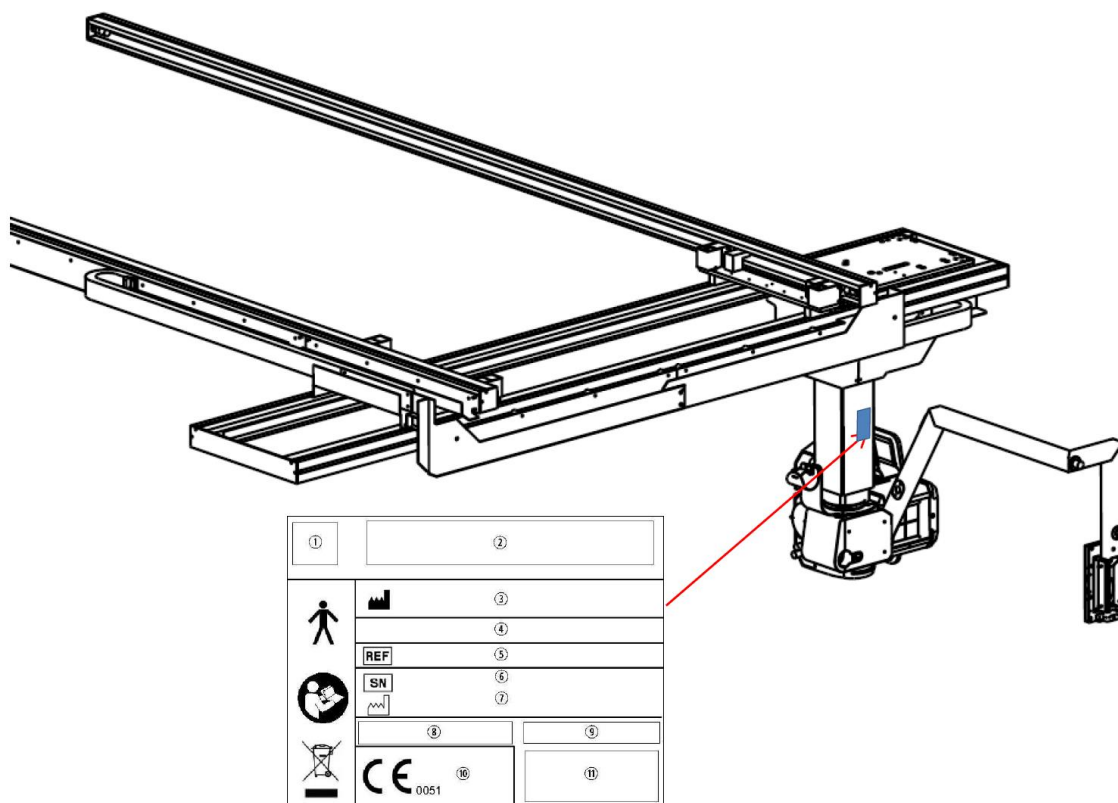
1.4 Identification Label

Label

Model / Type: Trauma DR+		Generator: Input voltage: 3 x 400 Vac 60A Frequency: 50/60 Hz Max current: 125A Power supply stand: Powered by generator, 230 Vac, 50 Hz, 6.3A.	 Oldelft Benelux A CANON COMPANY 
 (01)18720256816413 (11)yyymmdd (21)BUCxxxx	 mm-yyyy  BUCxxxx		
	Oldelft Benelux BV, Wiltonstraat 41, 3905KW, Veenendaal, The Netherlands.		

Identification label Trauma DR plus Rev.2024-07-03

Location



Identification label position Trauma DR plus Rev.2024-07-03

1.5 General description/indications for use

The Trauma DR+ is a multifunctional X-ray system equipped with a ceiling suspended trauma arm.

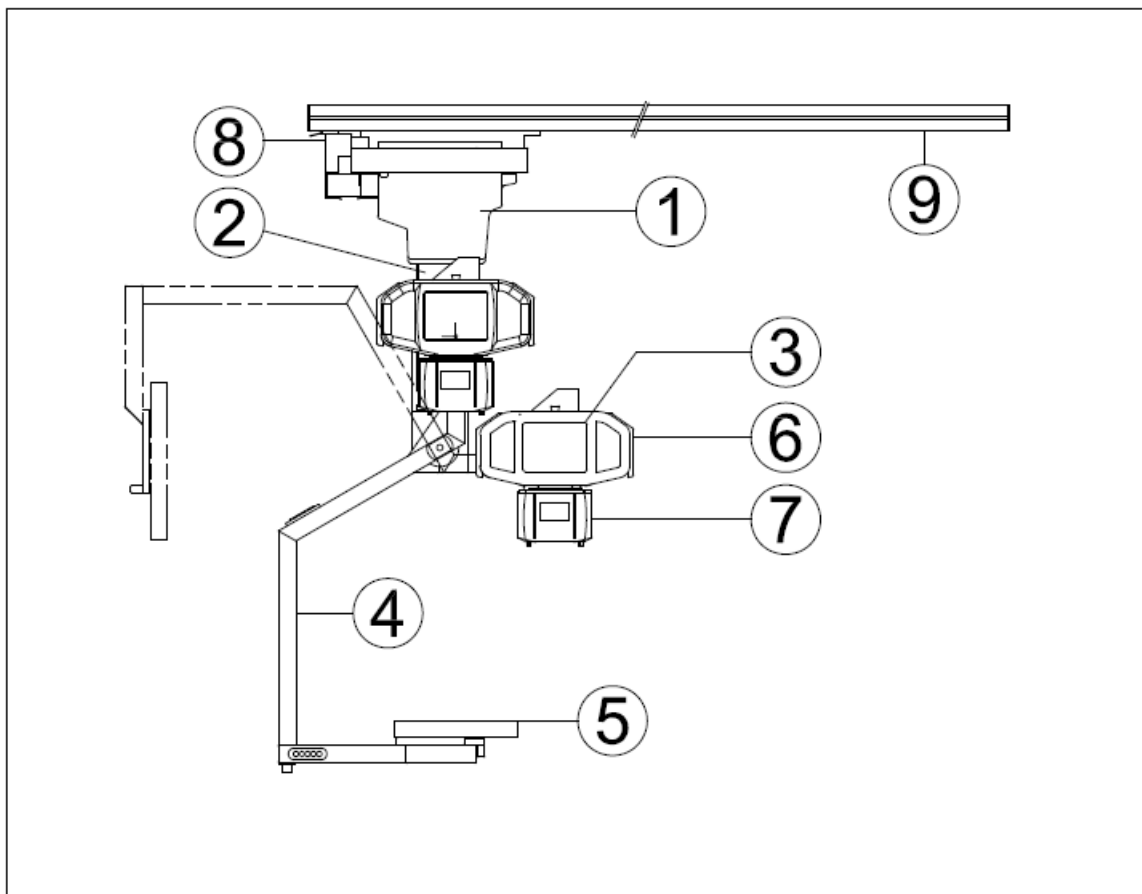


Figure 1: Trauma arm

1	Carriage
2	Telescope
3	Handlebar
4	C-Arm
5	Detector
6	X-ray tube
7	Collimator unit
8	Transversal bridge
9	Longitudinal support rails

The Control Computer is controlling the X-ray line. Depending of the Patient's body part the Control Computer configures the generator, X-ray tube and detector. When Patient data is filled in, the patient, X-ray tube and detector are positioned the operator can take an image by pressing the exposure button.

Depending on the configuration the Trauma DR plus system may contain the following sub-systems:

- The Trauma DR plus
- X-ray equipment and detectors (separate manuals)
- The Control Computer (separate manuals)
- Optional PACS and Diagnostic Viewing Station (separate manuals)
- Optional printer (separate manuals)
- Optional UPS/battery system (separate manuals)

1.6 Intended Purpose and Patient population

- Intended Purpose: The Trauma DR plus is a diagnostic X-ray system intended for imaging anatomical features.
- Intended indication: General Radiography.
- Intended Users: Trained professional radiographers.
- Intended environment: Dedicated X-ray room, in medical institutions.
- Patient Population: There are no restrictions in patient population.

1.7 Physical description

In this paragraph only the Trauma DR plus main hardware items are described. These are:

- Trauma arm
- Control Computer
- X-ray Tube
- X-ray Generator and Control Panel
- Collimator

The Trauma DR plus operates without X-ray film: the X-ray image is captured with the X-ray detector and stored in digital memory. Digitized images are stored as DICOM 3.0 compatible files on a local hard disk (short term) and automatically via a network link to the PACS and diagnostic viewer station.

After positioning the patient, the operator prompts the Trauma DR plus to make an X-ray exposure. Before the actual X-ray exposure can be made the patient's data have to be entered on the control computer. After patient data is entered the X-ray exposure is taken.



1.7.1 Trauma Arm

The Trauma Arm Holds the X-ray tube, collimator and detector. The Trauma Arm has many freedoms of movements to position the X-ray tube in almost all positions relative to the Patient Table. This allows for a wide range of radiological exams. The Trauma Arm is designed in such a way that in all positions patient placement is flexible and no cables or frames are in the way.

1.7.2 Control computer

The control computer is used to operate the detector and X-ray unit. The system comes with a wide range of preconfigured protocols for an optimal image quality. By selection of a protocol the correct X-ray settings (kV, mA and mS) are automatically set. The Automatic Exposure Control (AEC) will give the most optimal image quality at the lowest possible patient dose. The Control computer is also used to temporarily store the X-ray images. It is interfaced via a network to the PACS and Diagnostic Viewing stations.

The control computer is the controlling device for the Trauma DR plus rather than a means for advanced diagnostics, which would require very high resolution/brightness as in fast viewing stations. These are beyond the scope of this manual.

The control computer offers a graphical user interface to perform the following functions:

- Input of patient data, either directly (touch screen) or via import from peripheral systems (network, barcode scanner)
- Control of X-ray detector (Data acquisition, post processing and display)
- Control of X-ray unit (kV, mA and mS settings)
- Database management for patient data and related X-ray images
- Export of digital X-ray images to peripheral systems (network)

Radiation on/off switching is controlled by the Control computer and X-ray generator interface but always executed by the two stage exposure switch. Generator technique settings are controlled by the control computer. However Radiographers should follow well-known guidelines and procedures for operating the X-ray generator and are supposed to have read and understood the manufacturer's instructions for that device.

**Advice to consult the accompanying documentation:**

- Operation manual HF generator.

1.7.3 X-ray Tube unit

The X-ray tube unit consists of an X-ray tube, a filter unit and a collimator unit.

The filter unit contains a rotating disc with four types of X-ray filters.

The Collimator is used to collimate, limits the X-ray field to the Field of interest.

1.7.4 X-ray generator and control panel

The X-ray generator has its own documentation.

	Advice to consult the accompanying documentation: • Operation manual HF generator.
---	---

A two-step radiation exposure switch is connected to the (mini) generator interface panel. The following functions are available on the generator control panel:

- Switching the X-ray system ON or OFF
- Exposure switch and indicator (audible). Starting X-ray radiation.

1.7.5 Collimator unit

The collimator unit comprises an adjustable diaphragm and a field illumination indicator.

The adjustable diaphragm has two setting knobs: one to adjust two vertical shutters for symmetrical lateral adjustment, the other to move the horizontal shutters to adjust vertical field size.

The field illumination indicator delineates the total extent of the field to be exposed. It can be switched on and off by a push button. If left on, the light will turn off automatically.

1.7.6 Exposure interval

The actual exposure starts when the two stage exposure switch is pressed. The AEC automatically switch of the radiation as soon as the desired dose is measured by the system.

2 Safety precautions

2.1 General Warnings

The Trauma DR plus and X-ray line are equipped with safeguards to prevent injury to operator and patient during normal use and in case of system malfunctions. For the safeguards of the system parts refer to the corresponding documentation.

See par.: Reference manuals



The Trauma DR plus is a combination of subsystems that are built to work together.

All subsystems have accompanying technical documentation we advise to consult before starting repair or maintain the system.

In case of equipment malfunction, bring the patient into safety and seek the assistance of a qualified technician.

CAUTION:

Make sure that the operator always will remain in audio and visual contact with the patient when performing the X-rays in the safe zone.

2.2 Radiation precautions

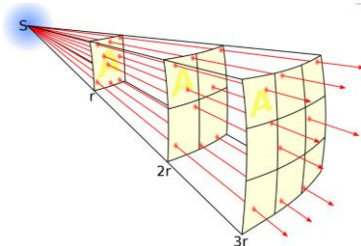
The X-ray equipment must be used in the radiological diagnosis by qualified personnel, who knows exactly what the risks are inherent in systems that use ionizing radiation and is able to take the necessary protective measures.

- The regulations and laws in force in the various countries for the type of equipment considered to be observed by the operator/user.
- The X-ray equipment must be operated only in areas designated for x-ray systems and for medical use (refer to local regulations).
- Operator and staff must only control the x-ray equipment from protected areas. There are no x-ray procedures requiring the Operator or staff to be in the x-ray room. All procedures can be operated from the control room (protected area), protected by lead window, lead walls or equivalent.
- To exclude the possibility of expose for patients to levels where deterministic effects may occur, an overall backup timer or the Body Part backup timer terminates the irradiation.

ALARA is a philosophy of excellence used in one's day-to-day work with radiation sources. It is when one strives to keep one's radiation exposure **As Low As Reasonably Achievable**.

In general, the dose for Operators and Staff can be reduced by the following means:

- Protection by distance
Square law: At large distances from the source, the dose is distributed over larger and larger spherical surfaces as the distance from the source increases. After all, the dose decreases square with the distance.



- Protection by shielding
All procedures can be operated from the control room (protected area), protected by lead window, lead walls or equivalent.

Patient Dose

When not using the AEC it is important to maximize the SID to minimize the patient radiation dose. The square law applies also for the patient dose as explained above.

2.3 Biocompatibility

To prevent infections, clean all parts of the equipment that are touched by the patient, before the next patient is examined. Do this according the instructions in par.: *Cleaning*

2.4 Moving the system

The Trauma DR plus Ceil stand can be moved by hand force or can also be moved motorized.

In order to prevent any possible collision, maintain a free area around the units. The manuals, should be studied to learn about the working zones and the limits of the device and to understand the risks of collisions when the unit is used with other devices and risks when used within the environment in which it is located.

 WARNING:	When moving the system by hand or motorized be alert to patients and objects in the direction of movement so that no collision can occur.
--	--

2.4.1 Emergency stop

The Emergency stop is located on top of X-ray tube. In case of an Emergency situation, the activation of the push-button will turns off the Trauma Arm's motorized movements. Once the emergency situation has been controlled, the pushbutton can be reset by twisting it in a clock-wise direction to reset the controller system and return to normal operations.

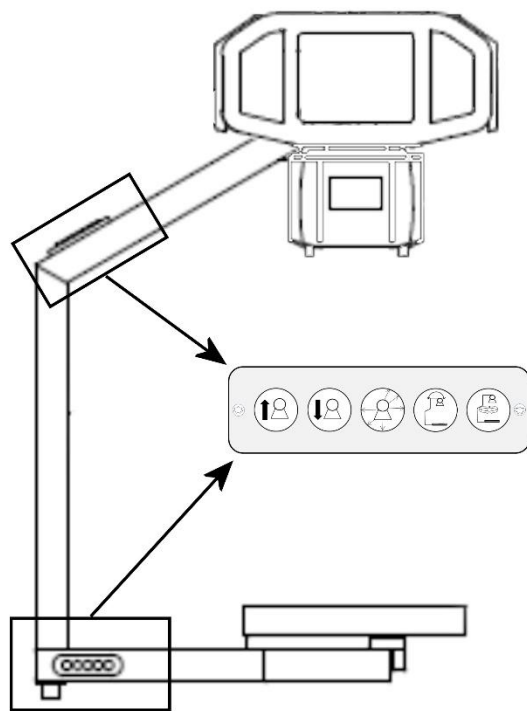
 WARNING:	This emergency stop control will not shut off the power to the interfaced equipment. Use the room emergency stop control for emergency situations.
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2.5 X-ray interlock

The Canon detector and X-ray generator are equipped with X-ray interlocks preventing the user from accidently making exposures. Please refer to the appropriate service manuals for a detailed description.

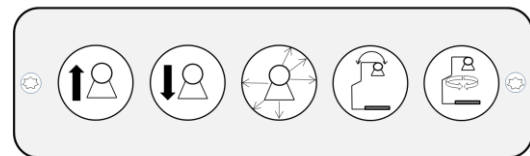
3 Operating controls

The Trauma Arm can be moved in all directions by hand force except for up and down movement. In order to prevent any possible collision, maintain a free area around the units. The manuals, should be studied to learn about the working zones and the limits of the device and to understand the risks of collisions when the unit is used with other devices and risks when used within the environment in which it is located.



The trauma arm contains two button groups which are conveniently positioned for AP position and for Lateral position.

Both button groups are identical but the order of buttons can be different as described here.



Movement buttons

Figure 2: Trauma arm with location of the movement buttons

Symbol	Description
	Motorized up
	Motorized down
	X and Y brake release
	AP-Lateral rotation
	Z-axis rotation



It is the operator's responsibility to observe movements carefully with the trauma arm. The system has a collision detection when motorized down is being used, but all mechanical movements and motorized up do not have collision detection.

It is not allowed to use the trauma arm as a tilting device!

3.1 X-ray tube display

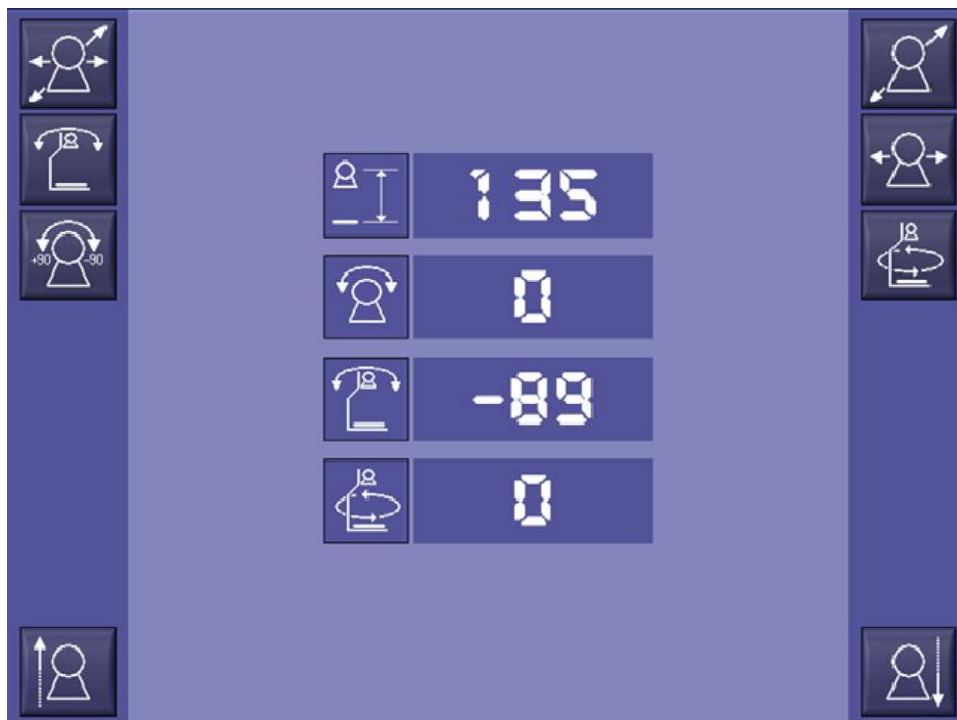


Figure 3: Tube Display

Touch handles release break mode		
Symbol	Description	
	X and Y movement disabled / enabled	
	AP-Lateral horizontal rotation disabled / enabled	
	Tube rotation disabled / enabled	
	Push – pull movement disabled / enabled	
	Left- right movement disabled / enabled	
	C-arm Vertical axis rotation disabled / enabled	
	Motorized up / down	
		Information: SID
		Information: Alpha angle
		Information: Horizontal angle
		Information: Vertical angle

3.2 Tube handles

The left and right side of the tube display contains handles to move the tube manually. Each tube handle contains three touch sensors at the back of the handle. If a person grabs one or both handles the sensors are being triggered, and the brakes, as set with the touch handle release break mode, will be released.

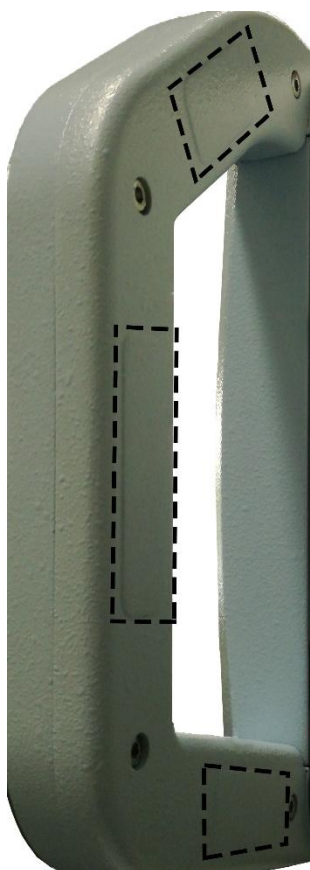


Figure 4: *Sensors at the back of the tube handle*

3.3 Detector case with anti-scatter grid

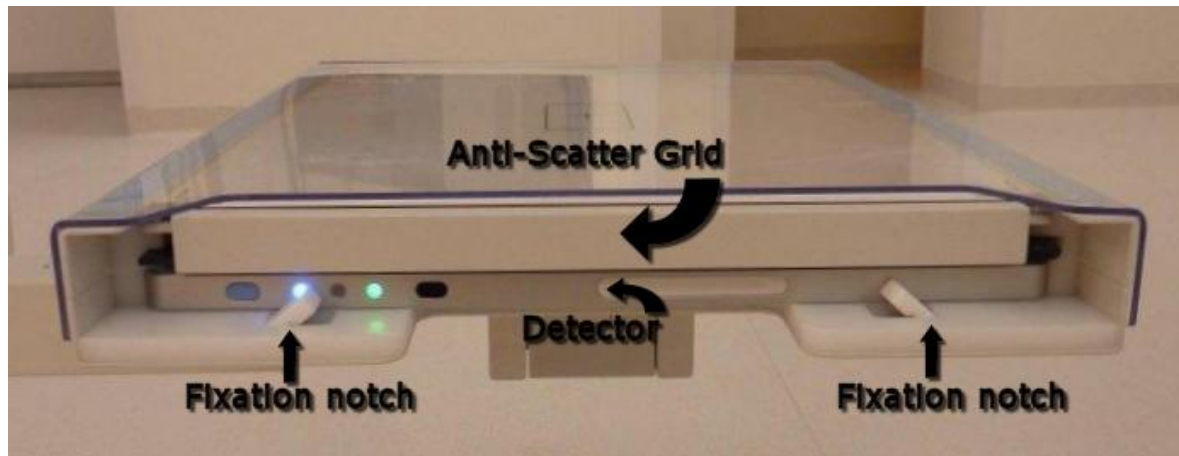


Figure 5: Detector Case with anti-scatter grid and detector

The detector case holds the detector and the anti-scatter grid. Pressing the grid bar will release the grid; press again to lock the grid.

The detector is being fixed by fixation notches to prevent the detector falling out of the detector case when rotated. Press both notches to release the detector. When inserting the detector shift the detector beyond the notches till they lock the detector.

It's advisable to insert the detector as shown on the image with the signal lights facing outside. This way the operator can check the status of the detector and can switch on or off the detector while inserted. It's possible that protocols have to be optimized for orientation.

3.4 X-ray generator control panel

The description in this paragraph applies to the CPI MP200 generator with a special interface to the Trauma DR plus system. For more details, see the documentation of the X-ray generator.



Advice to consult the accompanying documentation:

- Operation manual HF generator.

The X-ray unit may be dangerous to the patient and operator unless safe exposure factors and operating instructions are observed.

The generator mini control panel contains various indicators and controls. The relevant controls are:

- Switching the X-ray system ON or OFF
- Exposure switch and indicator (audible). Starting X-ray radiation.

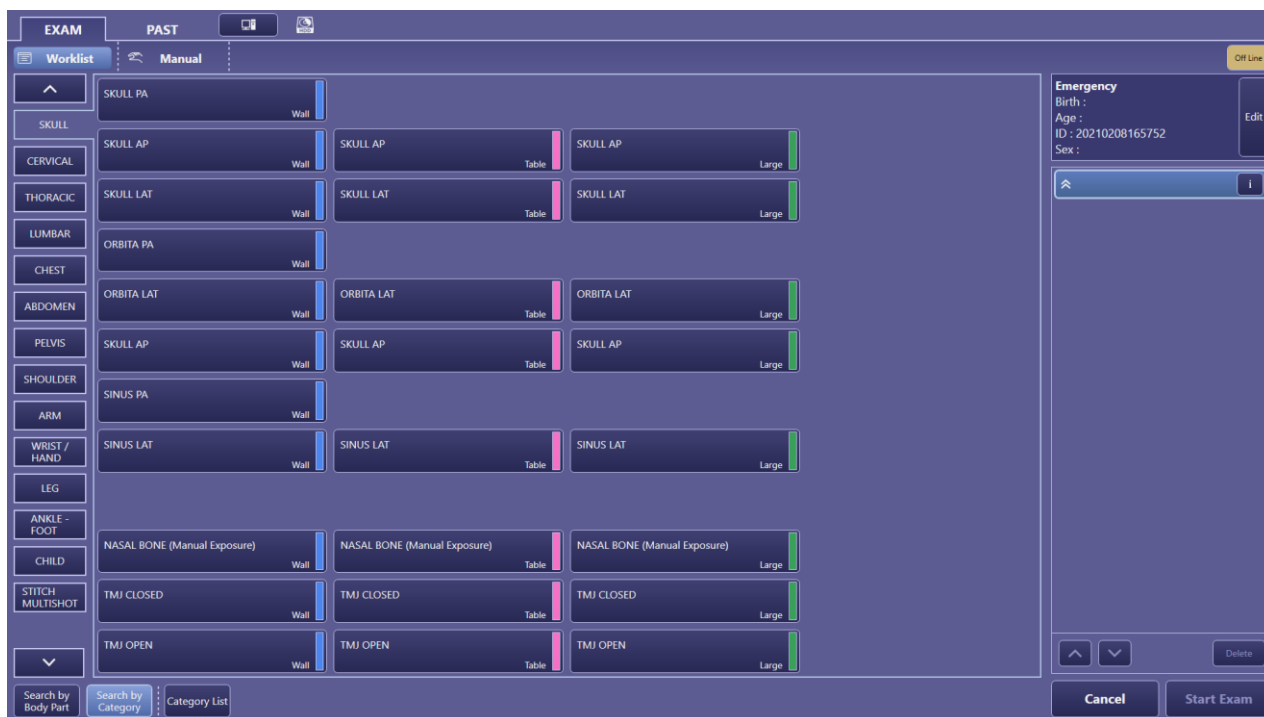
3.5 The user interface of control computer

Note:

Canon Inc. is the legal manufacturer of the CXDI Control Software used on the Trauma DR plus Acquisition/Control Computer.

The control computer is operated by;

- Touch screen buttons, or mouse



3.6 Wireless detector

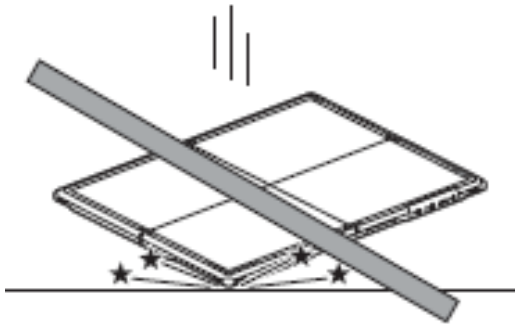
3.6.1 Handling

- Do not touch conductive parts of this equipment and the patient simultaneously. Otherwise, electric shock may result.
- Do not spill liquid or chemicals onto the equipment or, in cases where the patient is injured, allow it to come in contact with blood or other body fluids. Doing so may result in fire or electric shock. The detector is protected against dust (at the level of IP5X*) and liquids (at the level of IPX4**).

** Ingress of dust is not totally prevented, but dust shall not penetrate in a quantity to interfere with satisfactory operation of the apparatus or to impair safety (Dust protected). The equipment is protected so that specified operations and safety are not impaired even if the equipment is exposed to dust (75 µm in diameter) and dust gets inside the equipment and is left for eight hours.*

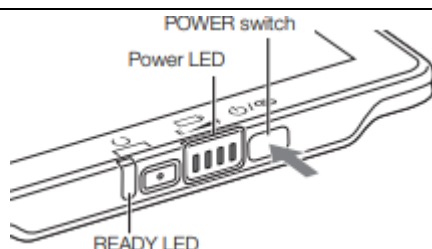
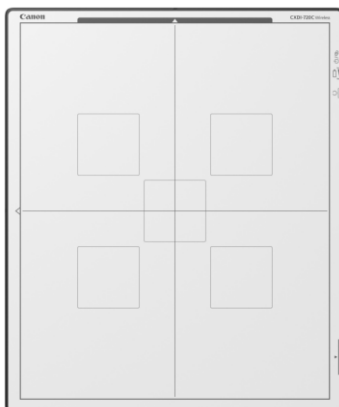
***Water splashing against the enclosure from any direction shall have no harmful effect (Protected against splashing water). There are no damaging effects when the equipment is sprayed with water from spray nozzles located in any direction at a pressure of 50 to 150 kPa by the 10 liter per minute over 5 minutes on the equipment.*

- Handle the equipment carefully as it is precision equipment. The inner device may be damaged if something hits against it, or if it is dropped, or, receives a strong jolt.



- Be sure to use the equipment on a flat surface while using it in horizontal position. If the detector is set on a diagonal and pressure is applied to it, the inner device may be damaged.
- Be sure to securely hold the equipment while using it in upright positions. Otherwise, the equipment may fall over, resulting in injury to the user or patient, or may flip over, resulting in damage to the inner device.
- Do not place excessive weight on the equipment. Do not use the equipment in a manner that will subject it to local loads of 100 kg or more. If the load exceeds the limit, the inner device may be damaged.
- Do not touch the electrode terminals of the equipment and the battery pack.

3.6.2 Detector

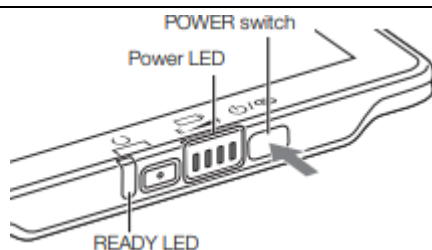


Press and hold the POWER switch (for at least 0,5 seconds) to turn the power on. The Power LED will light up.

Power LED					
Remaining battery charge	100% to 76%	75% to 51%	50% to 26%	25% to 5%	4% to 0%*

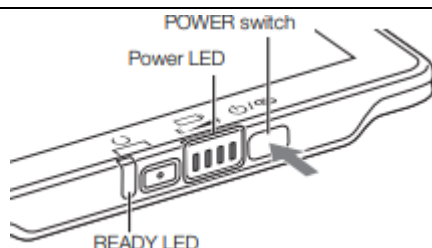
( : Lighted up,  : Flashing (1 sec. cycles),  : Off)

The Power LED to show the remaining battery power



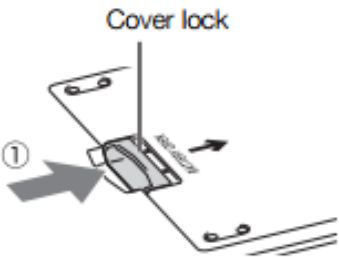
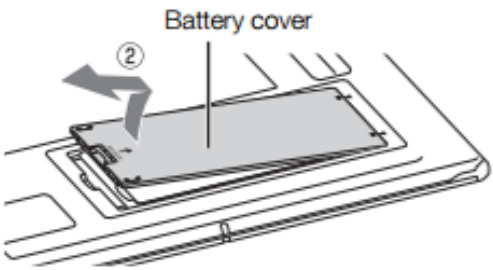
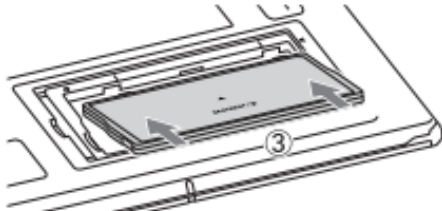
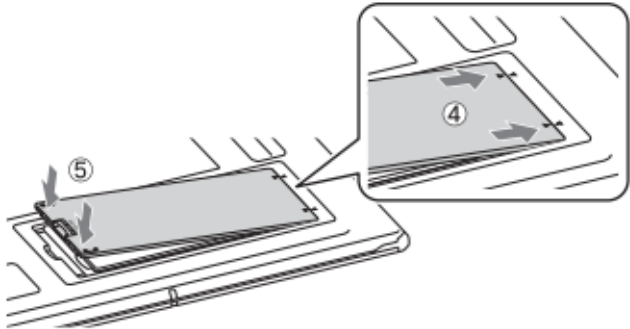
The ready LED to show whether the detector is ready to capture images

- Lights continually when detector status is "Ready"; exposure is possible.
- Flashes before exposure when the detector is in the "waiting" status.

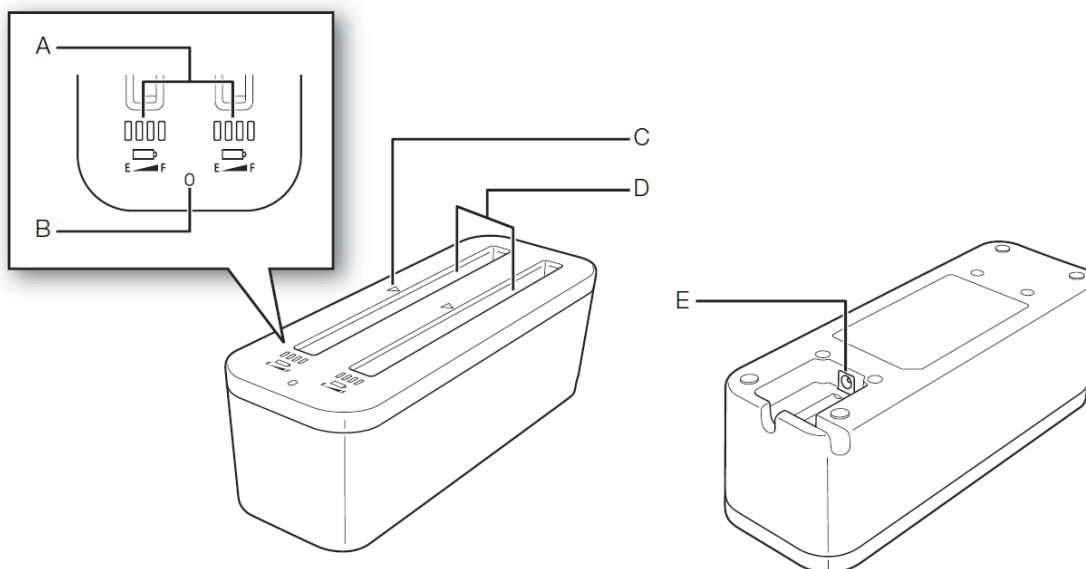


To switch off: press and hold the power switch till the Power LED extinguishes.

3.6.3 Battery pack

 Diagram 1 shows a close-up of the battery cover lock mechanism. A label 'Cover lock' points to the lock. A circled number '1' is next to an arrow indicating the direction to press and hold the lock.	1 - Press and hold the battery cover lock to release the lock.
 Diagram 2 shows the battery cover being lifted out of the device. A label 'Battery cover' points to the cover. A circled number '2' is next to an arrow indicating the direction to lift the cover.	2 - Lift up and pull out the Dust protected battery cover
 Diagram 3 shows the battery pack being inserted into the device. A circled number '3' is next to an arrow indicating the direction to insert the pack.	3 - Insert the battery pack fully
 Diagram 4 and 5 show the battery cover being inserted and locked. Diagram 4 is an inset showing the cover being inserted. Diagram 5 shows the cover being pressed down. A circled number '4' is next to an arrow indicating the direction to insert the cover. A circled number '5' is next to an arrow indicating the direction to press down on the cover.	
4 - Insert the battery cover fully 5 - Press down on the front of the battery cover to lock it.	

3.6.4 Battery pack charger

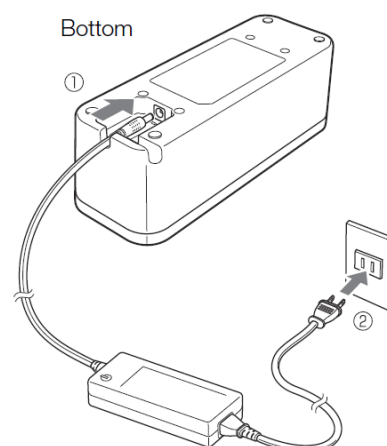


A	Charge status lamps	Shows the charge status of the battery packs inserted into each of the slots. For details, see “Charge status” (→ page 17).
B	POWER lamp	Lights up when the Battery Charger is turned on.
C	Direction marker	Serves as an index mark for inserting the batteries correctly to the slot.
D	Battery insertion slot	Accepts the Battery Pack LB-4A or LB-1A.
E	DC IN jack	Accepts the cable of the supplied AC adapter.

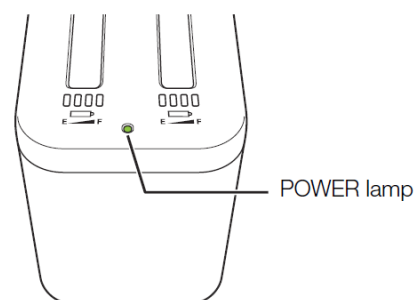
Note:	Fully charge the battery pack the first time it is used. The battery discharges gradually. The battery pack should be charged no more than two days before use.
--------------	--

1. Turn on the battery charger.

Insert the cable of the AC adapter into the DC IN jack (1) and the power cord plug into the AC outlet (2).



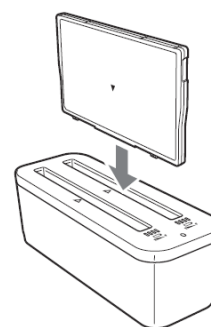
The POWER lamp lights green and the battery charger is ready for charging.



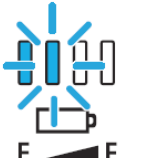
2. Charge the battery pack.

Insert the Battery Pack LB-4A or LB-1A as illustrated. Align the direction of the battery pack (▼) with the direction marker (▲) of the battery charger.

Insert the battery pack correctly and charging begins automatically.
The charge status lamps light blue when the battery is being charged.



Charge status

Charge status lamps					
Charging status	Charging 0% to 25%	Charging 26% to 50%	Charging 51% to 75%	Charging 76% to 99%	Charging finished 100%

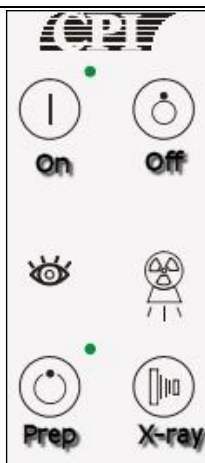
(■ : Lighted up  : Slowly flashing □ : Off)

Information:	Two battery packs can be charged at the same time.
Information:	It takes approximately 2.5 hours for the LB-4A and approximately 3.5 hours for LB-1A at a room temperature of 25°C to fully charge a completely discharged battery pack. The required charging time varies depending on the ambient temperature and the remaining battery level.

3. Remove the battery pack.

Remove the charged battery pack from the battery charger.

3.7 Generator mini console with external exposure button



Generator mini console

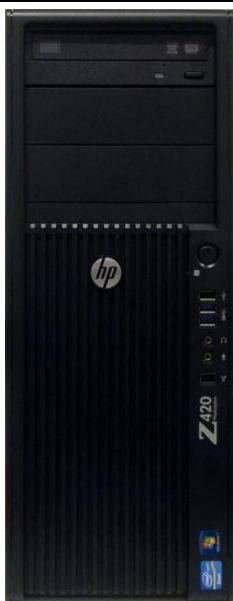
The generator mini console operates the X-ray generator.

The mini console contains an on and off button and a prep and exposure button.

Two indicators can light up during fluoroscopy or exposure.

All exposure settings, like kV, mAs or density are controlled within the acquisition software.

3.8 Acquisition Computer with CXDI NE software



Acquisition computer

The acquisition computer contains the CXDI software necessary for controlling the Trauma DR plus.

3.9 Touch Screen



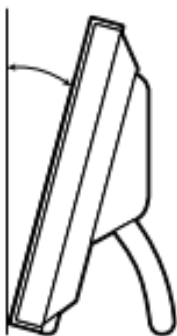
**This monitor is not for diagnostic purposes.
Use for diagnostics one Dicom Calibrated Diagnostic Workstation.**



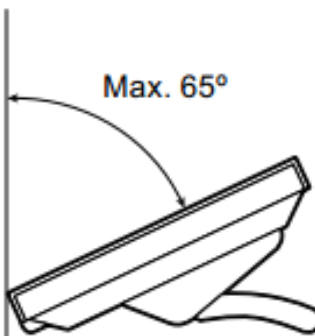
Touch Screen

The acquisition computer contains the CXDI software necessary for controlling the Trauma DR plus.

Min. 15°



Max. 65°



The touch screen can be tilted to angles of 15°, 25°, 35°, 45°, 55°, and 65°

4 Operating Instructions

Note:	When the system is on 24/7 it is recommended to restart the system once a day for cleaning up computer memory and automatically save the database. First turn on the x-ray generator and then the acquisition computer.
--------------	--

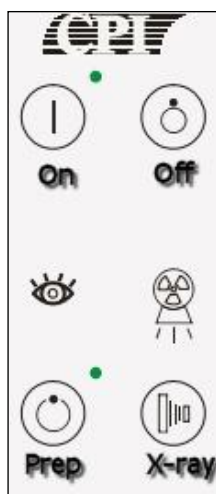
4.1 Powerup and Startup the system

Powerup the system

The Ceiling Stand does not have an on-board power switch. This equipment should normally be connected to the x-ray generator. The X-ray generator is connected to an independent circuit breaker switch located in the main electrical panel of the examination room. This switch should be placed into the RUN position to power the equipment ON.

Startup the system

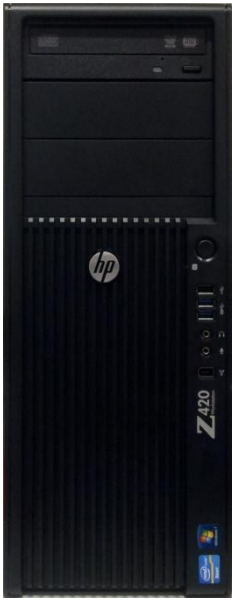
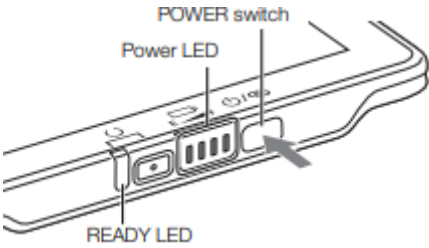
Turning on the x-ray generator



The generator mini console is a small box with on the upper side the on and off button and on the lower side the prep and exposure button.

In the middle there are indicators for fluoro and exposure when active the indicator will light.

Turn on the generator by pressing .

Turning on the Acquisition computer	Turning on the detector
	
Turn on the acquisition computer by pressing the power button.	Turn on the wireless detector by pressing and holding the Power switch. Release when power led lights up.

4.2 Tube warmup

To prolong the life of the X-ray tube it is important to warm-up the X-ray tube every day on the start of the day.

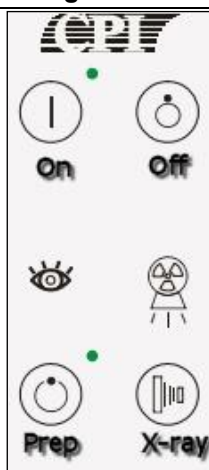
- Start an Emergency Patient.
- Select protocol: Tube warm-up in the Service: QA Category and click on Start Exam.
- Make 3 X-rays with the collimator fully closed with 30 seconds in between each X-ray.
- Click on Cancel. No images are captured; therefore, no images will be sent.

4.3 Shutdown and Power down the system

Shutdown the system

Note:	Before turning off the system park the system in its parking position. When the power is off the system cannot be moved.
--------------	---

Turning off the x-ray generator



Turn off the generator by pressing .

Turn off the acquisition computer by selecting  and . The computer will shut down automatically. The monitor will go standby.

Turn the detector off by pressing and holding the power button till the power led extinguishes.

Note:	Before switch off the main power, first bring down the equipment in the right sequence as described before.
--------------	--

Turn Off main circuit breaker

Switch off the main circuit breaker for the Trauma DR plus (depending on installation). This main circuit breaker is not a part of the Trauma DR plus and generally located on the power distribution panel from the hospital. The location of the power distribution panel is depending of the Installation.

4.4 CXDI Software

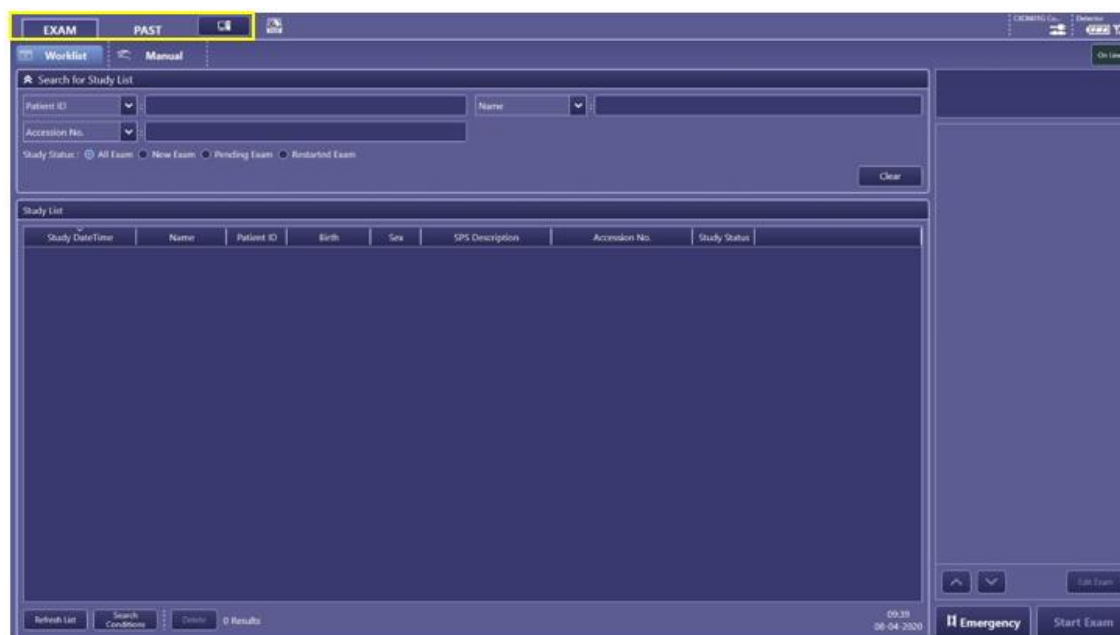
4.4.1 Logging in



It is possible to login to the CXDI software by entering the User Name and the Password and by clicking on Log In.

Note:	User and Log In setup wil be done in consultation with the User and wil be implemented during the application training.
--------------	--

4.4.2 Main screens

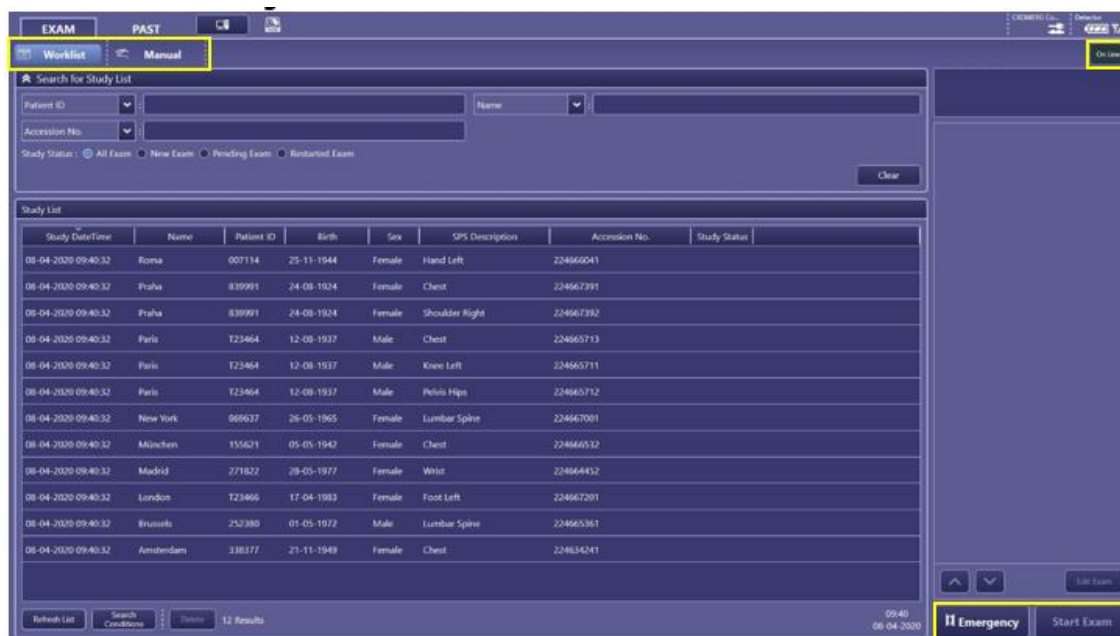


The user interface contains 3 main screens. Each main screen can be called from the upper left side of the screen.

EXAM	Worklist and manual input for image acquisition.
PAST	Patients who have been examined previously.
	To enhance easy reading this manual calls this screen the “system” screen. Here it is possible to adjust system settings, log out or shutdown the system.

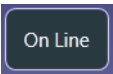
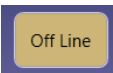
4.4.3 Exam screen

In the Exam screen it is possible to perform the examinations. There are three workflows for doing this:

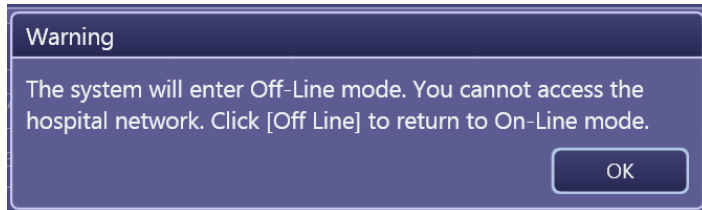


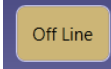
Study Date/Time	Name	Patient ID	Birth	Sex	SPS Description	Accession No.	Study Status
08-04-2020 09:40:32	Roma	007114	25-11-1944	Female	Hand Left	224665041	
08-04-2020 09:40:32	Praha	839991	24-08-1924	Female	Chest	224667391	
08-04-2020 09:40:32	Praha	839991	24-08-1924	Female	Shoulder Right	224667392	
08-04-2020 09:40:32	Paris	723464	12-08-1937	Male	Chest	224665713	
08-04-2020 09:40:32	Paris	723464	12-08-1937	Male	Knee Left	224665711	
08-04-2020 09:40:32	Paris	723464	12-08-1937	Male	Pelvis Hips	224665712	
08-04-2020 09:40:32	New York	869637	26-05-1965	Female	Lumbar Spine	224667091	
08-04-2020 09:40:32	München	155621	05-05-1942	Female	Chest	224665532	
08-04-2020 09:40:32	Madrid	271822	29-05-1977	Female	Wrist	224664432	
08-04-2020 09:40:32	London	723466	17-04-1983	Female	Foot Left	224667201	
08-04-2020 09:40:32	Brussels	252380	01-05-1972	Male	Lumbar Spine	224665361	
08-04-2020 09:40:32	Amsterdam	338377	21-11-1949	Female	Chest	224634241	

- By working with a worklist  **Worklist**
- By working with manual patient input  **Manual**
- By doing an Emergency examination  **Emergency**

Note:	The system can work in Online and Offline mode. During Offline mode the acquisition computer does not refresh the worklist and acquired images will not be send to a destination, PACS or DICOM printer.
	The system is Online and connected to the hospital information system and destinations. The worklist will be retrieved and when an exam has been ended, the acquired images are being sent automatically when a destination has been set.
	The system is Offline, not connected to hospital information system and destinations. The worklist will show the last patient list at the time the system is set to offline. Acquired images will stay in queue to be sent to their destinations when the system has been set to Online.

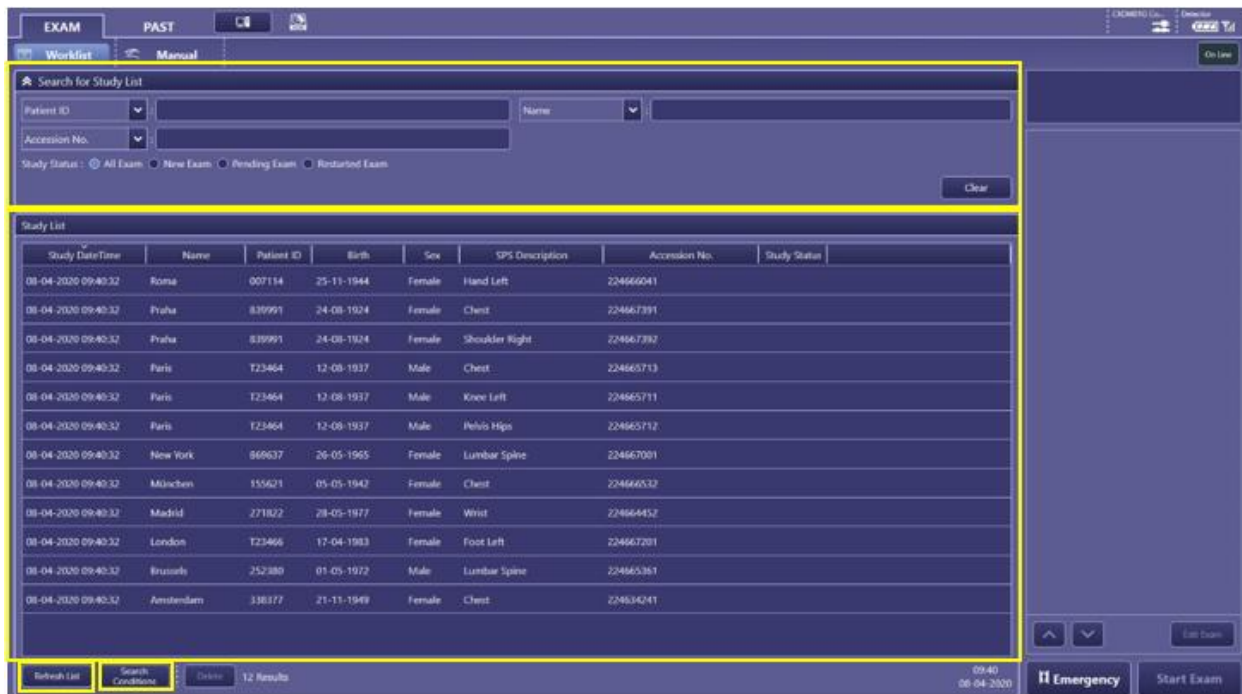
The system will show the following warning when entering offline mode:



It is possible to enter the On-Line mode again by clicking on  .

4.4.4 Worklist

In the worklist screen all patients who match the search conditions are retrieved from the worklist server and will be shown in the Study List.

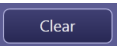


In the "Search for Study List" it is possible to filter the worklist by:

- Patient ID
- Name
- SPS Description
- Accession No.

It is also possible to filter by “Study Status”:

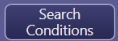
☒ All Exam	All Exams are shown
☐ New Exam	Only current worklist patients are shown
☐ Pending Exam	Only pending exams are shown
☐ Restarted Exam	Only restarted exams are shown

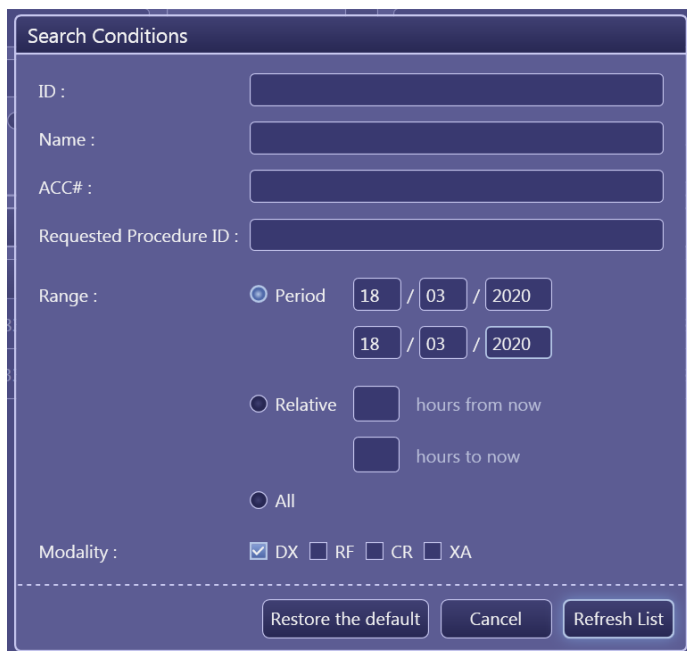
By clicking on  all filters will be cleared, by clicking on  it is possible to fold in the Search for Study List and by clicking on  it is possible to fold out the Search for Study List.

The different information columns (Study Date Time, Name, Patient ID etc.) in the Study List are adjustable in order, width and sorting.

- To change the order of a column: select the column name and drag the column to the desired place while keeping the left mouse button pressed
- To change the column width: select the column separator, when the mouse pointer changes in  press the left mouse button to drag and change the width.
- To change the information column on which the study list is sorting: click on the desired column. By clicking a second time on the column it is possible to change the sorting order, for instance from A to Z or from Z to A.

It is possible to refresh the Study List by clicking on  in the bottom left corner. This is only possible in Online mode; if the Offline mode is active this button will be greyed out.

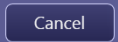
By clicking on  it is possible to change the following search conditions:



The 'Search Conditions' dialog box contains the following fields and options:

- ID :
- Name :
- ACC# :
- Requested Procedure ID :
- Range :
 - ☒ Period / /
 / /
 - ☐ Relative hours from now
 hours to now
 - ☐ All
- Modality : ☒ DX ☐ RF ☐ CR ☐ XA

At the bottom are three buttons: 'Restore the default', 'Cancel', and 'Refresh List'.

- By clicking on  the default settings will be restored. This will happen automatically by logging out.
- By clicking on  all changes will be cancelled.
- By clicking on  the net search conditions will be used to refresh the study list.

4.4.5 Worklist workflow

After selecting a patient from the worklist, start the examination by clicking on . When the patient has multiple studies planned, the software will select them all.

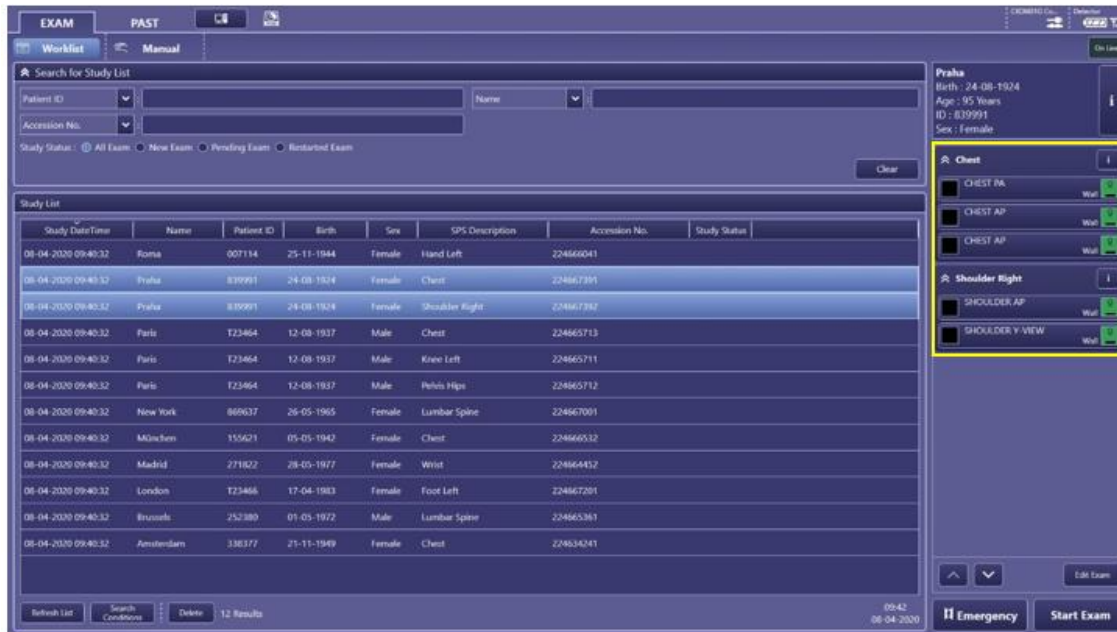
If the patient is not present in the list press refresh. Also, if not all studies aren't visible yet click refresh till all exams are present in the worklist.

When the exam is started by selecting a patient from the worklist there are two options:

- 1 - The system is programmed with prepacked protocols so studies will be filled automatically with protocols. This means the system will automatically go the exam after clicking on 
- 2 - No prepacked protocols have been added. This means you have to manually add protocols to a study after clicking on 

4.4.5.1 Prepacked protocols workflow

When prepacked protocols have been programmed the following screen will show after selecting the patient:



The screenshot shows the 'EXAM' screen with the 'Worklist' tab selected. The 'Study List' table contains the following data:

Study DateTime	Name	Patient ID	Birth	Sex	SPS Description	Accession No.	Study Status
08-04-2020 09:40:32	Roma	007114	25-11-1944	Female	Hand Left	224660041	
08-04-2020 09:40:32	Praha	839991	24-08-1924	Female	Chest	224667391	
08-04-2020 09:40:32	Praha	839991	24-08-1924	Female	Shoulder Right	224667392	
08-04-2020 09:40:32	Paris	T23464	12-08-1937	Male	Chest	224665713	
08-04-2020 09:40:32	Paris	T23464	12-08-1937	Male	Knee Left	224665711	
08-04-2020 09:40:32	Paris	T23464	12-08-1937	Male	Pelvis Hips	224665712	
08-04-2020 09:40:32	New York	869637	26-05-1965	Female	Lumbar Spine	224667001	
08-04-2020 09:40:32	München	155621	05-05-1942	Female	Chest	224666512	
08-04-2020 09:40:32	Madrid	271802	28-05-1977	Female	Wrist	224666452	
08-04-2020 09:40:32	London	T23466	17-04-1983	Female	Foot Left	224667201	
08-04-2020 09:40:32	Brussels	252389	01-05-1922	Male	Lumbar Spine	224665361	
08-04-2020 09:40:32	Amsterdam	338377	21-11-1949	Female	Chest	224634241	

The right panel shows the 'Chest' and 'Shoulder Right' protocols. The 'Start Exam' button is highlighted.

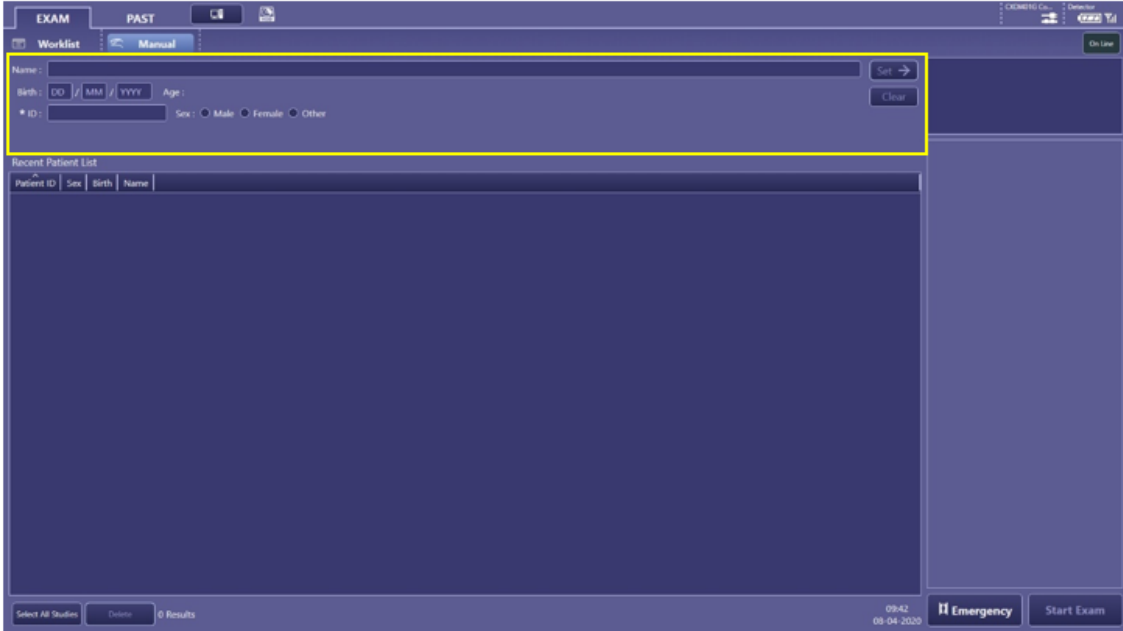
- When the prepacked protocols are sufficient for this patient you can start the exam by clicking on **Start Exam**
- When the prepacked protocols aren't sufficient for this patient you can manually add protocols by clicking on **Edit Exam**.

4.4.5.2 Manually adding protocols workflow

When no prepacked protocols have been programmed you will have to manually add protocols to the study. In chapter "Manually adding protocols" is described how you can do this.

4.4.6 Manual

In the Manual screen you can manually add patients by entering patient data. You can add multiple studies to the patient.



You can insert the patient data displayed in the top left part (Name, Birth etc.). The patient data displayed with an asterisk (*) are mandatory.

- By clicking on  you can set the entered patient data to start the examination.
- By clicking on  you can clear the entered patient data.

All manually added patients will show in the “Recent Patient List”.

Note:	In the screen Exam → Manual all patients are being displayed which have been selected from the worklist or entered manual in the past. It is possible to select a patient from this list but keep in mind that only patient information will be used. Other information, like accession number will not be present.
--------------	--

4.4.7 Manual adding patient workflow

After entering the correct patient data, the following screen will show:



In chapter “Manually adding protocols” is described how you can add protocols.

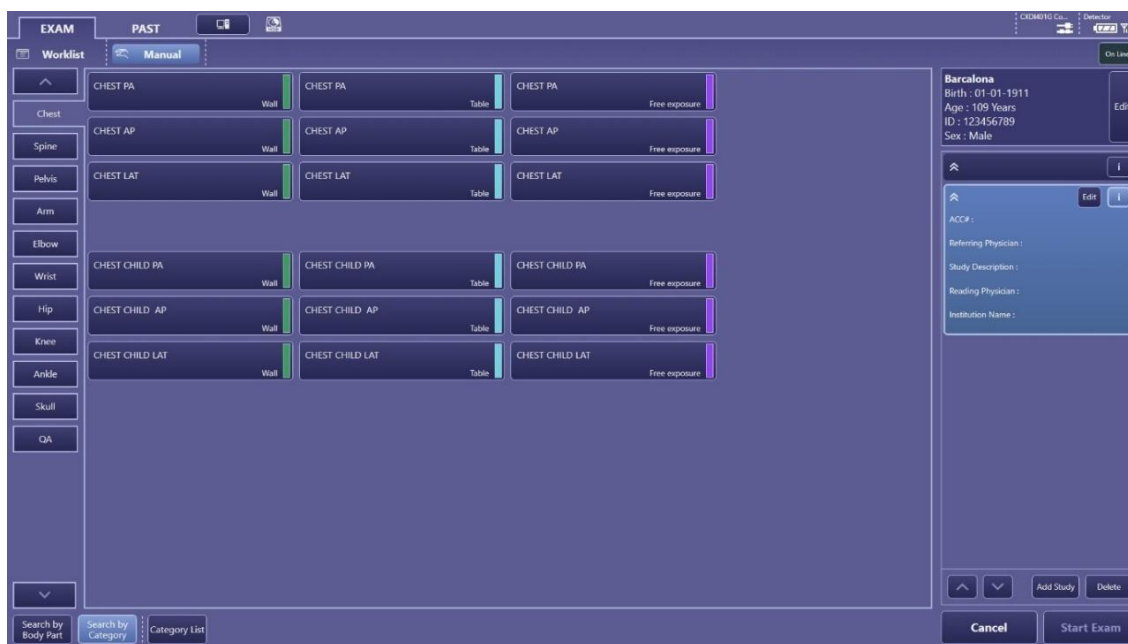
4.4.8 Adding multiple studies

You can add another study by clicking on **Add Study**. You can delete a study by clicking on **Delete**.

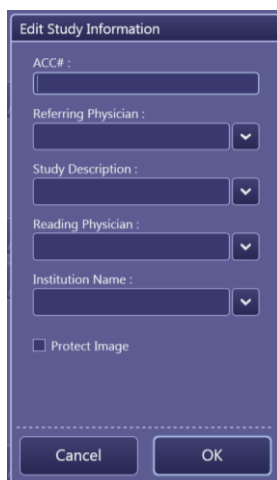


4.4.9 Editing study data

You can edit study data by clicking on . After this the following screen will show:



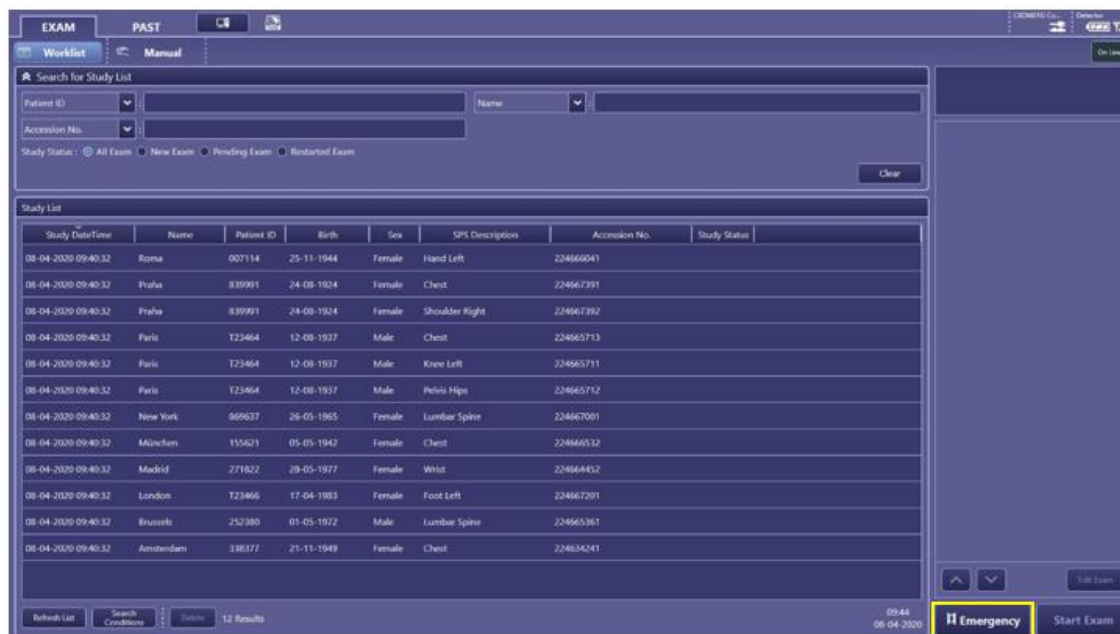
- By clicking on  the following screen will show:



- By clicking on , the study data will be added. Different data can be added to different studies.

4.4.10 Emergency

In both the Worklist screen and the Manual screen, you can perform an emergency examination when there is no patient data available. You can only add one study to a patient.



The screenshot shows the 'Worklist' screen with a search bar and a table of studies. The 'Emergency' button is highlighted in yellow at the bottom right.

Study DateTime	Name	Patient ID	Birth	Sex	SPS Description	Accession No.	Study Status
08-04-2020 09:40:32	Roma	007114	25-11-1944	Female	Hand Left	224665041	
08-04-2020 09:40:32	Praha	839991	24-08-1924	Female	Chest	224667391	
08-04-2020 09:40:32	Praha	839991	24-08-1924	Female	Shoulder Right	224667392	
08-04-2020 09:40:32	Paris	723464	12-08-1937	Male	Chest	224665713	
08-04-2020 09:40:32	Paris	723464	12-08-1937	Male	Knee Left	224665711	
08-04-2020 09:40:32	Paris	723464	12-08-1937	Male	Pelvis Hips	224665712	
08-04-2020 09:40:32	New York	869037	26-05-1965	Female	Lumbar Spine	224667091	
08-04-2020 09:40:32	München	155621	05-05-1942	Female	Chest	224666532	
08-04-2020 09:40:32	Madrid	271822	28-05-1977	Female	Wrist	224664432	
08-04-2020 09:40:32	London	723466	17-04-1983	Female	Foot Left	224667201	
08-04-2020 09:40:32	Brussels	252380	01-05-1972	Male	Lumbar Spine	224665361	
08-04-2020 09:40:32	Amsterdam	338377	21-11-1949	Female	Chest	224634241	

At the bottom right, the 'Emergency' button is highlighted in yellow, and the 'Start Exam' button is visible next to it.

Note:

- 1 - It is not possible to divide the images into several studies.
- 2 - It is not possible to send images with **Send**.

4.4.11 Emergency patient workflow

After clicking on **Emergency** the following screen will show:



- By clicking on **Edit** you can edit data. The following screen will show:



Here you can enter for example "room 1" in the Name field to differentiate between multiple emergency patients. After this click on **OK**.

In chapter "Manually adding protocols" is described how you can add protocols.

4.4.12 Manually adding protocols

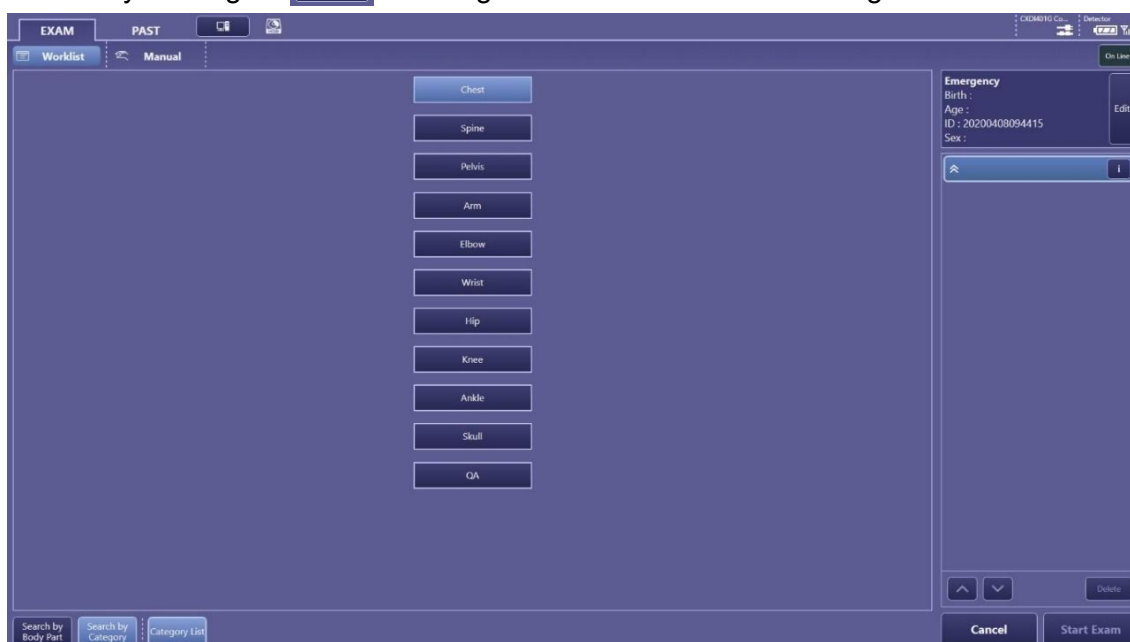
Protocols can be displayed in Categories and in Body Parts.

4.4.12.1 Categories

Different categories are shown. In this case for example: Chest, Spine etc. You can scroll through different tabs by clicking on  or .



- By clicking on  all categories will show in the following screen:

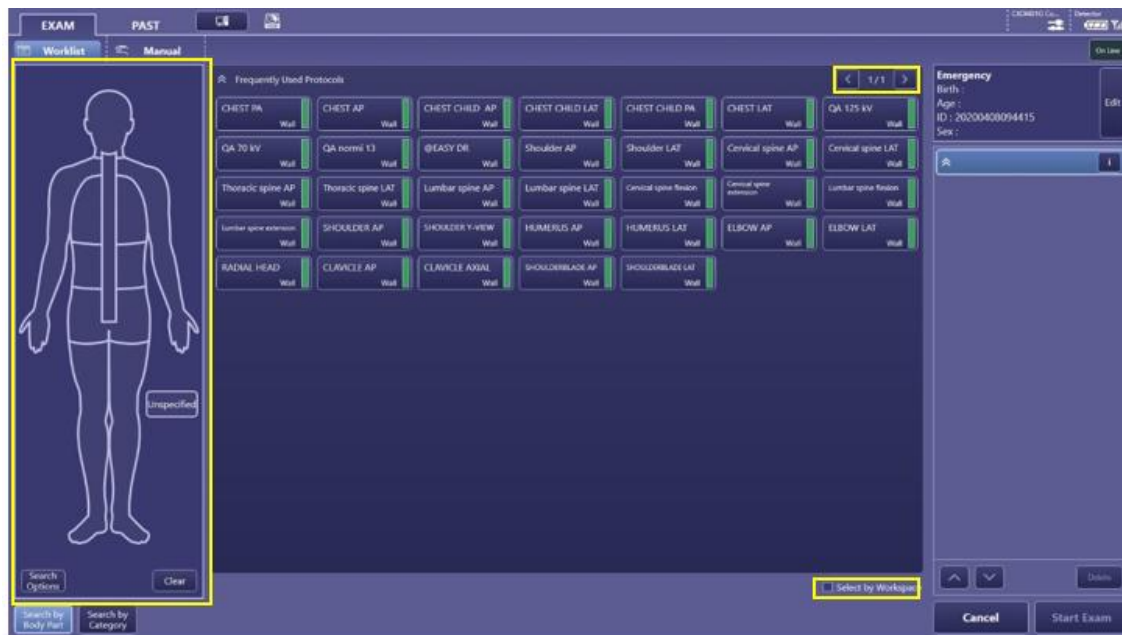


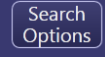
- By clicking on one of the categories this category will open, showing all protocols belonging to this category.

4.4.13 Body Parts

The system has been preloaded with clinical protocols. They have been optimized for optimal image quality and dose. They can be applied directly, but may require adjustments according to local regulations.

It is also possible to search by body part. Here you can find a protocol by using the anatomical diagram showing on the left. Selecting an anatomical region will show all protocols belonging to this region, showing the most frequently or most recent protocols first (depending on the system setup). You can scroll through the different tabs by using



- By clicking on  you can search by Exposure Type (radiography, stitch), View Position (AP, PA, etc.) or by Protocol Name.
- By clicking on  all Search Options and the selected anatomical regions will be cleared.

If ☐ **Select by Workspace** is not selected all different protocols combined with the default workspace (table, wall, free exposure) will show.

If ☒ **Select by Workspace** is selected all different protocols will show but you will have to select the workspace (table, wall, free exposure) after selecting the protocol.

4.4.14 Workspaces

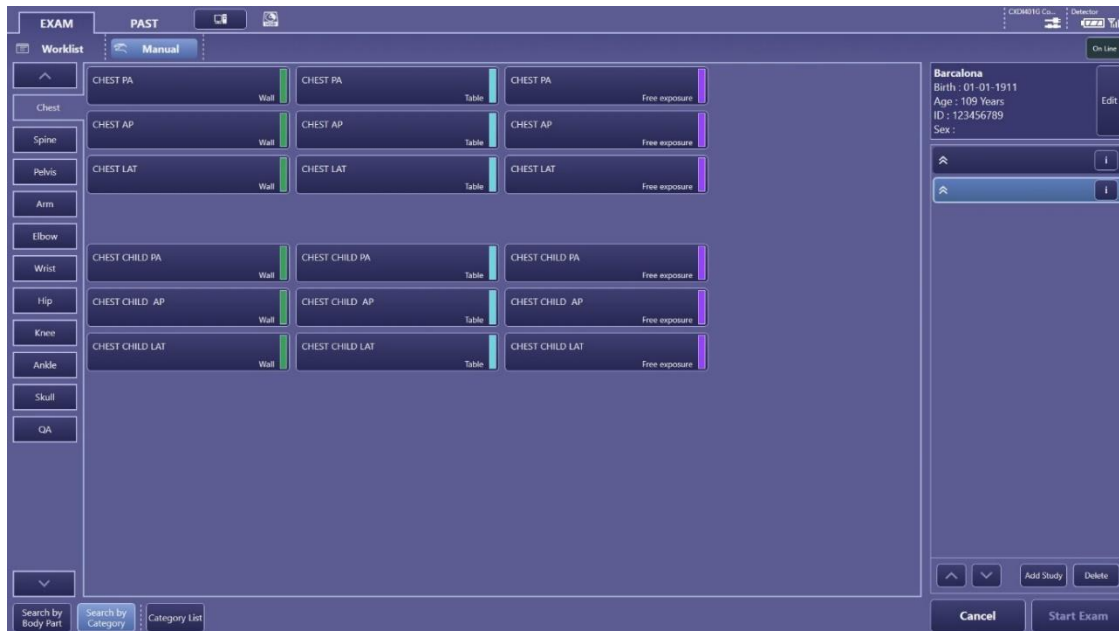
There are different workspaces possible. Every workspace corresponds with a color. In the below case CHEST PA has 3 different workspaces:

	Thorax AP on the wall stand
	Thorax AP on the table
	Thorax AP with free exposure

4.4.15 Adding the protocols

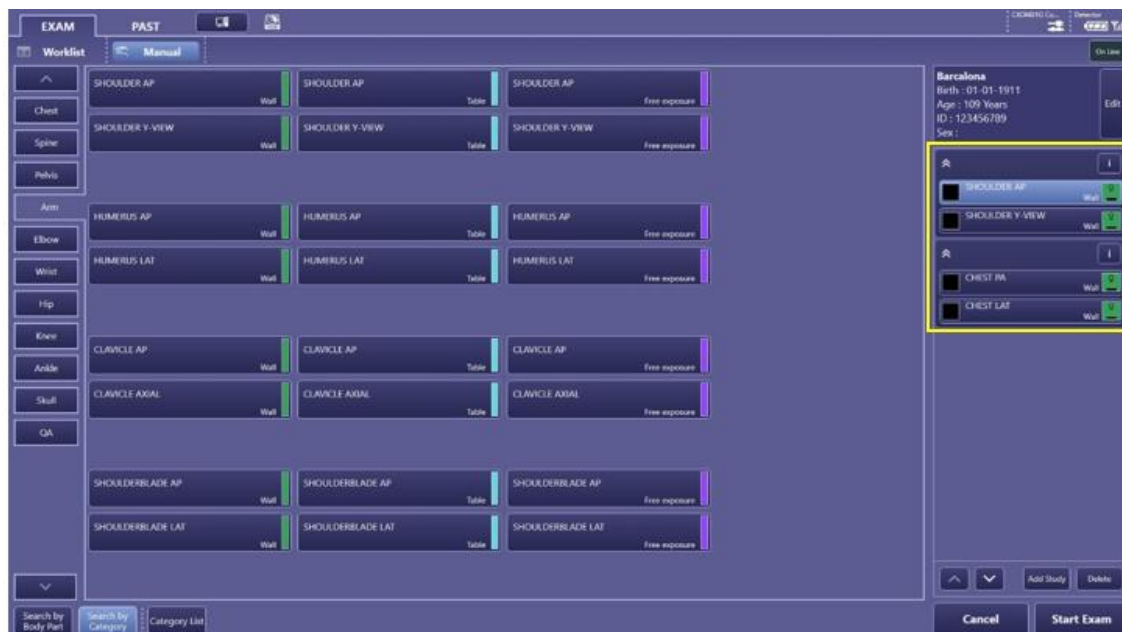
Note:

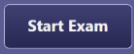
In this example we will use a manual added patient adding protocols for 2 studies using categories.



You can add a protocol to a study by first selecting the study and then by clicking on the desired protocol. You can add multiple protocols to one study. In the below case 2 protocols are added to 2 studies. You can change the order of the protocols by selecting the protocol you want to move (the light blue highlighted protocol is the selected protocol)

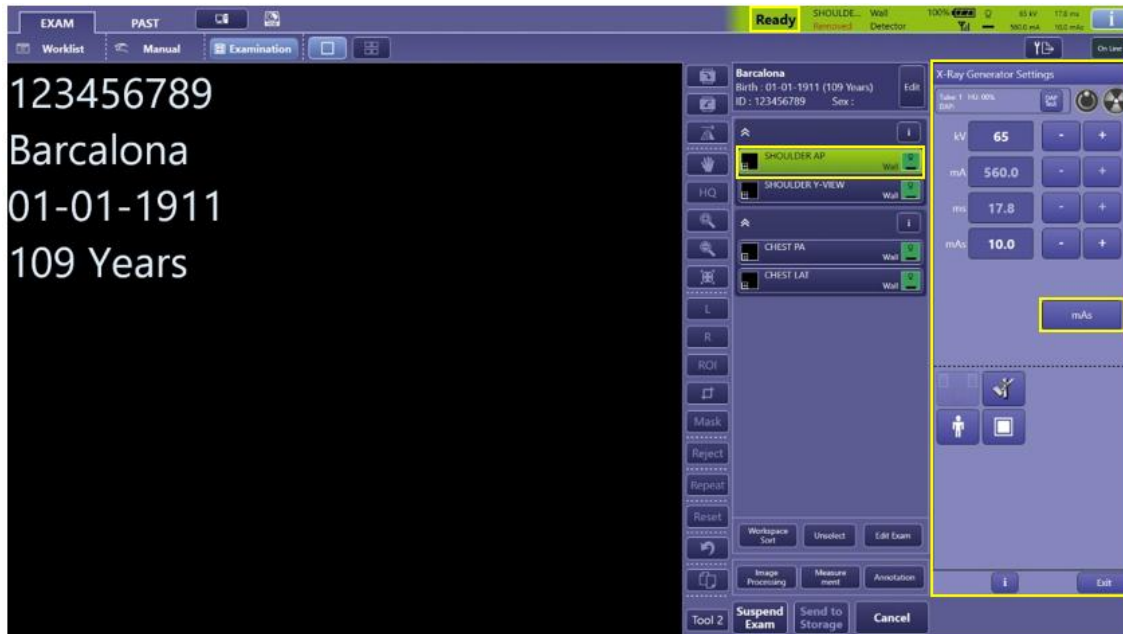
and by clicking on  . By clicking on  you can delete a protocol.



When all desired protocols are added, start the exam by clicking on .

4.4.16 Making x-rays

After starting the exam, the following screen will show. The green highlighted protocol is the selected protocol. You can select another protocol by clicking on it.



In the upper right part of the screen it is possible to see if the system is ready to make an x-ray

Ready	The system is ready to make the x-ray
Not Ready	The system is not ready to make the x-ray
Waiting...	The system is getting ready to make the x-ray

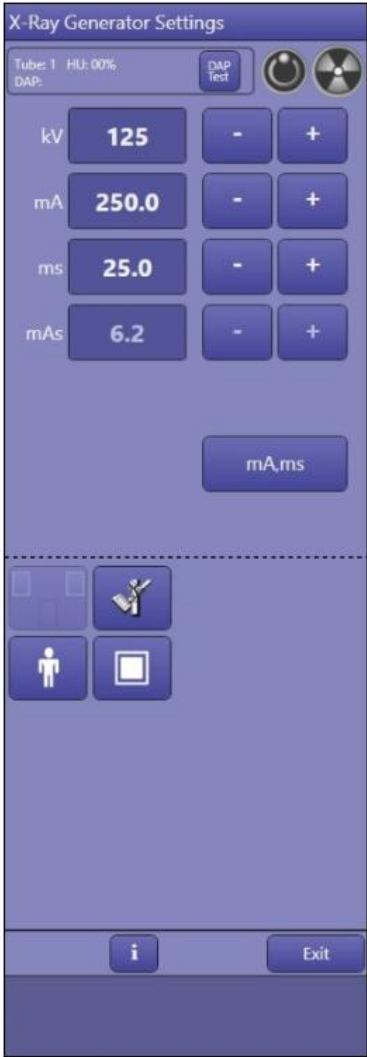
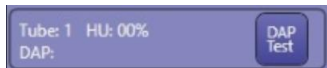
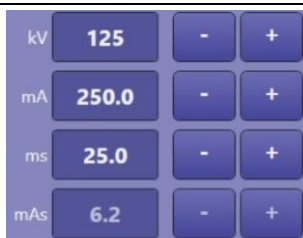
You can add a protocol by clicking on .

It is possible to change the workspace of a protocol by clicking on the selected protocol a second time and then selecting the desired workspace.

CAUTION:	Adding materials or objects into the X-ray beam will affect the dose and image quality.
-----------------	--

Generator Window

In the right part of the screen the x-ray generator screen is shown. Here it is possible to change the x-ray parameters (kV, mA, ms, mAs, exposure type, density, AEC fields, body size, focus size).

		Information window for: - Tube heat: (HU ...%) - DAP:
		Performs a DAP meter test and resets the DAP meter
		System is ready for exposure
		Exposure / irradiation indication
		X-ray parameters adjustment fields to optimize exposure parameters.
		Select exposure type: - 2-point (kV and mAs) exposure - Automatic Exposure Control (AEC) exposure - 3-point (kV, mA, ms) exposure.
		Select receptor type (C-arm or wireless)
		Select active AEC field. In de Trauma arm only the middle field is active. With the AEC in the detector you can make a combination from 5 fields.
		Select Body type
		Select focus type

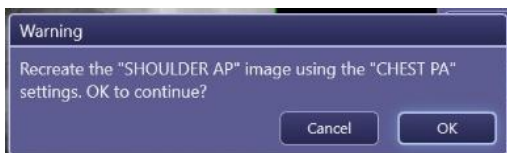
4.4.17 Moving an x-ray from one study to another study

When an x-ray is made in one study, but it belongs to another study it is possible to change it from one study to another. This is only possible from a workspace to the same workspace.

In the below case for example and x-ray of the thorax has been made with shoulder x-ray parameters and postprocessing. By clicking on the thumbnail of the x-ray that has been made you can drag the x-ray to the THORAX PA on the same workspace (wall).



After this, the following warning will show:



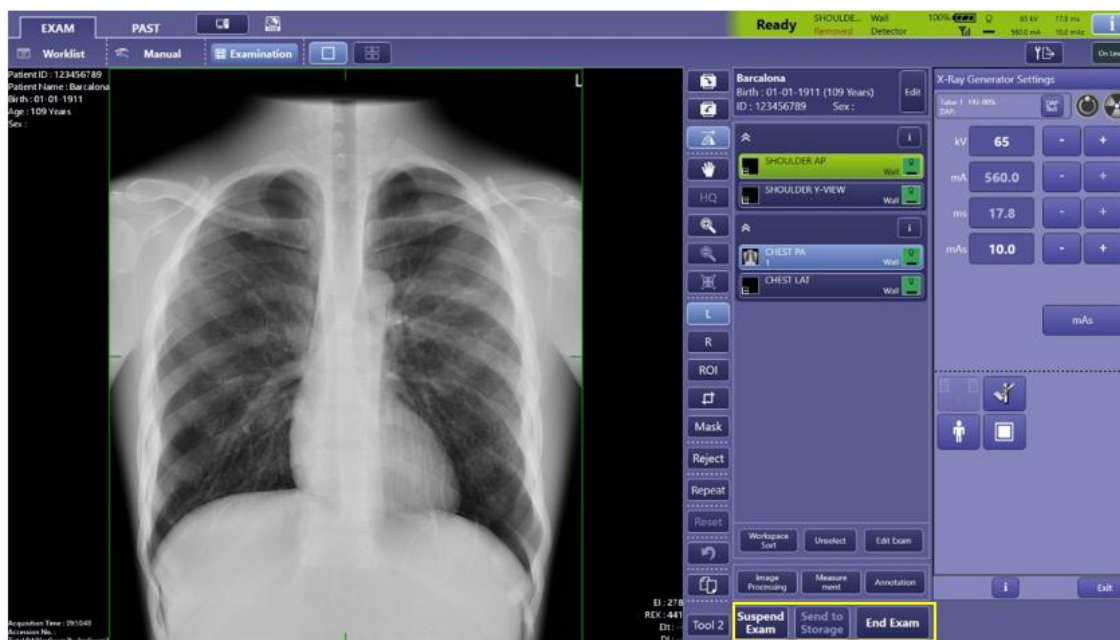
- By clicking on **OK**, the x-ray will move to the correct study and the x-ray will be processed with the correct image processing as shown in the image below.

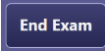
**Note:**

The x-ray has already been made with the x-ray parameters belonging to the wrong protocol. By moving it to the correct study, this can't be undone.

4.4.18 Ending the exam

After making the x-rays the following screen will show:



	Ending the exam temporarily. The made x-rays will not be send to the default destinations and the patient will sustain in the worklist with the study status "pending"
	Sending the x-rays that have been made till now to the default destinations. The exam will not end. This button is not available in an emergency examination.
	Ending the exam and automatically sending all acquired x-rays to the default destinations.

4.4.19 Dose Indication

REX Value

The Rex value given after every exposure is an indication that the dose received by the detector is sufficient to create a good image. This value needs to be between 100 and 400.

When this value is too low, the dose is too low, resulting in a grainy image. When the value is high, the dose is too high resulting in an unnecessarily high dose to the patient.

Note:	In digital imaging it is hard to see on the image itself when the dose is too high, because of dynamic range compression. That's why it is important to always check the REX value.
--------------	--

The REX value is shown in the lower right corner of the image:

DAP(mGycm2) : --
 REX : 148
 EI : 111

Check REX Value

Use the following procedure to check the REX value.

Use for all the exposures the following protocol and settings.

Category Service > protocol QA (table)

Post processing parameters that influences the REX Value must be always the following:

- Anatomical part
 - Category: Upper-Extremity
 - Anatomical Part: Humerus
 - Direction: Front
- LUT:
 - Curve Shape: LN (linear)
 - Brightness Adjustment: Auto
 - Base Brightness: 15
 - Base Contrast: 15

Make an exposure to test the automatic exposure control and measure the Screen dose.

Filter: 2mm Cu, measuring chamber in the middle above the middle AEC field.

Setting: 80 kV, large focus, density 0.

SID table: 120cm, SID wall: 200cm

Read the REX-value from the Canon monitor. REX (Reached EXposure index)

Note the mAs and REX-value of the left, middle and right field.

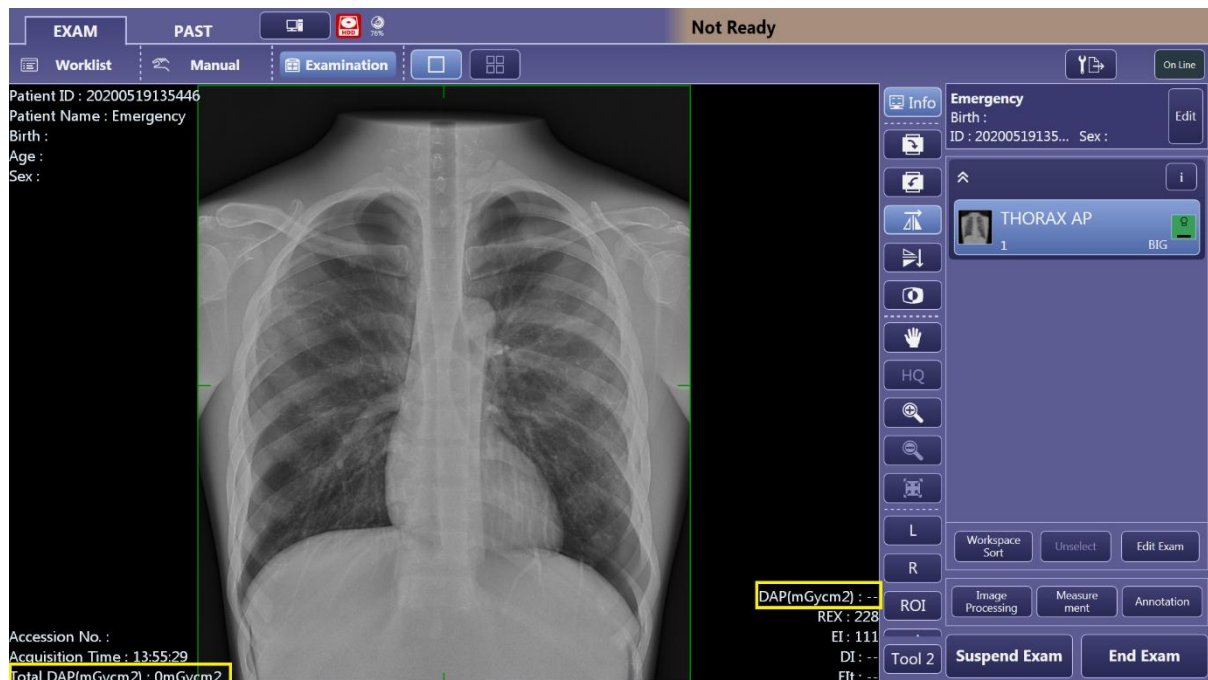
	SID	Left	Middle	Right	REX-value
Detector Table	1.20 m	mAs	mAs	mAs	
Detector Wall	2.00 m	mAs	mAs	mAs	

Compare the value with the reading made during installation or the last maintenance visit
 The reading should be $\leq 5\%$.

if the readings are outside the specs, Call your service representation

DAP meter

The systems contain an integrated Dose Area Product (DAP) measurement device. DAP values will be shown on the monitor of the system in mGycm² or in μGym^2 (depending on local settings).

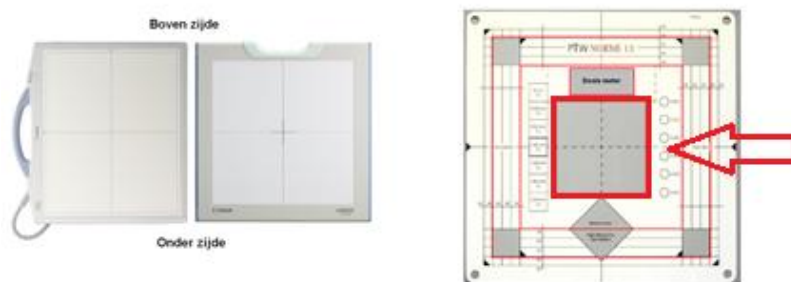


Check DAP meter

Setup the system with the following parameters

SID = 100cm

Generator: 100kV, 20mAs, large focus



Use the Normi 13 phantom to collimate the collimator on 10x10cm, collimated field 100cm²
 Take the x-ray and record the DAP reading

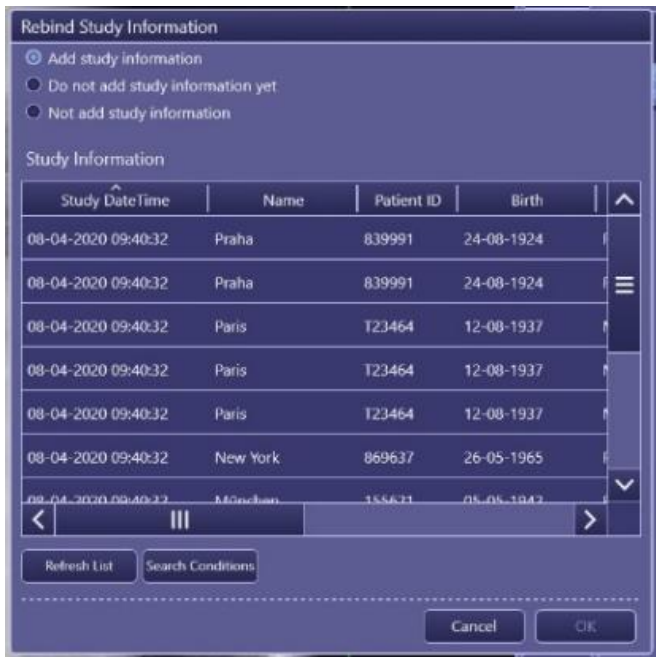
kV	50	70	80	100	125
mAs	10	10	10	10	10
DAP reading (mGy.cm ²)					
Calculated DAP (mGy)	0,0	0,0	0,0	0,0	0,0
Deviation	%	%	%	%	%
Calculation: Calculated DAP = DAP reading / 100					
Deviation = Calculated DAP – Measured DAP / Calculated DAP *100					

Compare the value with the reading made during installation or the last maintenance visit
 The value should be $\leq 25\%$.

If the readings are outside the specs, Call your service representation.

4.4.20 Ending an emergency exam

When patient data is available and inserted in RIS, click on  and the following screen will show:



The dialog box titled "Rebind Study Information" contains three radio buttons: "Add study information" (selected), "Do not add study information yet", and "Not add study information". Below is a table of study information.

Study DateTime	Name	Patient ID	Birth
08-04-2020 09:40:32	Praha	839991	24-08-1924
08-04-2020 09:40:32	Praha	839991	24-08-1924
08-04-2020 09:40:32	Paris	T23464	12-08-1937
08-04-2020 09:40:32	Paris	T23464	12-08-1937
08-04-2020 09:40:32	Paris	T23464	12-08-1937
08-04-2020 09:40:32	New York	869637	26-05-1965
08-04-2020 09:40:32	Moscow	155631	05-05-1942

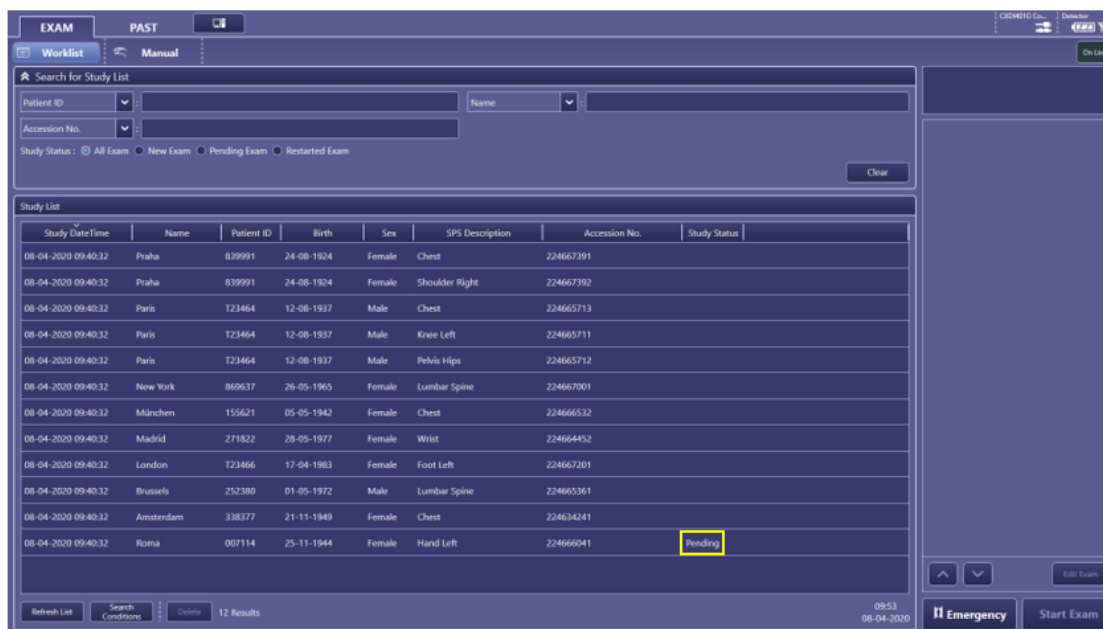
At the bottom are buttons for "Refresh List", "Search Conditions", "Cancel", and "OK".

Click on the correct patient (that has been entered in RIS) and click on . To verify, click on  again.

Note:	This action can't be undone.
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4.4.21 Pending patients

Patients who have been suspended with the **Suspend Exam** button will have the Study Status **Pending** in the worklist.



Study DateTime	Name	Patient ID	Birth	Sex	SPS Description	Accession No.	Study Status
08-04-2020 09:40:32	Praha	839991	24-08-1924	Female	Chest	224667391	
08-04-2020 09:40:32	Praha	839991	24-08-1924	Female	Shoulder Right	224667392	
08-04-2020 09:40:32	Paris	723464	12-08-1937	Male	Chest	224665713	
08-04-2020 09:40:32	Paris	723464	12-08-1937	Male	Knee Left	224665711	
08-04-2020 09:40:32	Paris	723464	12-08-1937	Male	Pelvis Hips	224665712	
08-04-2020 09:40:32	New York	869637	26-05-1965	Female	Lumbar Spine	224667001	
08-04-2020 09:40:32	München	155621	05-05-1942	Female	Chest	224666532	
08-04-2020 09:40:32	Madrid	271822	28-05-1977	Female	Wrist	224664452	
08-04-2020 09:40:32	London	723466	17-04-1983	Female	Foot Left	224667201	
08-04-2020 09:40:32	Brussels	252380	01-05-1972	Male	Lumbar Spine	224665361	
08-04-2020 09:40:32	Amsterdam	338377	21-11-1949	Female	Chest	224634241	
08-04-2020 09:40:32	Roma	007114	25-11-1944	Female	Hand Left	224666041	Pending

Pending patients will sustain in the worklist. To delete a pending patient, select the pending patient, start the exam and then click on **End Exam** when images are acquired or on **Cancel** if no images are acquired.

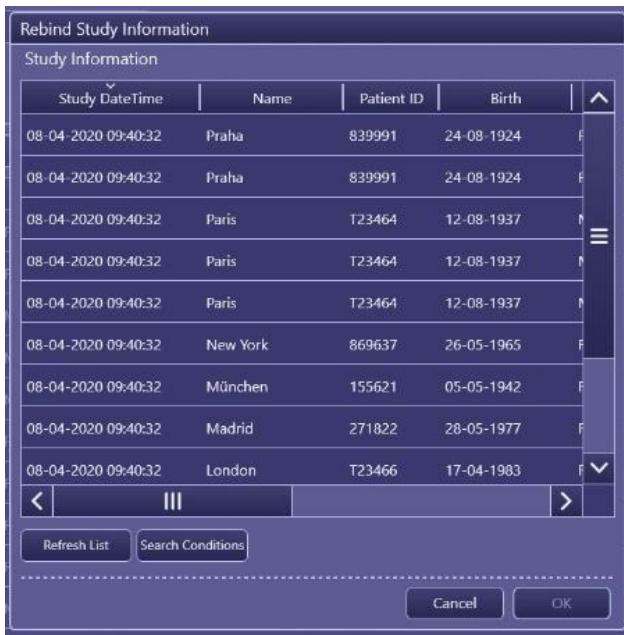
4.4.22 Correcting patient data after making images with the wrong patient data

Note:

This is only possible when the images have not been sent to PACS/has not been ended

After selecting the patient from the worklist and making images it is possible to correct the patient data. Click on **Suspend Exam** after making the images. The study will remain in the worklist.

Select the patient and click on  in the upper right part of the screen. After this the following screen will show:



Rebind Study Information

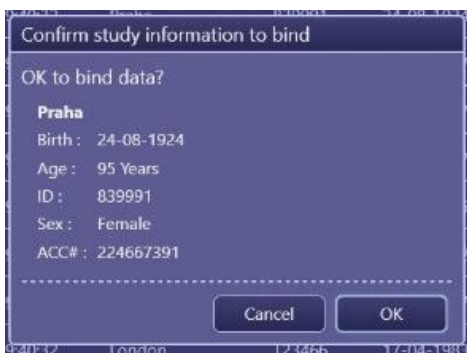
Study Information

Study DateTime	Name	Patient ID	Birth	
08-04-2020 09:40:32	Praha	839991	24-08-1924	F
08-04-2020 09:40:32	Praha	839991	24-08-1924	F
08-04-2020 09:40:32	Paris	T23464	12-08-1937	F
08-04-2020 09:40:32	Paris	T23464	12-08-1937	F
08-04-2020 09:40:32	Paris	T23464	12-08-1937	F
08-04-2020 09:40:32	New York	869637	26-05-1965	F
08-04-2020 09:40:32	München	155621	05-05-1942	F
08-04-2020 09:40:32	Madrid	271822	28-05-1977	F
08-04-2020 09:40:32	London	T23466	17-04-1983	F

Refresh List Search Conditions

Cancel OK

Select the correct patient and click on **OK**. After this the following screen will show:



Confirm study information to bind

OK to bind data?

Praha

Birth : 24-08-1924

Age : 95 Years

ID : 839991

Sex : Female

ACC# : 224667391

Cancel OK

Click on **OK** when the new patient data is correct. After this the study will appear in the worklist with the new correct patient data. To end the study, select the study and click on

Start Exam

End Exam

4.4.23 Post Processing

It is possible to perform different kind of postprocessing on the images. This chapter will explain them.

Note:	The quality of an image is to a large extent determined by the digital image processing. Keep in mind that the digital image processing is strongly dependent on sufficient exposure parameters.
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4.4.24 Toolbar

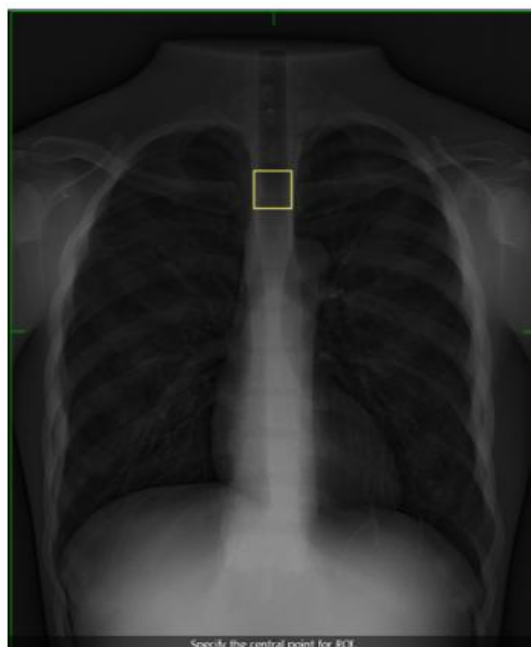
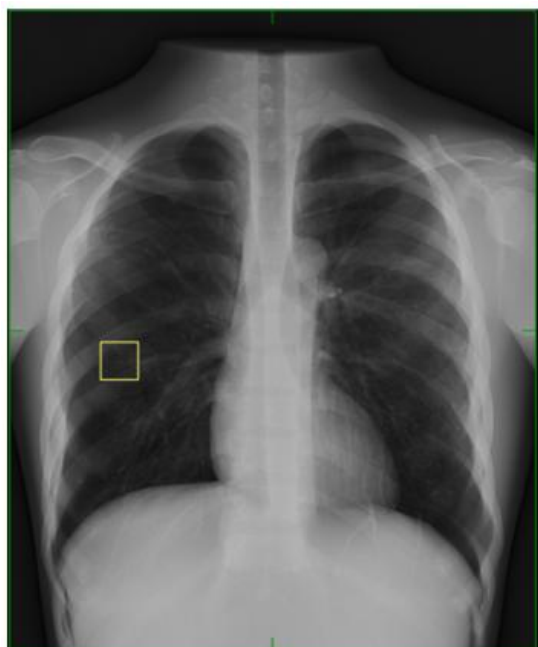
Next to the image the toolbar is displayed. The Toolbar can be adjusted in the system properties. The order or presence of buttons within the toolbar may differ in this manual by the system. A toolbar button will turn light blue  when activated. You can deactivate it by clicking on the button again.

	Overlay on/off (Patient name, DoB, REX etc.)
	Rotate the image 90° clockwise
	Rotate the image 90° anti-clockwise
	Free rotation
	Flip the image horizontal
	Flip the image vertical
	Invert the contrast of the image from positive to negative or vice versa
	Pan the image
	High Quality mode
	Zoom in
	Zoom out
	Make the image fit to the screen
	Left marker
	Right marker
	Region of Interest. The brightness (density) of the photo using the ROI can be modified.
	Crop box. The system will automatically put a crop box around the X-ray field. The content of this crop box is sent to the specific destinations. With this button the crop box size can be changed.
	Normally, the area outside the X-ray field will be blackened, called black mask. With this button the black mask can be customized.
	The currently selected photo is rejected and the same protocol is

	automatically added tot the study to retake the x-ray. The rejected image will not bet sent to a destination, but will remain on the workstation. The rejected image can be restored by clicking on Restore .
Reject	The currently selected photo is rejected. The rejected image will not be sent to a destination, but will remain on the workstation. The rejected image can be restored by clicking on Restore .
Repeat	Repeat the selected x-ray. This will automatically add the same protocol to the study without deleting the image.
Reset	All actions will be undone
	Undo the last action
	Make a copy of the image
	Change WW/WL
	Activate/deactivate Advanced Edge Enhancement

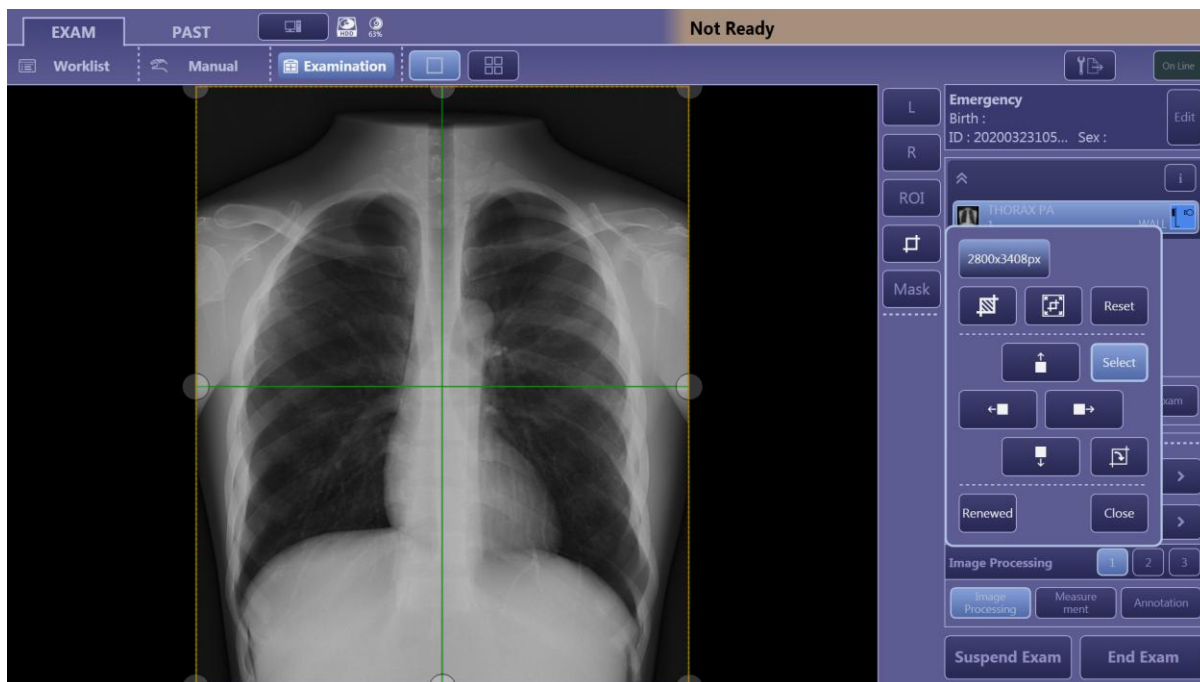
4.4.24.1 ROI (Region Of Interest)

When the **ROI** button is activated you can adjust the brightness (density) of the image by replacing the ROI. By clicking on the region of interest (for example a lung or a vertebra in the below examples) the brightness will change matching the new ROI. Click on **OK** to save the new ROI/brightness



4.4.24.2 Crop box

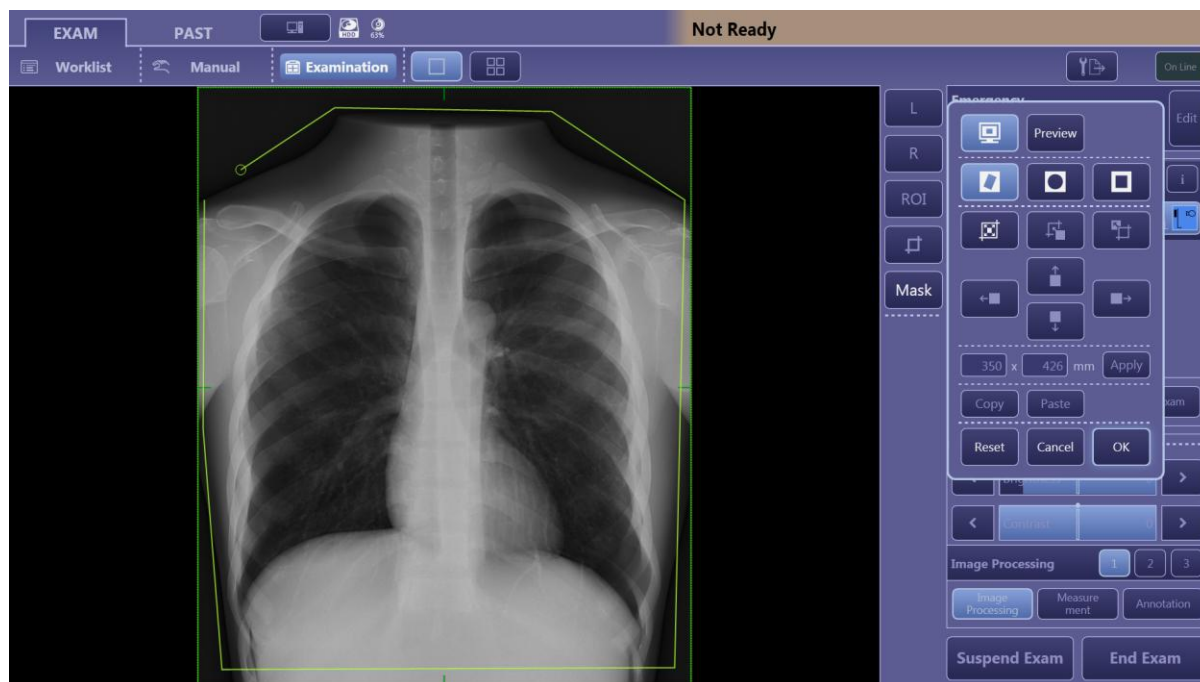
The system automatically puts a crop box around the x-ray field. Only the content inside this crop box is sent to the specific destinations. With this button the crop box size can be changed. You can change this crop box by clicking on 2 opposite corners with the left mouse button. You can also change the crop box by dragging the dots around the crop box.

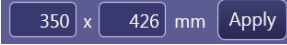


	Crops the irradiated field only
	Crops the whole detector
	All actions will be undone
	Select/deselect the crop box
	Move the crop box upwards
	Move the crop box downwards
	Move the crop box to the left
	Move the crop box to the right
	Turn the crop box 90° clockwise
	Select renewed to make a new crop box by clicking on two opposite corners.

4.4.24.3 Mask

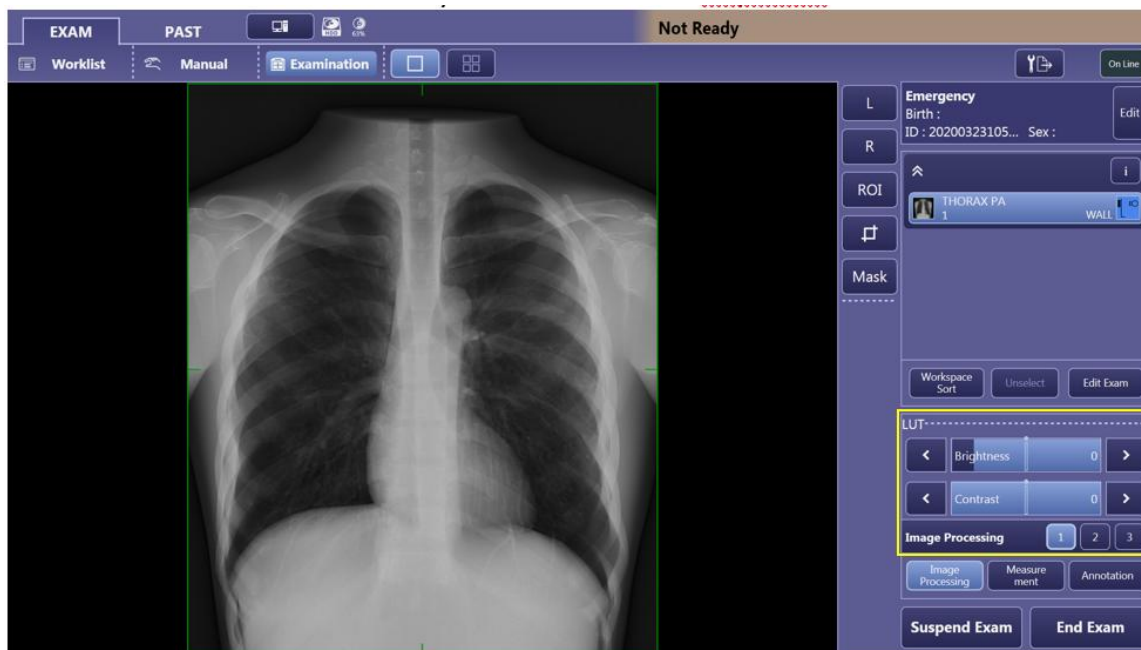
The system automatically gives the area outside the exposed area a black mask. With this button it is possible to change this black mask area. When the **Mask** button is activated you can draw a self-defined field. By pressing the left mouse button, you can create up to 12 anchor points to define the field. The black mask on the system is shown semitransparent but on a PACS screen or on a hard copy it will be shown completely black. Click on **OK** to confirm.



	Polygonal, round or square mask
	Fit the mask to the current crop box
	Move the mask box to the center of the crop box
	Move the crop box the center of the mask box
	Move the mask upwards, downwards, left or right
	Define a specific size of the mask
	Copy/paste the mask
	All actions will be undone

4.4.25 Image Processing

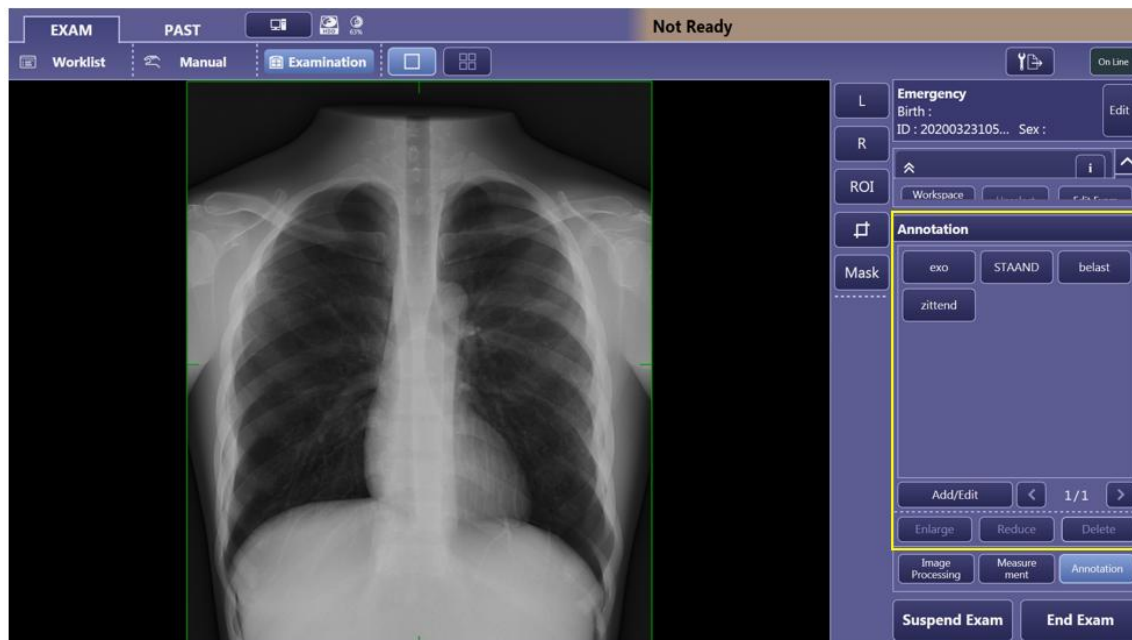
In image processing 3 tabs are available. The first tab is available for every user, the second and third tab is only available for the key users.



The first tab contains 2 sliders for Brightness and Contrast. You can increase the brightness of the image by sliding the slider to the right and you can decrease the brightness by sliding the slider to the left. To increase the contrast of the image you can slide the slider to the right. To decrease the contrast, move the slider to the left.

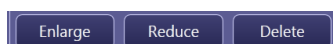
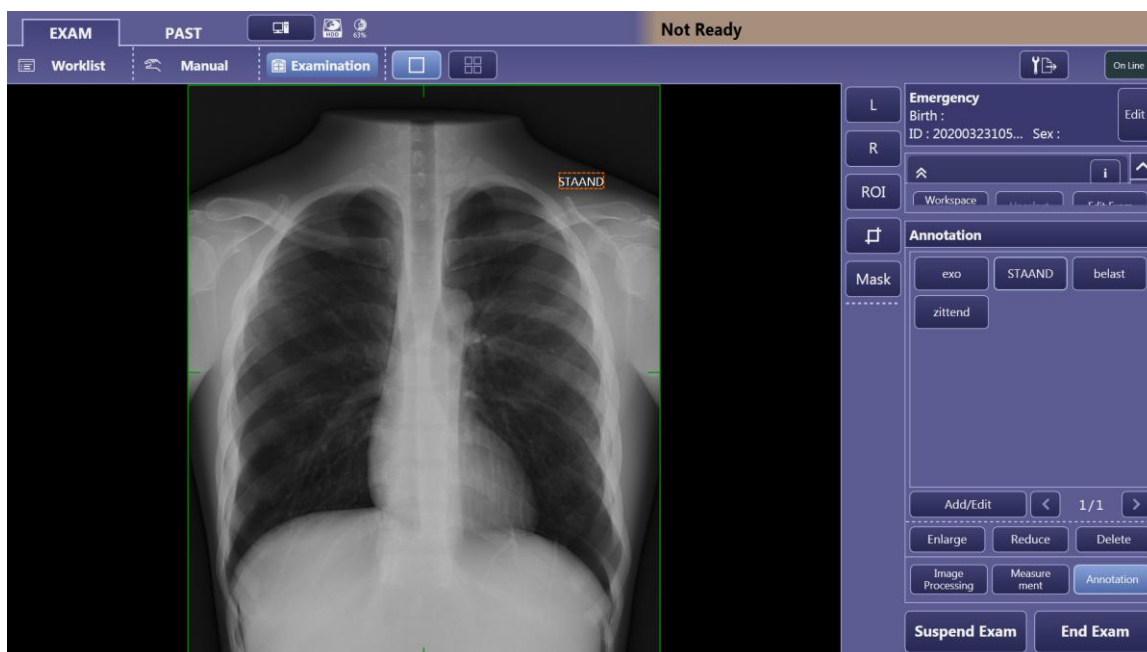
4.4.26 Annotation

It is possible to add annotation to the image. You can either add existing annotation or you can type annotation yourself.



Add existing annotation

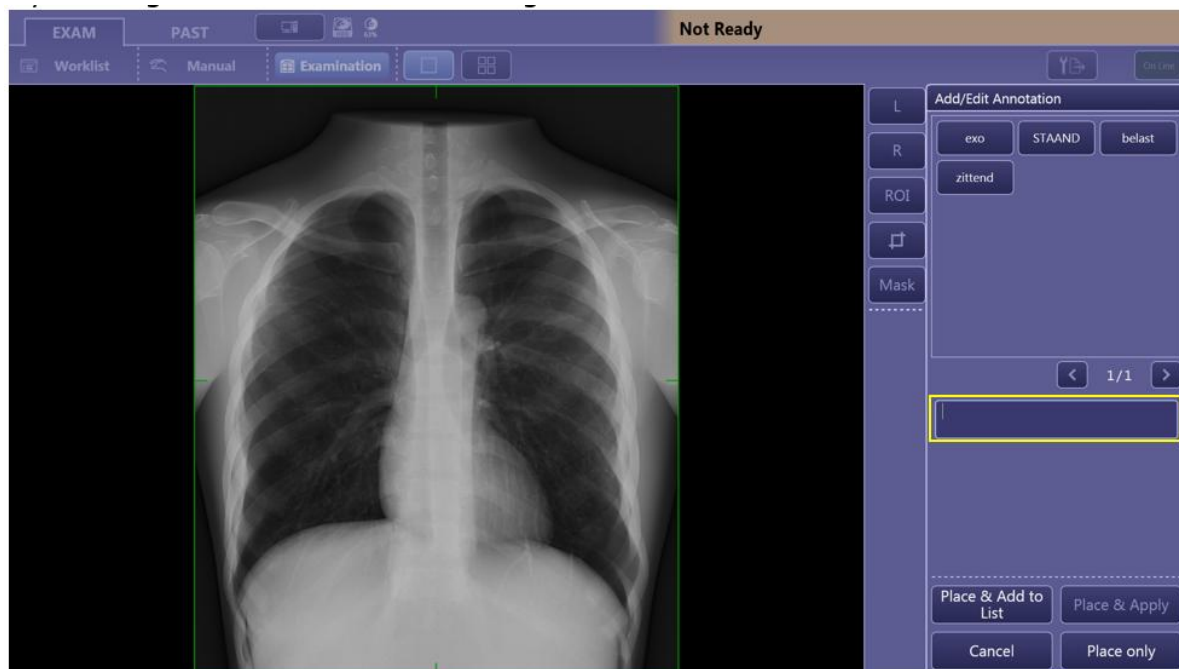
By clicking on the desired annotation, the annotation will appear in the upper right corner of the image. Move the annotation to the desired place by clicking on the desired space. You can move the annotation again by clicking on the annotation (an orange box will appear) and then clicking on the desired space.



Enlarge, Reduce or Delete the annotation.

4.4.26.1 Add new annotation

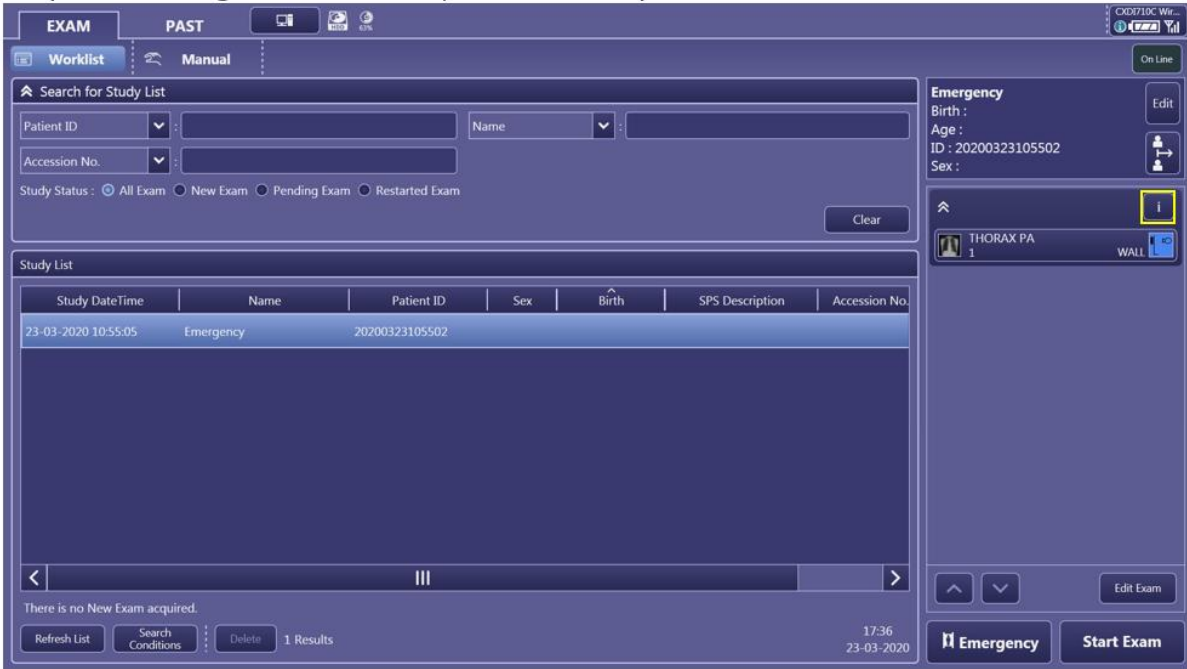
By clicking on **Add/Edit** the following screen will show:



Type the desired annotation. By clicking on **Place & Add to List** the annotation will be added to this image and to the annotation list for future use. By clicking on **Place only** the annotation will only be added to this image.

4.4.27 Protecting images for deletion

To protect images for deletion, click on the patient and click on .



EXAM **PAST**   

Worklist **Manual**

Search for Study List

Patient ID: Name:

Accession No.:

Study Status: ☒ All Exam ☐ New Exam ☐ Pending Exam ☐ Restarted Exam

Clear

Study List

Study DateTime	Name	Patient ID	Sex	Birth	SPS Description	Accession No.
23-03-2020 10:55:05	Emergency	20200323105502				

There is no New Exam acquired.

Refresh List Search Conditions Delete 1 Results

17:36
23-03-2020

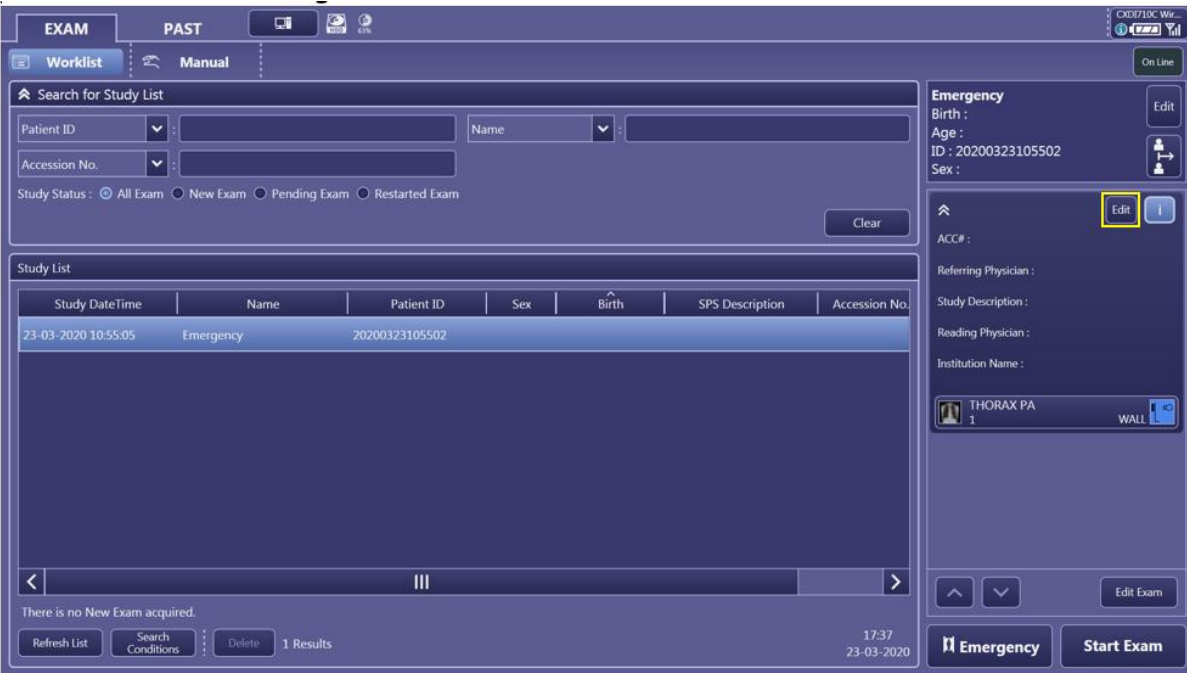
Emergency Birth: Age: ID: 20200323105502 Sex: Edit

THORAX PA 1 WALL

Edit Exam

Emergency **Start Exam**

After this the following screen will show:



EXAM **PAST**   

Worklist **Manual**

Search for Study List

Patient ID: Name:

Accession No.:

Study Status: ☒ All Exam ☐ New Exam ☐ Pending Exam ☐ Restarted Exam

Clear

Study List

Study DateTime	Name	Patient ID	Sex	Birth	SPS Description	Accession No.
23-03-2020 10:55:05	Emergency	20200323105502				

There is no New Exam acquired.

Refresh List Search Conditions Delete 1 Results

17:37
23-03-2020

Emergency Birth: Age: ID: 20200323105502 Sex: Edit

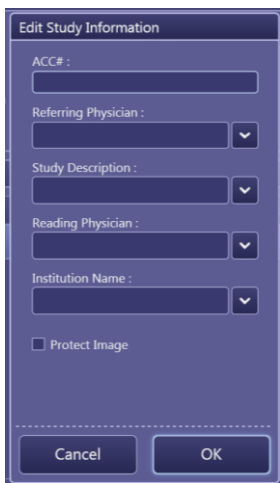
ACC#: Referring Physician: Study Description: Reading Physician: Institution Name:

THORAX PA 1 WALL

Edit Exam

Emergency **Start Exam**

Click on  and the following screen will show:



Edit Study Information

ACC# :

Referring Physician :

Study Description :

Reading Physician :

Institution Name :

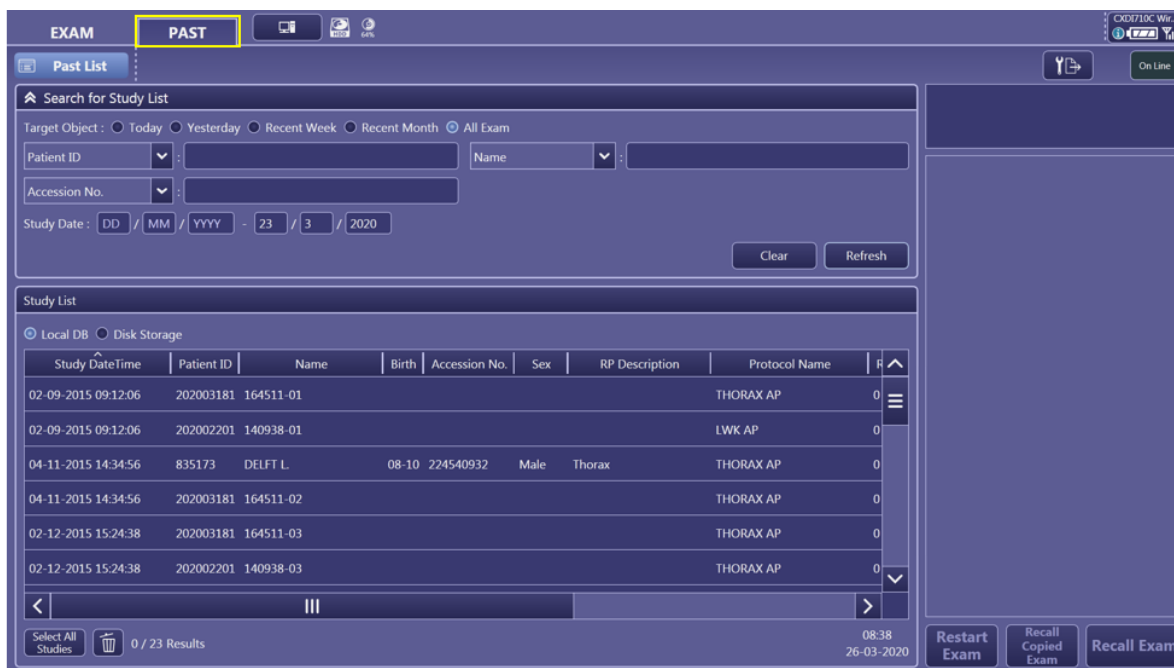
☐ Protect Image

Cancel OK

Click on  and on .

4.4.28 Past Screen

In the Past List old images can be retrieved. The retrieved images can be changed and resend.



PAST

Past List

Search for Study List

Target Object : ☐ Today ☐ Yesterday ☐ Recent Week ☐ Recent Month ☒ All Exam

Patient ID : Name :

Accession No. :

Study Date : DD / MM / YYYY - 23 / 3 / 2020

Clear Refresh

Study List

☒ Local DB ☐ Disk Storage

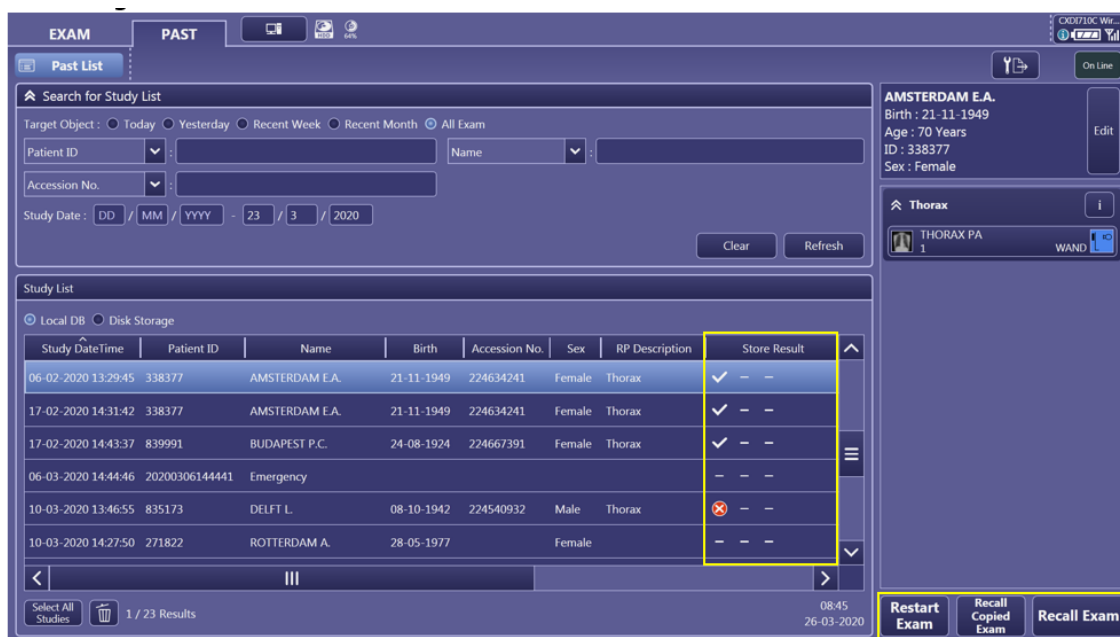
Study DateTime	Patient ID	Name	Birth	Accession No.	Sex	RP Description	Protocol Name	
02-09-2015 09:12:06	202003181	164511-01					THORAX AP	0
02-09-2015 09:12:06	202002201	140938-01					LWK AP	0
04-11-2015 14:34:56	835173	DELFT L	08-10	224540932	Male	Thorax	THORAX AP	0
04-11-2015 14:34:56	202003181	164511-02					THORAX AP	0
02-12-2015 15:24:38	202003181	164511-03					THORAX AP	0
02-12-2015 15:24:38	202002201	140938-03					THORAX AP	0

Select All Studies 0 / 23 Results

08:38 26-03-2020

Restart Exam Recall Copied Exam Recall Exam

It is possible to search in this list by study date (today, yesterday, recent week, recent month, all exam or a specified date), by Patient ID, name, Accession No., RP description and Protocol Name. After selecting the correct patient, the following screen will show:



EXAM PAST

Past List

Search for Study List

Target Object: ☐ Today ☐ Yesterday ☐ Recent Week ☐ Recent Month ☒ All Exam

Patient ID: Name:

Accession No.:

Study Date: DD / MM / YYYY - 23 / 3 / 2020

Clear Refresh

Study List

☒ Local DB ☐ Disk Storage

Study Date/Time	Patient ID	Name	Birth	Accession No.	Sex	RP Description	Store Result
06-02-2020 13:29:45	338377	AMSTERDAM E.A.	21-11-1949	224634241	Female	Thorax	✓ - -
17-02-2020 14:31:42	338377	AMSTERDAM E.A.	21-11-1949	224634241	Female	Thorax	✓ - -
17-02-2020 14:43:37	839991	BUDAPEST P.C.	24-08-1924	224667391	Female	Thorax	✓ - -
06-03-2020 14:44:46	20200306144441	Emergency					- - -
10-03-2020 13:46:55	835173	DELFT L.	08-10-1942	224540932	Male	Thorax	✗ - -
10-03-2020 14:27:50	271822	ROTTERDAM A.	28-05-1977		Female		- - -

Select All Studies 1 / 23 Results

08:45 26-03-2020

AMSTERDAM E.A.
Birth : 21-11-1949
Age : 70 Years
ID : 338377
Sex : Female

Thorax
THORAX PA 1
WAND

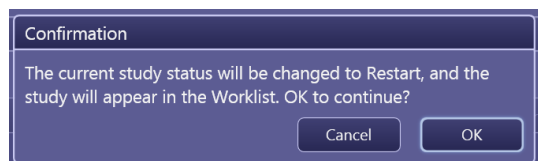
Restart Exam **Recall Copied Exam** **Recall Exam**

In the **Store Result** column it is possible to see whether the image(s) have been sent to the destination(s)

✓ - -	All images have been sent to one destination.
- - -	No images have been sent.
✗ - -	Failed to send images to the destination.

4.4.29 Restarting an exam

After ending an exam, it is possible to restart an exam and add protocols to this study. This is possible by selecting the correct study and by clicking on **Restart Exam**. After this the following screen will show:



Confirmation

The current study status will be changed to Restart, and the study will appear in the Worklist. OK to continue?

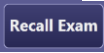
Cancel OK

By clicking on **OK**, the study will reappear in the worklist. After selecting the patient again in the worklist and starting the exam you can add protocols to this study. Images that already have been taken will appear with a yellow overlay . It is not possible to post process these images in the exam screen. After ending the exam this will be possible again in the past list.

4.4.30 Recalling an exam

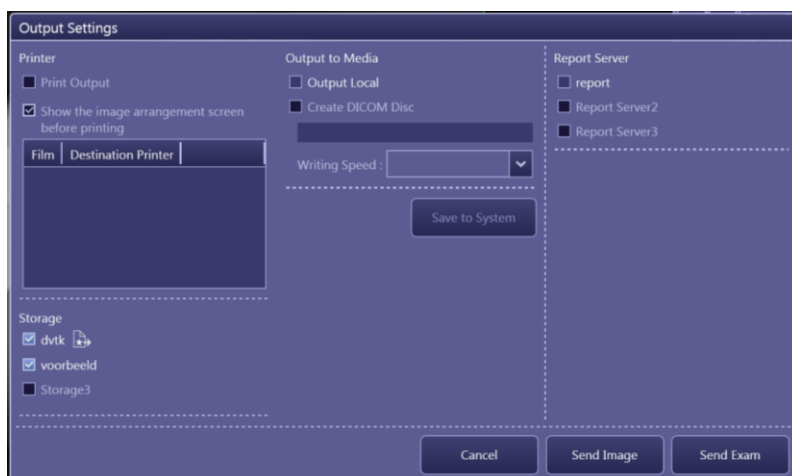
After ending an exam, it is possible to recall an exam to perform post-processing on the image(s). This is possible by selecting the correct study and by clicking on  or .

By clicking on  the system will make a copy of the whole exam and will recall this copy.

By clicking on  the system will only recall the exam. After this it is possible to post-process these image(s).

4.4.31 Resending images

It is possible to resend images from both the Past List Screen and after recalling an exam with the  button. Select the study (in the past list) or the image (after recalling the exam) and click on . The following screen will show:



The destinations with a checkmark are the active destinations. Changing the active destination will change it for this time only.

By clicking on , only the selected image will be resent (this button will only be available when you are resending after recalling an exam).

By clicking on , the whole exam will be resent.

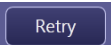
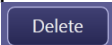
4.4.32 System Screen

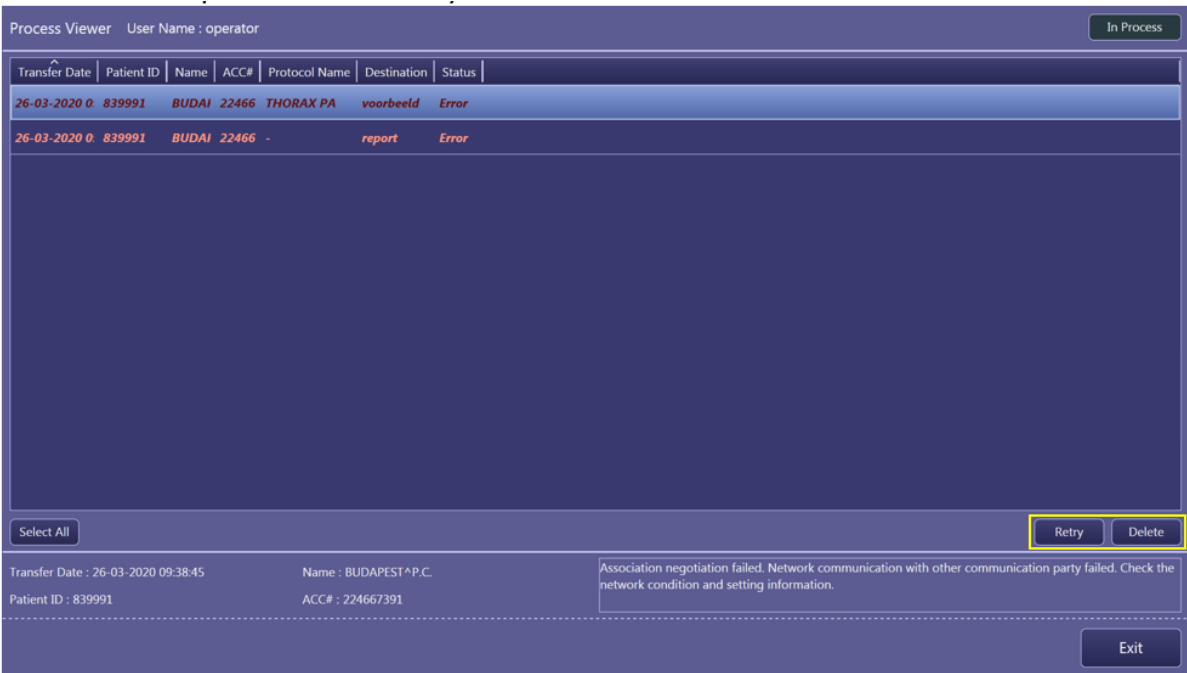
In the system screen you can log out, shut down the system or consult the Process Viewer.

4.4.33 Logging out

Click on  to log out.

4.4.34 Process Viewer

In the Process Viewer it is possible to see errors transferring data. By selecting the process, you can either  or  the process. By clicking on  you can select all processes to retry or delete.



The screenshot shows the 'Process Viewer' window with the user name 'operator'. It contains a table with two rows of data, both marked as 'Error'. Below the table are buttons for 'Select All', 'Retry', and 'Delete'. At the bottom, there is a detailed error message and an 'Exit' button.

Transfer Date	Patient ID	Name	ACC#	Protocol Name	Destination	Status
26-03-2020 0	839991	BUDAI 22466	THORAX PA	voorbeeld	Error	
26-03-2020 0	839991	BUDAI 22466	-	report	Error	

Buttons:   

Transfer Date : 26-03-2020 09:38:45
Patient ID : 839991
Name : BUDAPEST^P.C.
ACC# : 224667391

Association negotiation failed. Network communication with other communication party failed. Check the network condition and setting information.



4.4.35 Filtration Indication

Filters are materials (for example metal sheets) placed in the x-ray beam between the window and the patient that are used to attenuate the low-energy (soft) x-ray photons from the spectrum. Filtering is the removal of these low energy x-rays from the beam spectrum which would otherwise not contribute to image quality but would add to patient dose and scatter.

There are two types of filtration.

- Permanent filtration: inherent filtration from components in the x-ray tube, collimator, docking, AEC etc....
- Added filtration: added filtration from interchangeable metal sheets (Al, Cu) integrated to the collimator.

All Items in the x-ray beam shall marked with the mm AL equivalent:

Item	AL Equivalent (mm)
Tube assembly:	0.7 mm AL
Collimator permanent	2.0 mm AL
Collimator selectable	- 1mmAL+0.1mmCu - 1mmAL+0.2mmCu - 2mmAL - none

4.4.36 Software options

There are several software options available for the trauma arm.

Free rotation

Is a free option on every system. With this option is it possible to angulate de angle that you need with the image. This button will be visible in the toolbar.

Advanced Edge Enhancemnt

Improved visualizqation of tubes, catheters and bone details. This option is ideal for clearly displaying catheters and lines. You can set it als default in the protocols or it is possible to set the button in the toolbar.

Scatter Correction

Is a digital grid. You can use this options when you have wireless examinations without a physically grid, like chest in bed. The software works by creating a scatter model, which is subsequently subtracted from the image. The result is an image with reduced scatter and increased contrast. You can set this option on or off in the image processing.

Intelligent Noise Reduction

Intelligent NR is a substantial step up from conventional noise reduction. Intelligent NR will provide radiologists high-quality diagnostic images containing significantly less grainy noise with no noticeable loss of detail. This should make improved diagnosis possible for their patients.

Intelligent NR makes it possible to reduce the noise content without losing the fine details of the anatomy, even in low dose regions. For this reason, it is superior to conventional noise reduction. Intelligent NR results in an optimal diagnostic environment, especially for infants and pediatric patients where dose is a prevailing concern. But the image improvement will benefit all patients and exam types by providing superior images even in inherently noisy conditions such as the dense anatomy of the abdomen. With Intelligent NR, there is the potential for significant dose reduction while retaining equivalent image quality (noise content).

It is possible to set the Intelligent Noise Reduction on or off in the protocols. When you set the INR of a protocol the system will use the conventional noise reduction.

Built-in AEC Assistance

Whit the Elite (20 series) detector is possible to make automatically terminaded exposures without the use of an addintional AEC receptor. You can set it on or off in the protocol. You should also specify the AEC backup, desired AEC fields, and detector orientation in your protocol so that the correct fields are used. You can also enable autorotation to ensure the AEC fields are always in the correct position.



5 Cleaning and disinfection

Note:	Test cleaning products and disinfectants to ensure there is no adverse effect on the cover.
--------------	--

5.1 Cleaning

Note:	The mains power must be disconnected before proceeding with the cleaning operations.
--------------	---

- Clean the equipment with a damp cloth or a cotton swab. Use a non-abrasive cleaning material. To moisten use water or lukewarm water with a diluted solution of liquid detergent of the common household type.
- Do not use abrasive cleaners or organic solvents or cleaning agents containing solvents (e.g. alcohol, ether, etc.). Do not use hydrogen peroxide which enhances the corrosion of metals.
- Do not use spray liquids! Agents for cleaning fluids must not enter under any circumstances within the enclosures!
- For cleaning the screen, use a microfiber cloth. Any conventional screen cleaner can be used. Be aware that buttons can be pressed accidentally on the touchscreen while cleaning when the system is powered on.
- For cleaning the wireless detector we refer to the User's Manual of the detector.

5.2 Disinfecting

- For surface disinfection, it is recommended to use aqueous solutions commercially available for the disinfection of surfaces based on aldehyde or amphoteric surfactants (e.g., Tego 103, Korsolin). Disinfectant substitutes based on phenolic and chlorine-containing compounds are partially aggressive and therefore normally to be avoided. The same limitations apply to agents which have an alcohol content when they are applied in undiluted form (e.g. for the disinfection of hands).
- Strictly prohibited are spray disinfectants, because the spray could penetrate inside the device. As a result, the safety of the appliance can no longer be guaranteed (damage to electronic components, formation of flammable mixtures of air and solvent vapors).
- Where the entire exam room is to be disinfected, remember to cover the unit with plastic sheeting.
- For disinfecting the wireless detector we refer to the User's Manual of the detector.



6 Troubleshooting

6.1 The detector does not become “Ready”

Possible Solution:

Is the correct detector selected?

Is the battery from the detector depleted?

Is the detector turned on?

Is the appropriate protocol selected?

Shows the Wi-Fi access point the blue light? Yes = connection. No = No connection

6.2 Image acquisition fails

Possible Solution:

Is there ready in the screen? Image acquisition is possible only when the status is on “Ready”.

6.3 Worklist cannot be retrieved

Possible Solution:

Is the acquisition computer Online (upper right corner)

Are there troubles with the network?

6.4 Image cannot be sent

Possible Solution:

Is the acquisition computer Online (upper right corner)

Are there troubles with the network?

6.5 Some columns are missing from the worklist

Possible Solution:

The columns are perhaps resized to small. Check width of the columns.

6.6 The exposure values are not set correctly for the selected protocol.

Possible Solution:

Is the system on “Ready”? The exposure settings are being set after the check if the detector is available.

Check the generator status with . If the button  is visible. If so, press this button the recover generator communication.

6.7 Incomplete protocols exist appears

This means that not all added protocols are used for exposure, check if are necessary exposures are being made.

6.8 Error while retrieving worklist or sending images

Possible Solution:

Probably there is a malfunction on the hospital network. Check all possible network settings, restart the acquisition computer. If problem remains call hospital IT department.

6.9 PACS is temporarily not available.

Possible Solution:

Press  and select the secondary PACS if available, or uncheck the PACS and save settings. Check back on when PACS is available again and resend from PAST list all un-send images.

6.10 DAP accumulated value has been exceeded

Possible Solution:

Press in the generator screen on DAP test (above kV). Select DAP reset. If error remains: temporarily end this patient and select a new one for resetting the DAP.

6.11 Error Image analysis processing failed

Cause

The image processing tries to identify the ROI in the image and it fails to do so.

Possible Solution:

Check image brightness and adjust if necessary, by setting the ROI.

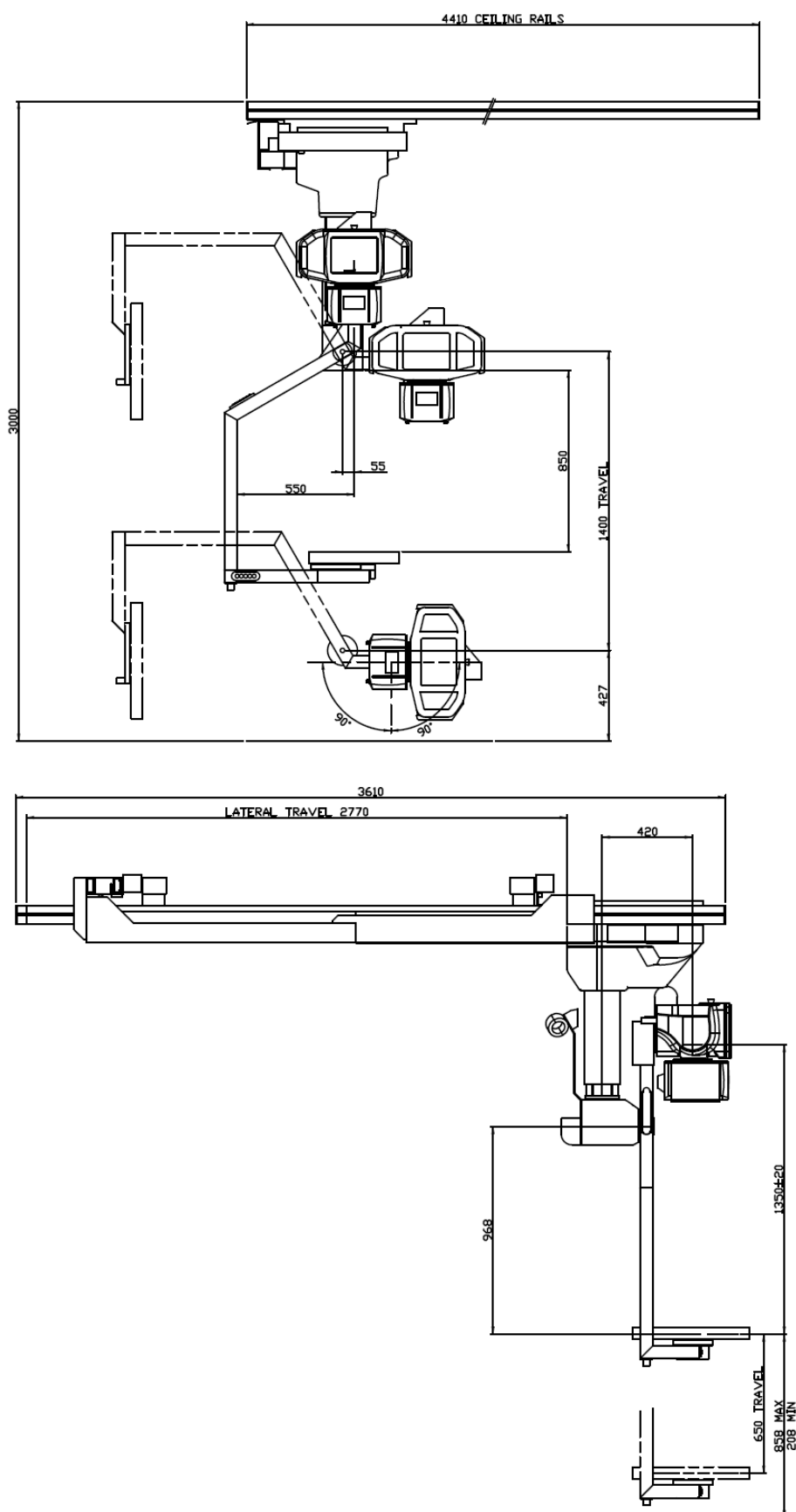
7 Technical Specifications

1. Specifications Detector			
	CXDI-720C Wireless	CXDI-420C Wireless	CXDI-820C Wireless
Purpose	General Radiography		
Scintillator	CsI (Cesium Iodide)		
Effecttime imaging area	350 x 426 mm	426 x 426 mm	350 x 274 mm
Resolution	4.0 lp/mm		
DQE	Typical 74% (0 lp/mm) / 67% (0.5 lp/mm)		
Gray scale	A/D: 16 bit		
Pixel size	125 x 125 μ m		
Image matrix size	2800 x 3408 pixels	3408 x 3408 pixels	2800 x 2192 pixels
Dimensions Sensor	384 x 460 x 15.7 mm	460 x 460 x 15.7 mm	384 x 307.5 x 15.7 mm
Weight Sensor	2.5 kg	3.0 kg	1.8 kg
Load capacity	310 kg (over the whole area of the detector surface)		
Preview image access time	+/- 1 sec		
Cycle time	+/- 4 sec		

2. Specifications X-ray generator	
Type, CMP200DR, 150 kV / 50 kW (optional 65 or 80 kW available)	Line voltage: 3 phase 400/480 VAC +/- 10% / 65 kVA / 50/60Hz
	High frequency generator 200kHz
kV range	50 to 150 kV +/- 5% + 1 kV
mA range	10 to 630 mA (50kW)
mAs range non-AEC	Tube dependent, 0,1 to 630 mAs
Time range	1.0 to 6300 ms
Technique selections	kV with AEC option
	kV/mAs
	kV/mA/time

3. Specifications X-ray tube and Collimator		4. Specifications Ceil stand
X-ray tube	400 kHU high capacity	Motorized vertical column up to 1750mm vertical travel (depending on ceiling height)
	0,6/1,2 focal spot	Manual X/Y/Alpha/Beta and Tube rotation with electromagnetic brakes
	12° target angle 150 kV	X-ray tube rotation (Alpha) 270°
	Low speed 24/58 kW	C-arm rotation around the telescope vertical axis (Beta) 300°
	High speed 38/91 kW	C-arm rotation around the telescope horizontal axis (Alpha) +120° -120°
Collimator	Manual	Large U-shape depth of 550mm for optimal positioning
	LED lamp	SID of 1350 mm
	DAP integration	Dockingstation with AEC and removable grid

8 Dimensions



9 Glossary

Units of Measurement

°C	degrees in Celsius
cm	centimeters (metric)
hPa	hectopascal. Related to measurement of barometric pressure.
Kbit/sec	kilobits per second. Related to communication protocols.
kg	kilogram (metric)
m / M	meters (metric)
N	Newton. It is the International System of Units (SI) derived unit of force.
s	Second. A measurement of Time.

Radiographic Terminology

AEC	Automatic Exposure Control (AEC)
Anode	The positive terminal of an x-ray tube
APR	Anatomical programmed radiography
Bucky	It is a radiology device that holds the x-ray film cassette or Detector and moves the grid during x-ray exposure.
Bucky grid	A component used in radiography to prevent the scattered rays from reaching the film, aiding in the clarity and definition of the picture. May also be called Potter–Bucky grid.
CAN / CAN bus	Controller Area Network (CAN). CAN bus is a message-based protocol, designed for use in areas such as industrial automation and medical equipment
Cathode	A negative electrode from which electrons are emitted. In x-ray tubes, the cathode usually consists of a helical tungsten filament behind which a molybdenum reflector cup is located to focus the electron emission toward the target of the anode.
Central Ray	The center point of the X-ray beam.
Collimator	A device, sometimes called a diaphragm, which filters and controls the direction and the dimensions of the x-ray beam.
CR	Computed Radiography (CR). Uses an imaging plate made of photo stimulated phosphor storage in place of a film to create the image.
DAP	Dose Area Product meter. An electro-mechanical device used in assessing the radiation risk from diagnostic X-rays examination and interventional procedures.
Diaphragm	A collimation device, which filters and controls the direction and the dimensions of the x-ray beam.
DICOM	Digital Imaging and Communications in Medicine
DR	DR Digital Radiography (DR). Also called Direct Radiography. Employs the use of a special electronic x-ray detector which converts the radiographic image into a digital picture for review on a computer monitor.
FFD (DFF)	Focal Film Distance. More common term for use of Focal Distance with traditional film imaging systems. This is determined by the Focal Point of the X-ray tube to the surface of the media in use. (see also SID)
Filament	A coiled tungsten wire, which emits electrons when heated to incandescence.
Filter	The material (usually aluminium) placed in the useful primary beam of x radiation.

flat panel	Generic term for use to describe a digital-type media radiography component to create a digital image
Focal Distance	Also seen as 'FFD' or 'SID'. The measurement between the Focal Point of the X-ray tube and the type of media in use.
Ionizing Radiation	Electromagnetic radiation (e.g., x rays or gamma rays) or particulate radiation (e.g., electrons, neutrons, and protons) capable of ionizing air directly or indirectly.
kVp	Peak kilovoltage (kVp). It is the maximum voltage applied across an X-ray tube.
mA	Milliamperage refers to the current flow from the cathode to the anode, which, in turn, regulates the intensity of radiation emitted by the x-ray tube and, hence, directly influences the radiographic density.
mAs	Milliampere-Seconds (mAs): The product of the x-ray tube operating amperage and exposure time, in seconds.
MPD	Maximum Permissible Dose (MPD). For radiation workers, 50 mSv per year is permissible. Oblique An angular view of a surface or object.
OFD	Object-to-Film Distance (OFD). Distance between the object or skin and the cassette or Film/detector.
PACS	Picture Archiving and Communication System. A Technology which provides economical storage of, and convenient access to, images.
Projection	A term for the position of a part of the patient with relation to the x-ray film and the x-ray beam.
Rad	Radiation Absorbed Dose (RAD). A unit of measurement for the absorbed dose of any type of ionizing radiation in any medium.
Radiation	The emission and propagation of energy, in the form of waves or particles, through space or a material medium.
Radiograph	A visible image on a radiation-sensitive film emulsion to ionizing radiation that has passed through an area, region, or substance of interest.
Radiography	The technical process of positioning, exposing, and processing radiographs.
RIS	Radiology Information System
SID	Source to image distance. More common term of Focal Distance for use with digital imaging systems.
Source	The point of emanation of gamma or x rays when used as an origin of radiation. The X-ray tube.
TFD	Target-Film Distance (TFD). This is the same as focal-film distance (FFD).
Wall Bucky	Stand Also written as: Wall Bucky, Wall stand, Wall Receptor Stand. Mobile radiology receptor component mounted to vertical column that is usually attached to the wall. This term is generically used also if different media is used such as 'bucky', 'DR' or CR' receptor components. (see also Teleradiograph)
Workstation	Operator working station. Mainly in reference to a work area dedicated to the X-ray generator console and Digital Imaging PC workstation for use in DR or CR applications.
X-ray	A type of electromagnetic radiation characterized by wavelengths of 100 angstroms or less.
X-ray Beam	The radiation emerging from an x-ray generator or source.

Patient Orientation

Normal Anatomical Position	To avoid misunderstanding, a standard position of the human body is arbitrarily taken to be the erect (standing) position with feet flat on the floor, heels together, upper extremities at the sides, and palms, toes, and eyes directed forward.
AP	Anteroposterior (AP). The prefix indicates the surface from which the Central Ray (CR) enters the part and the suffix indicates the surface from which the CR emerges.
PA	Posteroanterior (PA). The prefix indicates the surface from which the Central Ray (CR) enters the part and the suffix indicates the surface from which the CR emerges.
Anterior, frontal, or ventrum	The front side of the body.
Posterior or dorsum	The back, or dorsum, of the body.
Median	Pertaining to the midline of the body.
Lateral	Away from the midline or lateral side of the body.
Proximal	Nearest to a point under consideration or the point of origin.
Superior	Above
Inferior	Below
Cephalic	Towards the head
Caudal	Toward the feet
Sagittal plane	plane Any vertical plane that divides the body into right and left unequal portions.
Transverse plane	Also known as Axial or Horizontal plane, it is an X-Z plane that is parallel to the ground which divides the body into upper and lower portions (head from feet).
Coronal plane	Also known as Frontal plane, it is a Y-X plane that is perpendicular to the ground. A vertical plane that divides the body into front and rear portions.
Sagittal plane	Also known as Lateral plane, it is a Y-Z plane that is perpendicular to the ground. The vertical plane that divides the body into right and left halves.

Body Axes

H = Head
 F = Feet
 L = Left
 R = Right
 P = Posterior
 A = Anterior

