

ENGLISH

ORTHOPANTOMOGRAPH® OP300

3D Dental X-Ray System

User Manual



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1 Introduction

1.1 ORTHOPANTOMOGRAPH® OP300

INSTRUMENTARIUM DENTAL™

ORTHOPANTOMOGRAPH® OP300 x-ray unit (hereafter called “OP300”) is a dental x-ray system for producing high quality digital images of dentition, TM-joints and skull. In order to take images with OP300 you need a suitable PC hardware connected to the OP300 unit and CLINIVIEW™ software (or suitable third party software via TWAIN driver) to capture and manage images.

OP300 performs the following procedures:

Panoramic

- Standard panoramic
- Pediatric panoramic
- Wide arch panoramic
- Bitewing
- TMJ, PA projection
- Ortho TMJ, axial corrected lateral projection
- Maxillary sinus
- Ortho Zone enhanced panoramic
- Orthogonal panoramic

Cephalometric (optional)

- Cephalometric lateral projection
- Cephalometric pediatric lateral projection
- Cephalometric postero-anterior (PA) projection
- Reverse Towne projection
- Waters view
- Carpus program (optional) (Not available in USA and Canada)

3D SFOV (optional) H x W

- 61x41 mm Field of View
- 61x78 mm Field of View

3D MFOV (Maxio) (optional) H x W

- MFOV (Maxio) 50 x 50 mm Field of View
- MFOV (Maxio) 61 x 78 mm Field of View
- MFOV (Maxio) 78 x 78 mm Field of View
- MFOV (Maxio) 78 x 150 mm Field of View
- MFOV (Maxio) 130 x 150 mm Field of View (optional)

1.2 References

The following instructions are delivered with in the OP300 installation manual:

- Firmware update instructions
- Calibration instructions
- Cephalostat upgrade instructions
- Cephalostat side changing instructions

The following instructions are separate and can be ordered from customer service:

- 3D upgrade instructions are delivered with the 3D upgrade kit.

1.3 Intended use

OP300 must only be used and operated by healthcare professionals and other qualified professionals. OP300 must only be used to take panoramic, cephalometric and 3D images of the dento-maxillofacial complex of the human skull. It must not be used to take images of any other part of the human body.

Panoramic and 3D exposures should not be used if conventional intraoral radiographic images (like bitewing exposures) would be sufficient.

Cone beam computed tomography images are not adequate for the analysis of soft tissue.

CAUTION! *USA only: Federal law restricts this device to sale by or on the order of a dentist or other qualified professional.*

1.4 Associated documentation

- OP300 user manual
- OP300 installation manual
- The CLINIVIEW™ software user manual
- The CLINIVIEW™ software installation manual
- The user manual supplied with the dental imaging software
- The installation manual supplied with the dental imaging software
- The user manual supplied with the 3D imaging software
- The installation manual supplied with the 3D imaging software

1.5 Abbreviations used in this manual

FOV = Field Of View. The cylindrical 3D volume that is reconstructed by the system.

ROI = Region Of Interest. The anatomical area or region of the patient that you are interested to examine.

FH = Frankfort-Horizontal

H = Horizontal

1.6 Warnings and precautions

1.6.1 Warnings to be observed during use

The unit may be dangerous to the user and the patient, if the safety regulations in this manual are ignored, if the unit is not used in the way described in this manual and/or if the user does not know how to use the unit.

The unit must only be used to take the dental x-ray exposures described in this manual. The unit must NOT be used to take any other x-ray exposures. It is not safe to use the unit to take x-ray exposures, that it is not designed for.

Only professionally qualified dental and/or medical personnel are allowed to operate the unit and carry out any diagnoses based on output from the unit.

Because the x-ray limitations and safety regulations change from time to time, it is the responsibility of the user to make sure that all the valid safety regulations are fulfilled.

When taking an x-ray exposure of a patient with exceptional anatomy (typically very tall or large) use the Test-mode (no x-rays) first to make sure that patient can be positioned correctly to the unit and for checking that the unit doesn't hit the patient.

Operator should maintain visible contact with the patient and technique factors. This allows immediate termination of radiation by the release of the exposure button in the event of a malfunction or disturbance.

It is the responsibility of the doctor to decide whether x-ray exposure or any additional exposures are justified and necessary.

The minimum height of patient that can be x-rayed is 120 cm (3.9ft / 47.2in) and the maximum is 200 cm (6ft /78in). These heights only apply to patients with normal anatomy.

Always use the lowest suitable x-ray dose to obtain the desired level of image quality.

Avoid taking x-ray exposures of pregnant women.

When taking an x-ray exposure of a child always use the lowest possible x-ray dose, the smallest possible image area and the lowest possible resolution that allows you to perform the required diagnostic task.

If the patient is using a pacemaker, consult the manufacturer of the pacemaker before taking an exposure to confirm that the x-ray unit will not interfere with the operation of the pacemaker.

Decontaminate all the surfaces that the patient is in contact with after every patient to prevent cross infection.

Decontaminate all device accessories that contact the patient during a radiographic examination.

Do not open or remove any of the unit's enclosures. No user serviceable parts inside.

The customer must ensure that the siting environment fulfills the requirements listed in the Installation manual. Special attention must be paid to the strength of the floor and wall materials, electrical mains and radiation protection. It is the responsibility of the customer to ensure that the site is large enough for the patients.

The unit contains toxic materials that need to be handled properly when disposing the unit. Return the unit to the dealer in the end of its life cycle.

Excessive dust should be cleaned from the unit for free airflow and cooling. Switch of the unit before cleaning.

Always follow the instructions for patient positioning and imaging procedures instructed in the User Manual.

In case of water damage/water dropping over the product, call for service technician to ensure the product is fully operational according to specification.

1.6.2 Warnings for cross infection

Always use available disposable protective covers with the patient positioning accessories:

- Bite fork cover
- Chin support cover
- Head support cover
- Nose support cover
- Ear holder cover

1.6.3 General warnings

Personnel operating the device must be adequately trained with respect to the technological principles of operation and radiation protection when using cone beam computed tomography (CBCT) imaging.

This unit complies with the EMC (Electromagnetic Compatibility) according to IEC 60601-1-2. Radio transmitting equipment, cellular phones etc. shall not be used in close proximity of the unit as they could influence the performance of the unit.

The correct software and settings in the workstation are essential to the performance of the unit. Consult technical support to ensure correct setup.



Danger: Explosion hazard - do not use in the presence of flammable anesthetics, gases or vapors.

The unit is factory set to operate using a 230-240 $\pm$ 10 VAC power supply. Never connect the unit to a power supply different to the voltage marked on the unit.

The site must fulfill the environmental requirements in the installation manual chapter technical specifications.

There should be free space around the unit for safe operation.

To maintain patient safety it is mandatory to use an unshielded CAT6 Ethernet cable between the unit and the network or workstation, so that multiple chassis are not connected. Non-medical grade PC should not be used in patient environment.

This product itself complies with IEC 60601-1 medical safety standard but in order to the system incorporating also a PC to comply the standard, EITHER the PC has to be a medical PC OR the PC has to be located over 1,5 meters apart from the unit. The installer and the user of the system shall confirm that at least one of the above requirements is fulfilled. A PC is a medical one if it complies IEC 60601-1 standard and that is indicated in the accompanying documents of the PC. See chapter Technical specifications, Minimum PC Requirements, in user manual.

The unit shall be connected directly to the acquisition PC with an Ethernet cable. Connection through the LAN-network of the site is not allowed. Two network ports are needed in the PC in order to connect also to the site network.

All service operations must be made by authorized service personnel only.

The annual service as described in manual is mandatory for the correct and safe operation of the unit.

When taking exposures, operators and service personnel must protect themselves from radiation and remain at least two meters (six feet) away from the unit during exposure.

Protect the patient from scattered radiation by placing a protective lead apron over the patient.

The unit must be installed and serviced according to the unit Installation & adjustments manual by a qualified technician.

Only personnel trained and approved by the manufacturer of the unit are allowed to service the unit. 3D should not be used for routine or screening examinations in which a radiograph is taken regardless of the presence or absence of clinical signs and symptoms. 3D imaging examinations must be justified for each patient to demonstrate that the benefits outweigh the risks.

Where it is likely that evaluation of soft tissues will be required as part of the patient's radiological assessment, the imaging should be done using conventional medical CT or MR, rather than 3D imaging using Cone Beam technology.

Make sure that patient's thyroid glands are protected by a lead apron during the exposure.

The place where the unit is to be installed and the position from where the user will take exposures must be correctly shielded from the radiation that is generated when the unit is operated. Ensure to fulfill or exceed the requirements of your local regulations.

The unit or its parts must not be changed or modified in any way without approval and instructions from the manufacturer.

When servicing use only approved replacement parts supplied by the manufacturer.

The use of accessories not complying with the equivalent safety requirements of this equipment may lead to a reduced level of safety of the resulting system.

If this device is used with 3rd party imaging application software not supplied by the manufacturer, the 3rd party imaging application software must comply with all local laws on patient information software. This includes the Medical Device Directive 93/42/EEC and/or relevant legal requirements in the USA.

Do not connect any equipment to the unit that has not been supplied with the unit or that is not recommended by the manufacturer. The use of accessory equipment not complying with the equivalent safety requirements of this

equipment may lead to a reduced level of safety of the resulting system.

All protective covers must be properly installed before handing unit to the user or when operating the unit. Correct sharp layer should be chosen when using multilayer PAN images. See user manual chapter Multilayer PAN images for correct procedure.

1.7 Disclaimer

The manufacturer shall have no liability for consequential damages, personal injury, loss, damage or expense directly or indirectly arising from the use of its products. No agent, distributor or other party is authorized to make any warranty or other liability on behalf of the manufacturer with respect to its products.

1.8 Disposal

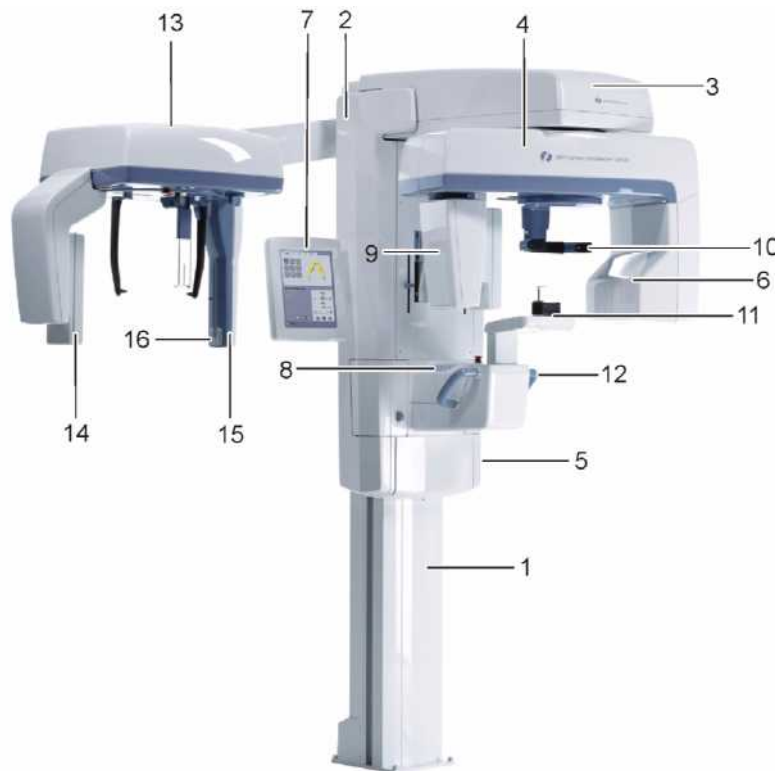
The device, its spare parts, its replacement parts and its accessories may include parts that are made of or include materials that are non-environmentally friendly or hazardous. These parts must be disposed of in accordance with all local, national and international regulations regarding the disposal of non-environmentally friendly or hazardous materials.

Unit has at least the following parts that should be regarded as non-environmental friendly waste products:

- Tubehead (Pb, oil)
- Collimator (Pb)
- All electronic circuits, electronic boards inside
- Sensor covers (EMC painted)

2 Unit description

2.1 Main parts and controls



1. Column
2. Carriage
3. Main support
4. Rotating unit
5. On/off switch (rear of carriage) and main fuses
6. Tubehead assembly
7. Touch screen display
8. Positioning panel
9. Sensor head
10. Head support
11. Chin rest
12. Handles
13. Cephalostat unit
14. Cephalostat sensor
15. Secondary collimator
16. Positioning panel



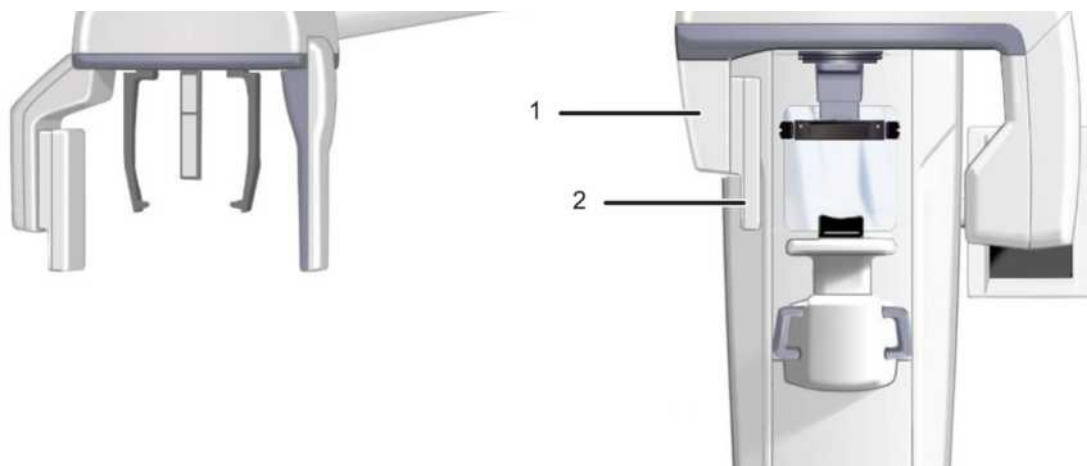
Fig 1.1. On/off switch and main fuses



PC with MDD approved dental imaging software and 3D viewing software (not included).

All software must conform to the MDD and the relevant legal requirements in the USA.

The PC must conform to all the unit and dental imaging software requirements.

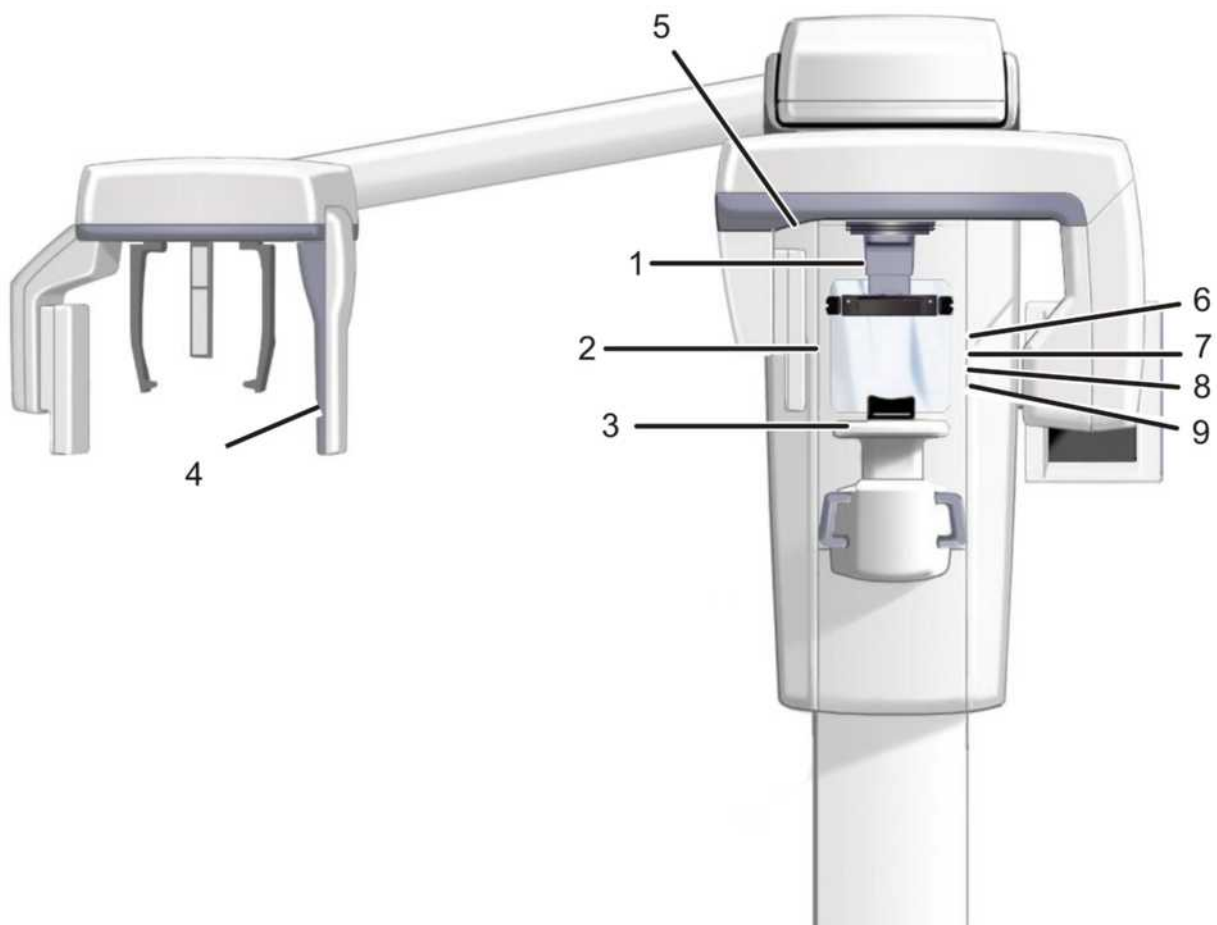


1. Sensor holder (units without 3D option)
2. Panoramic sensor



1. 3D sensor (units with 3D option)
2. Panoramic sensor

2.2 Patient positioning lights



1. Midsagittal light
2. Frankfort horizontal (FH) light /
Horizontal light, top of 130 mm high FOV (3D MFOV
(Maxio) option only)
3. Image layer light
4. Cephalometric FH light
5. TMJ light
6. Horizontal light, top of 78 mm high FOV
(3D MFOV (Maxio) option only)
7. Horizontal light, top of 61 mm high FOV
(3D option only)
8. Horizontal light, top of 50 mm FOV
(3D MFOV (Maxio) option only)
9. Horizontal light, bottom of FOV (3D option only)

Panoramic lights

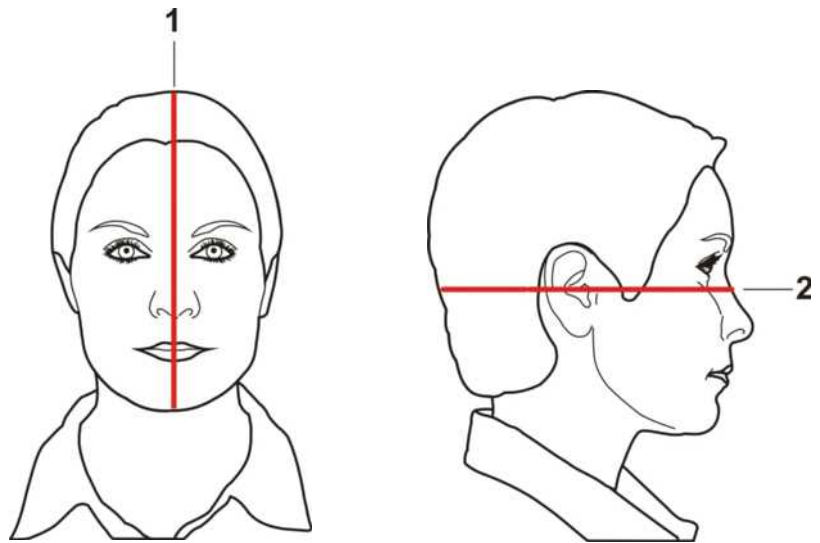


Fig 1.1.

1. Midsagittal light
2. FH light

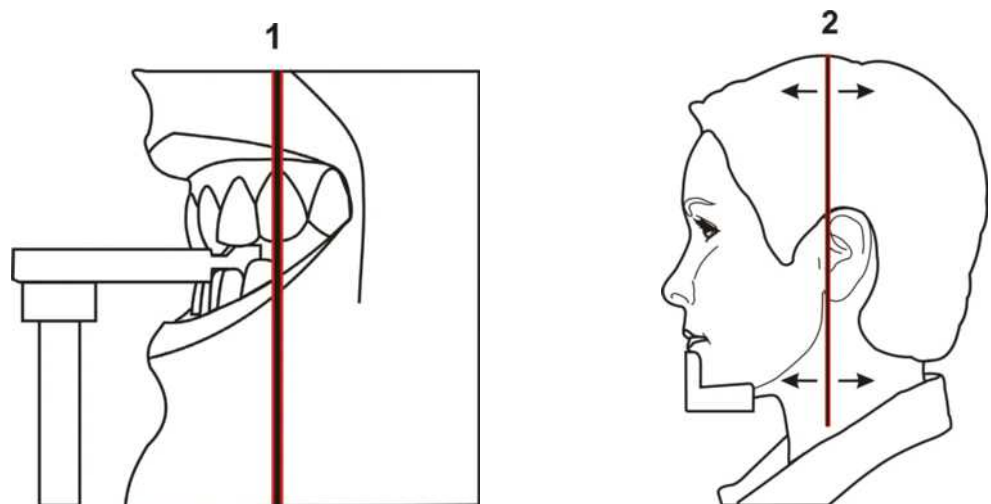


Fig 1.2.

1. Image layer light
2. TMJ light

Cephalometric lights

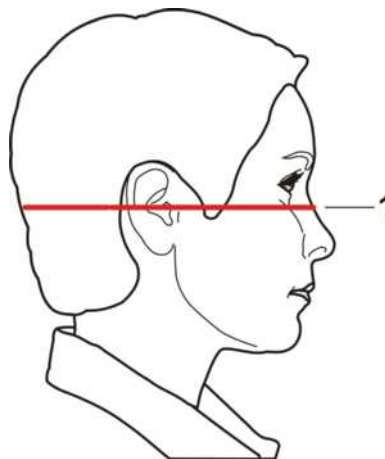


Fig 1.3.

1. FH light

3D lights (optional)

Note! Appropriate lasers are turned automatically on based on selected FOV.

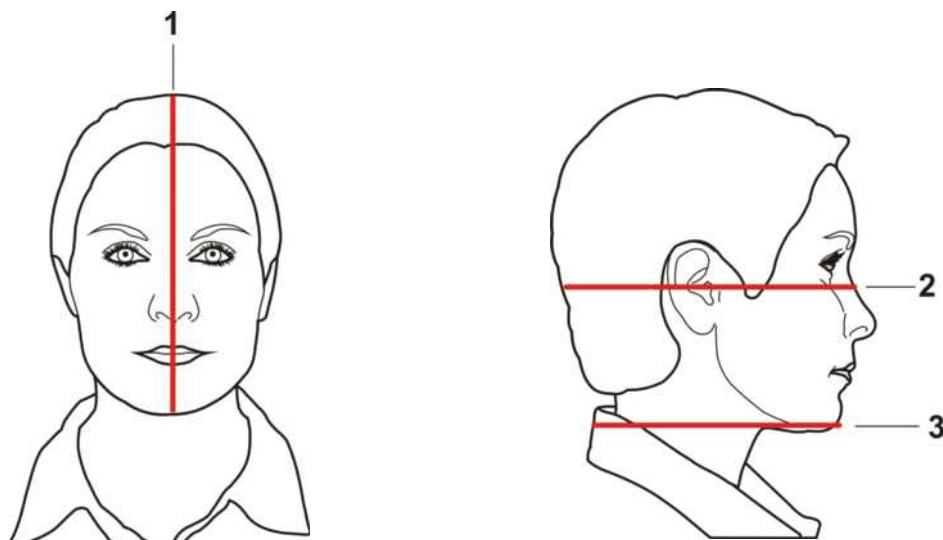


Fig 1.4.

1. Midsagittal light

2. Horizontal light, top of FOV

Note! With 3D MFOV (Maxio) option height 130 mm is indicated with Frankfort horizontal (FH) light. Move FH light to 130 mm position (locked in up-position).

3. Horizontal light, bottom of FOV

2.3 Patient positioning panel

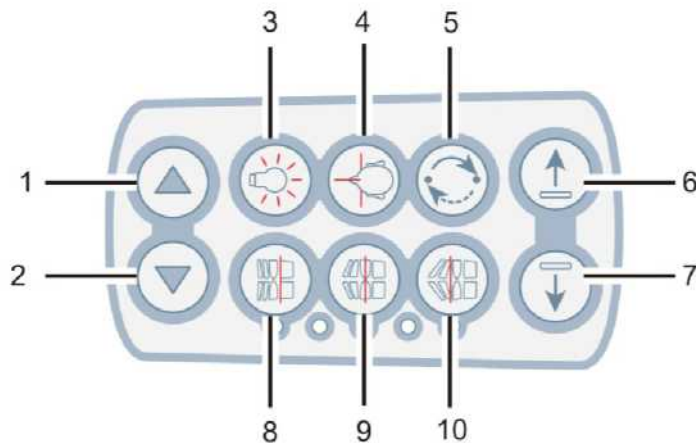
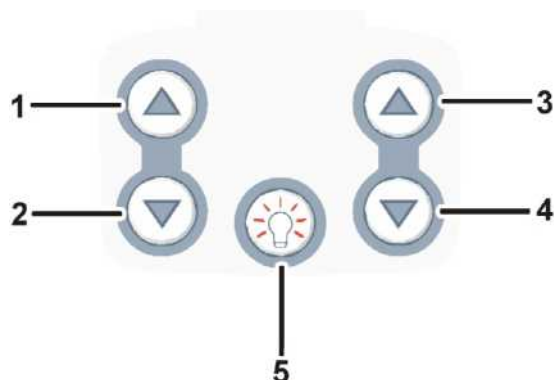


FIG. 4.5

1. Carriage UP
2. Carriage DOWN
3. Positioning lights ON/OFF
4. Patient positioning
5. Start positioning
6. Chin support UP
7. Chin support DOWN
8. Move the image layer anterior before exposure 3 mm, with sinus program 10 mm
9. Normal occlusion/ reset position
10. Move the image layer posterior before exposure 3 mm, with sinus program 10 mm

2.3.1 Cephalometric unit (optional)



1. Carriage UP
2. Carriage DOWN
3. Carriage UP
4. Carriage DOWN
5. Positioning lights ON/OFF

2.4 Emergency stop switch

In case of malfunction of the exposure button or other protective devices of the unit, an emergency stop switch is provided near the handles and on the roof of the cephalostat head so that the patient can reach it.

If the emergency stop switch is pressed during an exposure, the exposure is terminated immediately and the x-ray unit is completely stopped. An interrupted exposure cannot be continued later, but has to be retaken from the beginning.

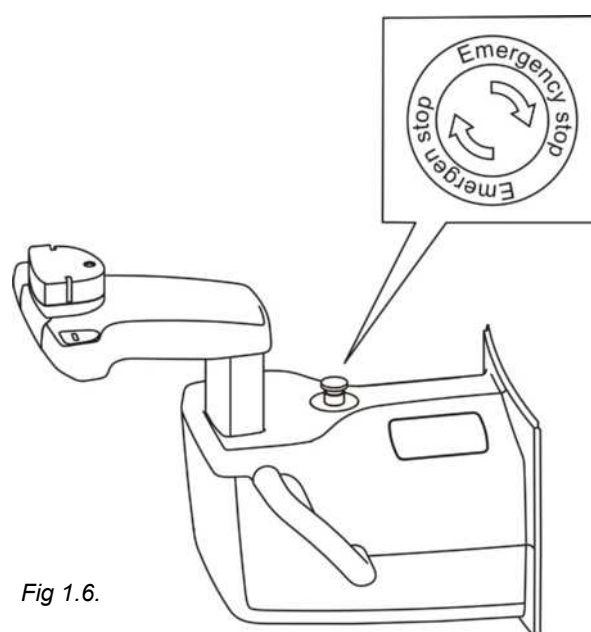


Fig 1.6.

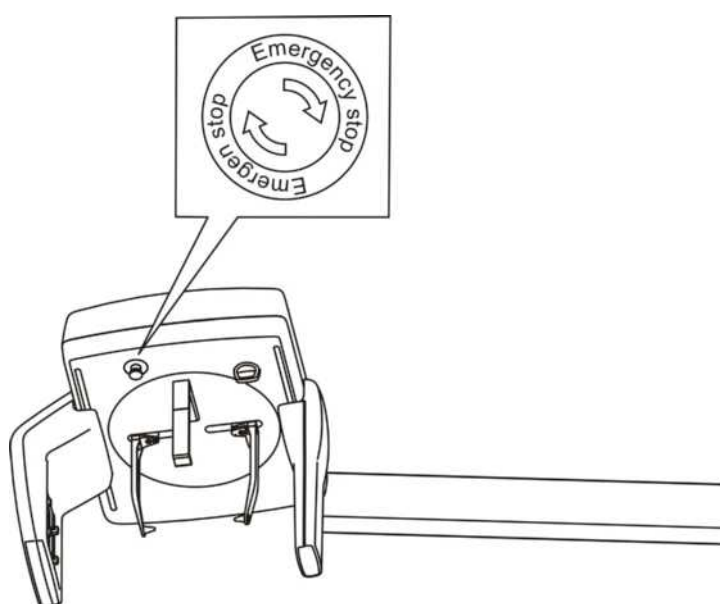


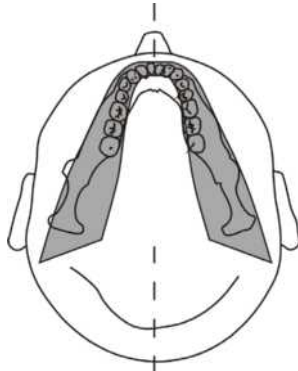
Fig 1.7.

Press to stop the unit, rotate to release.

3 Imaging programs

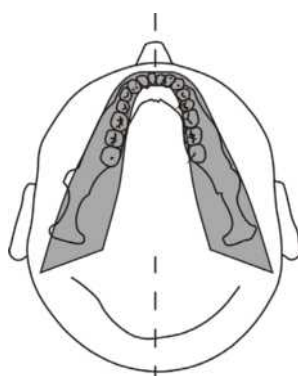
3.1 Panoramic programs

Standard: Magnification 1.3

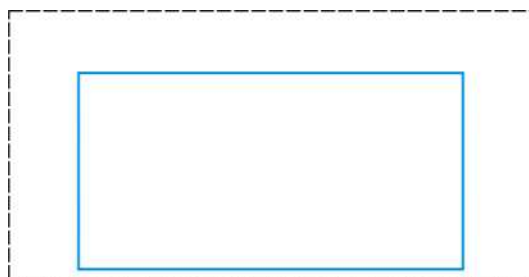


Exposure settings for panoramic program

100 VAC	66 kV/5 mA	66 kV/8 mA	66 kV/10 mA	70 kV/13 mA
230 VAC	66 kV/5 mA	66 kV/8 mA	66 kV/10 mA	70 kV/13 mA
			< default >	



Pediatric: Magnification 1.3

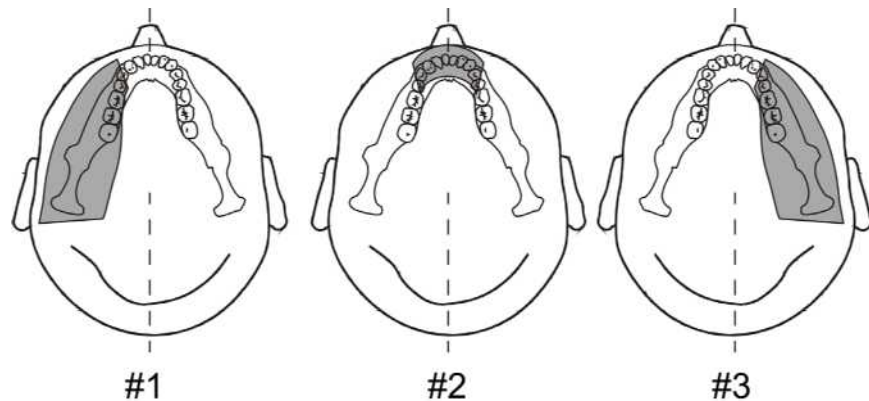


Exposure settings for pediatric program

100 VAC	66 kV/4 mA	66 kV/6.3 mA	66 kV/8 mA	70 kV/10 mA
230 VAC	66 kV/4 mA	66 kV/6.3 mA	66 kV/8 mA	70 kV/10 mA
			< default >	

Pediatric patients can be imaged with less radiation dosage and shorter exposure time. Patients with jaw more narrow than average jaw can be exposed with this procedure too.

Ortho Zone: Magnification 1.25



The Ortho Zone program produces two different scanning geometries combined in the same image.

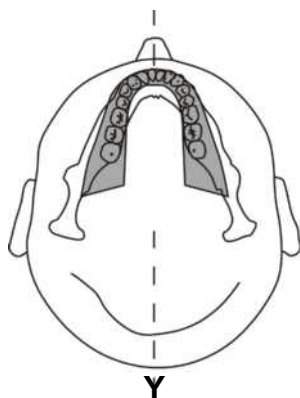
The first geometry (#1 and #3 in the figure) gives a standard panoramic view of the molar region.

The result of this scanning location will allow for views of the TM joint and molar area without redundant shadows from the opposite side ramus obscuring the image. Patients with prosthetic condyles or other posterior radio opaque objects can have the opposite side successfully imaged.

The second view (#2 in the figure) produces an image of the anterior region with a very wide layer of focus (approx. 35 mm). This view may be helpful when diagnosing trauma, wired shut, severe class III malocclusion and uncooperative patients.

Exposure settings for Ortho Zone program

				
100 VAC	66 kV/5 mA	66 kV/8 mA	66 kV/10 mA	70 kV/13 mA
230 VAC	66 kV/5 mA	66 kV/8 mA	66 kV/10 mA	70 kV/13 mA
			< default >	



Orthogonal: Magnification 1.3

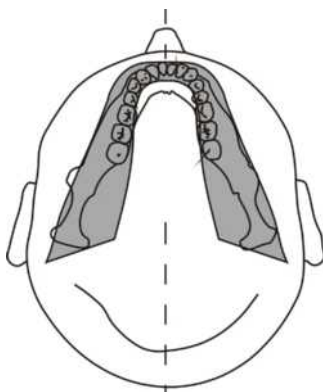
An optimized view of the dentition only with optimized angulation and reduced radiation.



Orthogonal program produces a panoramic view with modified projection geometry. The Y axis of the rotation path is changed to improved the beam angle to be closer to 90° to the interproximal surfaces. With this improvement, other trade off's must be made. The ascending rami may be lost and in adult patients and redundant shadows will be increased.

Exposure settings for Orthogonal program

				
100 VAC	66 kV/5 mA	66 kV/8 mA	66 kV/10 mA	70 kV/13 mA
230 VAC	66 kV/5 mA	66 kV/8 mA	66 kV/10 mA	70 kV/13 mA
			< default >	



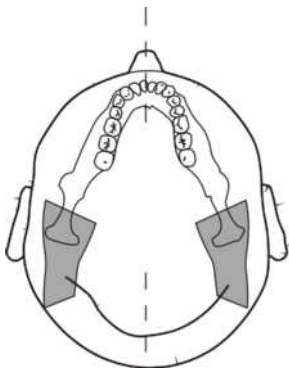
Wide arch: Magnification 1.3

Used when the patient has a wider than normal dental arch.

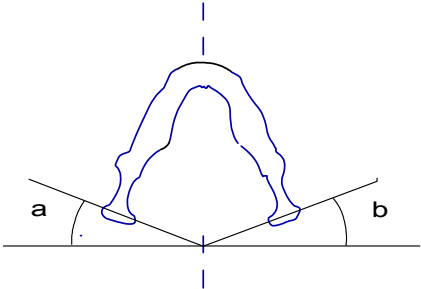


Exposure settings for Wide arch program

				
100 VAC	66 kV/5 mA	66 kV/8 mA	66 kV/10 mA	70 kV/13 mA
230 VAC	66 kV/5 mA	66 kV/8 mA	66 kV/10 mA	70 kV/13 mA
			< default >	

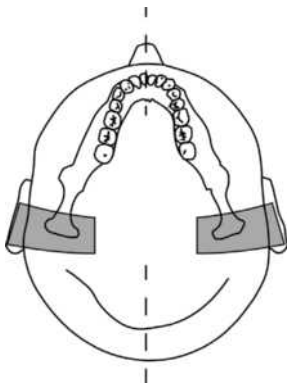


Ortho Lateral TMJ: Magnification 1.23



Ortho TMJ program provides a wide layer axially corrected views for the patient's left and right temporomandibular joints.

Exposure settings for Ortho Lateral TMJ program				
				
100 VAC	73 kV/6.3 mA	73 kV/10 mA	73 kV/13 mA	73 kV/16 mA
230 VAC	73 kV/6.3 mA	73 kV/10 mA	73 kV/13 mA	73 kV/16 mA
			< default >	

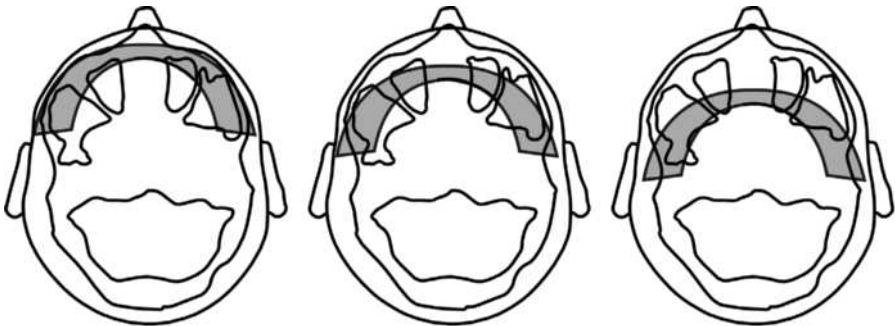


PA TMJ: Magnification 1.55



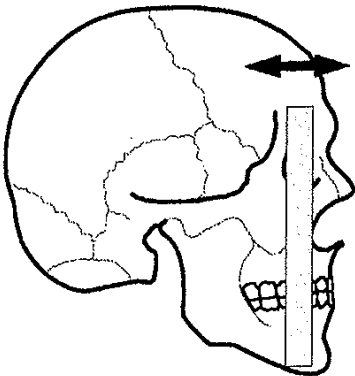
Exposure settings for PA TMJ program				
				
100 VAC	73 kV/6.3 mA	73 kV/10 mA	73 kV/13 mA	73 kV/16 mA
230 VAC	73 kV/6.3 mA	73 kV/10 mA	73 kV/13 mA	73 kV/16 mA
			< default >	

Maxillary Sinus: Magnification 1.3

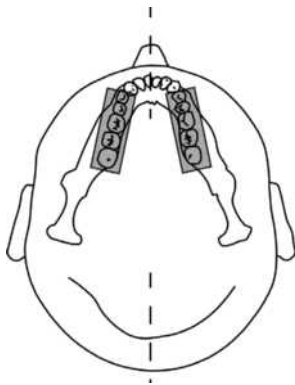


Mesial 10 mm Start Distal 10 mm

Maxillary Sinus program produces a pan - tomographic layer through the posterior maxillary sinus. The layer is flatter than the standard panoramic programs and is moved 18 mm backward. These images are helpful in visualizing the mid and posterior maxillary sinus.



Exposure settings for Maxillary Sinus program				
100 VAC	66 kV/6.3 mA	66 kV/10 mA	66 kV/13 mA	73 kV/13 mA
230 VAC	66 kV/6.3 mA	66 kV/10 mA	66 kV/13 mA	73 kV/13 mA
			< default >	



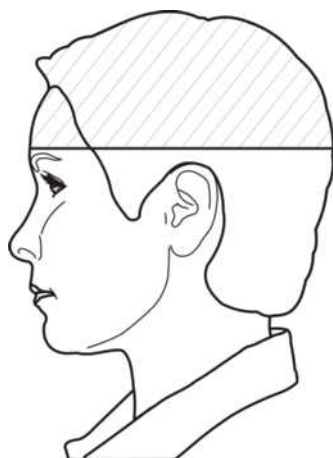
Bitewing: Magnification 1.3

An orthogonal view of the dentition from the canine and posterior.



Exposure settings for Bitewing program				
				
100 VAC	66 kV/5 mA	66 kV/8 mA	66 kV/10 mA	70 kV/13 mA
230 VAC	66 kV/5 mA	66 kV/8 mA	66 kV/10 mA	70 kV/13 mA
			< default >	

3.2 Cephalometric programs

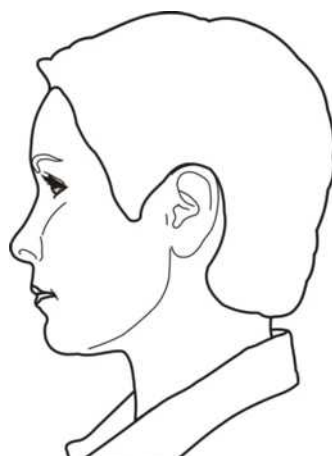


Cephalometric pediatric lateral projection

Pediatric Lateral Cephalostat has an optimized image height (180 mm) that is used e.g. for pediatric patients but also adult patients to reduce the radiation dose. The pediatric lateral projection covers all the typical cephalostat landmarks from Nasion down to the spine and the starting point of the lateral scan is adjustable with both Standard and pediatric lateral cephalostat programs.

Exposure settings for cephalometric pediatric lateral program

100 VAC	90 kV/8 mA/10 s	90 kV/8 mA/13 s	90 kV/8 mA/16 s	90 kV/8 mA/20 s
120 VAC	85 kV/10 mA/10 s	85 kV/10 mA/13 s	90 kV/8 mA/16 s	90 kV/8 mA/20 s
230 VAC	85 kV/10 mA/13 s	90 kV/10 mA/13 s	90 kV/13 mA/16 s	90 kV/13 mA/20 s
			< default >	



Cephalometric lateral projection

Lateral Cephalostat uses a full height image field (223 mm).



Exposure settings for cephalometric lateral program

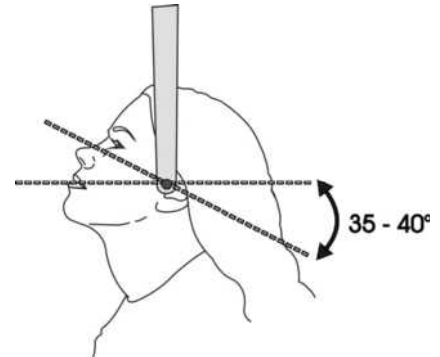
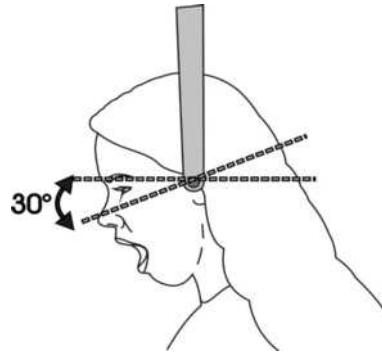
100 VAC	90 kV/8 mA/10 s	90 kV/8 mA/13 s	90 kV/8 mA/16 s	90 kV/8 mA/20 s
120 VAC	85 kV/10 mA/10 s	85 kV/10 mA/13 s	90 kV/10 mA/16 s	90 kV/10 mA/20 s
230 VAC	85 kV/10 mA/13 s	90 kV/10 mA/13 s	90 kV/13 mA/16 s	90 kV/13 mA/20 s
			< default >	

Cephalo posterior-anterior (PA) projection



Reverse towne projection

Waters view



Exposure settings for cephalometric PA program

100 VAC	90 kV/8 mA/10 s	90 kV/8 mA/13 s	90 kV/8 mA/16 s	90 kV/8 mA/20 s
120 VAC	85 kV/10 mA/ 10 s	85 kV/10 mA/ 13 s	90 kV/10 mA/ 16 s	90 kV/10 mA/ 20 s
230 VAC	85 kV/10 mA/ 13 s	90 kV/10 mA/ 13 s	90 kV/13 mA/ 16 s	90 kV/13 mA/ 20 s
			< default >	



Carpus view (Not available in USA and Canada)

Exposure settings for Carpus view program

100 VAC	66 kV/3,2 mA/ 8 s	70 kV/3,2 mA/ 8 s	73 kV/3,2 mA/ 8 s	73 kV/6,3 mA/ 8 s
120 VAC				
230 VAC				
			< default >	

3.3 3D SFOV programs

61 x 41 mm FOV



High resolution (133 μ m voxel size)



Standard resolution (200 μ m voxel size)

Program for optimized endodontic imaging:



Endo program (85 μ m voxel size)

61 x 78 mm FOV



High resolution (200 μ m voxel size)



Standard resolution (300 μ m voxel size)

Exposure settings for 3D SFOV imaging (default values)				
Resolution	FOV (h x w)	kV	mA	Exposure time
Endo program	61 x 41 mm	90	10	6,1 s
High Res	61 x 41 mm	90	8	6,1 s
Std Res	61 x 41 mm	90	10	2,3 s
High Res	61 x 78 mm	90	6,3	13 s
Std Res	61 x 78 mm	90	10	4,9

3.4 3D MFOV (Maxio) programs

50 x 50 mm FOV



High resolution (125µm voxel size)



Standard resolution (200µm voxel size)



Low Dose Scan (280µm voxel size)

Program optimized for endodontic imaging:



Endo program (85µm voxel size)

61 x 78 mm FOV



High resolution (200µm voxel size)



Standard resolution (300µm voxel size)



Low Dose Scan (320µm voxel size)

78 x 78 mm FOV

High resolution (200µm voxel size)



Standard resolution (300µm voxel size)



Low Dose Scan (320µm voxel size)

78 x 150 mm FOV

High resolution (250µm voxel size)



Standard resolution (350µm voxel size)



Low Dose Scan (400µm voxel size)

130 x 150 mm FOV

High resolution (320µm voxel size)



Standard resolution (380µm voxel size)



Low Dose Scan (420µm voxel size)

3.5 Selecting resolution and FOV

The resolution selection has an effect to the image quality and to the patient dose. For example High resolution will give better image quality than Standard resolution, but on the other hand the patient dosage is also higher. The unit offers a Low Dose resolution (see Tables 1.1 & 1.4) which can be used for example in treatment follow-up cases. The Low Dose resolution will result in images of reduced image quality (in other words image quality is proportional to dose) and it is up to the dental professional to decide when it is sufficient to use this mode. As small as possible FOV size should be selected for the patient case in order to follow the ALARA (As Low As Reasonably Achievable) principle.

Table 1.1 General guidelines for selecting 3D resolution. It is always up to the dental professional to select the appropriate mode.

Resolution setting	General recommendations for the use
Low Dose Resolution	Treatment follow up, children
Standard Resolution	Implants, 3 rd molars, TMJ, impacted teeth, resorptions
High Resolution	Pathologies, alveolar bone defects, root fractures
Endo Resolution	Endodontic cases (periapical infections, root canals, fractures, etc.)

Table 1.2 General guidelines for selecting 3D FOV. It is always up to the dental professional to select the appropriate mode.

Resolution setting	General recommendations for the use
SFOV 61 x 41 mm MFOV 50 x 50 mm	Optimized for single site implants or localized diagnostics, for example 3 rd molar extractions, impacted teeth's, single TMJ analysis, endodontics, and children.
SFOV 61 x 78 mm MFOV	Multiple implant placement using surgical guides, covers complete dental arch, optimized for one jaw.
MFOV 78 x 78 mm	Entire dentition, both mandibula and maxilla as well as a portion of maxillary sinus.
MFOV 78 x 150 mm	Both mandibula and maxilla including airway and upper cervical spine or the sinus, both TM joints.
MFOV 130 x 150 mm	Covers entire maxillofacial region, from maxilla to frontal sinus or from mandibula to maxillary sinus.

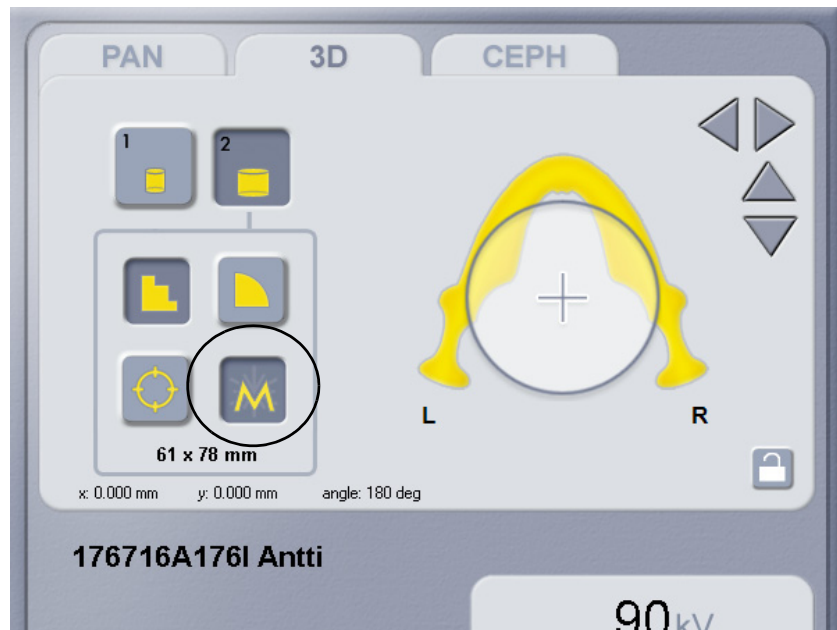
•

3.6 MAR, Metal Artifact Reduction

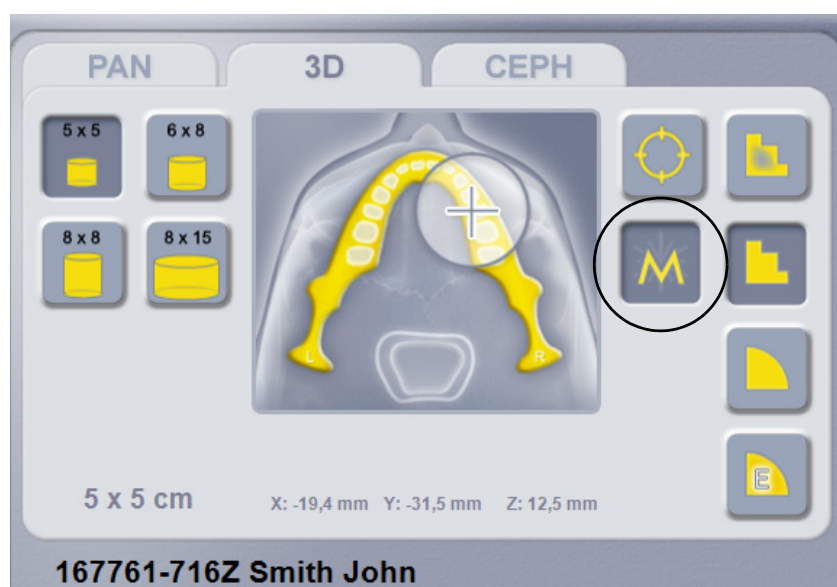


MAR-button is OFF.

MAR, Metal Artifact Reduction software can be used to reduce the effect of metals and other dense radiopaque objects on the 3D image. These create artifacts that are seen typically as stripes and shadows from the above-mentioned objects. To utilize MAR on a 3D image may have affect to image reconstruction time



3D SFOV touch screen: MAR-button is ON.
MAR-button becomes visible on the 3D modality.



MFOV (Maxio) touch screen: MAR-button is ON.
MAR-button becomes visible on the 3D modality.

3.7 Exposure settings for 3D imaging

Table 1.3

Exposure settings for 3D SFOV imaging							
Note! Voltage settings is always 90kV with the unit 3D modality.							
Resolution	FOV (h x w)	Exposure time	Scanning time	Amount of projections	mA Low dose (DAP mGycm ²)	mA default (DAP mGycm ²)	mA High-quality (DAP mGycm ²)
High Res	61 x 41 mm	6,1 s	10 s	609	6.3 mA (240)	8 mA (385)	13 mA (601)
Std Res	61 x 41 mm	2,3 s	10 s	234	8 mA (148)	10 mA (184)	13 mA (231)
Endo Res	61 x 41 mm	6,1 s	10 s	609	8 mA (385)	10 mA (476)	13 mA (601)
High Res	61 x 78 mm	12,6 s	20 s	1262	5 mA (498)	6.3 mA (619)	10 mA (996)
Std Res	61 x 78 mm	4,9 s	20 s	486	8 mA (306)	10 mA (372)	13 mA (479)

Table 1.4

Exposure settings for 3D imaging, MFOV (Maxio)								
Note! Voltage settings is always 90kV with the unit 3D modality.								
3D program	FOV (h x w)	Exposure time	Project- ions	Scanning time	Default mA	mAs	DAP (mGycm ²)	Voxel (um)
Low Dose	50 x 50 mm	1,17 s	234	10,96 s	3,2 mA	3,7	32	280
Std Res	50 x 50 mm	2,34 s	234	10,96 s	8 mA	18,7	162	200
High Res	50 x 50 mm	6,09 s	609	17,4 s	6,3 mA	38,4	332	125
Endo Res	50 x 50 mm	8,7 s	870	17,4 s	5 mA	43,5	377	85
Low Dose	61 x 78 mm	1,17 s	234	10,96 s	3,2 mA	3,7	58	320
Std Res	61 x 78 mm	2,34 s	234	10,96 s	8 mA	18,7	288	300
High Res	61 x 78 mm	6,09 s	609	17,4 s	6,3 mA	38,4	591	200
Low Dose	78 x 78 mm	1,17 s	234	10,96 s	3,2 mA	3,7	72	320
Std Res	78 x 78 mm	2,34 s	234	10,96 s	8 mA	18,7	358	300
High Res	78 x 78 mm	6,09 s	609	17,4 s	6,3 mA	38,4	735	200
Low Dose	78 x 150 mm	2,25 s	450	19,5 s	3,2 mA	7,2	138	400
Std Res	78 x 150 mm	4,5 s	450	19,5 s	6.3 mA	28,4	543	350
High Res	78 x 150 mm	8,5 s	850	24,3 s	5 mA	42,5	814	250
Low Dose	130 x 150 mm	4,5 s	900	40 s	6.3 mA	14,4	276	420
Std Res	130 x 150 mm	9,0 s	900	40 s	4 mA	36,0	690	380
High Res	130 x 150 mm	9,0 s	900	40 s	6,3 mA	56,7	1086	320

DAP values vary from unit to unit in relation to the x-ray tube output. Thus above values indicate average DAP values. In addition to these recommended values, there is a possibility to use the whole mA range if the user prefers.

Available mA ranges for each field of view sizes and resolution settings (3D SFOV)							
Resolution	FOV (h x w)	4 mA	5 mA	6.3 mA	8 mA	10 mA	13 mA
Endo Res	61 x 41 mm	x	x	x	x	x	x
High Res	61 x 41 mm	x	x	x	x	x	x
Std Res	61 x 41 mm			x	x	x	x
High Res	61 x 78 mm	x	x	x	x	x	
Std Res	61 x 78 mm			x	x	x	x

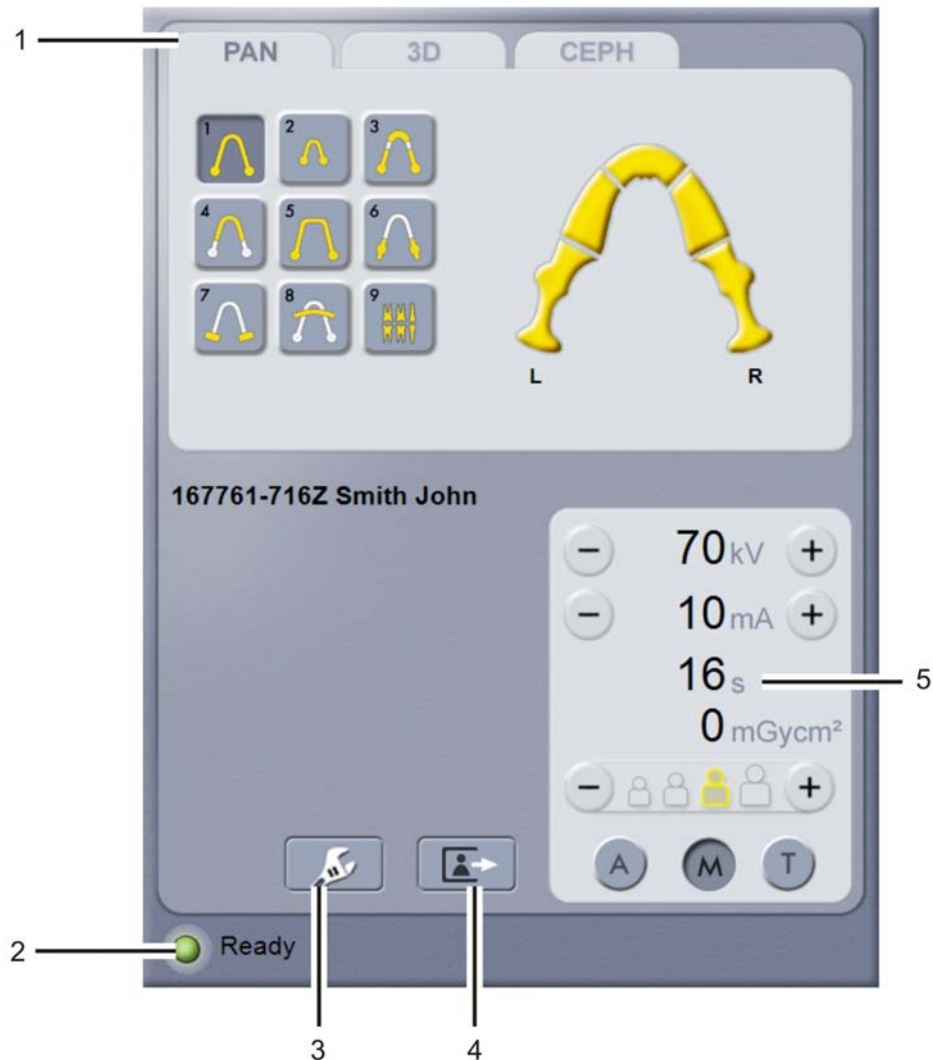
Available mA ranges for each field of view sizes and resolution settings (MFOV (Maxio))								
Resolution	FOV (h x w)	3.2 mA	4 mA	5 mA	6.3 mA	8 mA	10 mA	12,5 mA
Low Dose	50 x 50 mm	x	x	x	x			
Std Res	50 x 50 mm			x	x	x	x	x
High Res	50 x 50 mm		x	x	x	x	x	x
Endo Res	50 x 50 mm		x	x	x	x	x	x
Low Dose	61 x 78 mm	x	x	x				
Std Res	61 x 78 mm			x	x	x	x	x
High Res	61 x 78 mm		x	x	x	x	x	
Low Dose	78 x 78 mm	x						
Std Res	78 x 78 mm			x	x	x	x	x
High Res	78 x 78 mm		x	x	x	x	x	
Low Dose	78 x 150 mm	x						
Std Res	78 x 150 mm		x	x	x	x	x	
High Res	78 x 150 mm	x	x	x	x	x		
Low Dose	130 x 150 mm	x						
Std Res	130 x 150 mm		x	x	x	x	x	
High Res	130 x 150 mm	x	x	x	x	x		

Exposure settings for scout imaging (3D SFOV, default values)				
Resolution	FOV (h x w)	kV	mA	Scanning time
Scout	61 x 41 mm	90	13	0,02 s
Scout	61 x 78 mm	90	13	0,04 s

Exposure settings for scout imaging (MFOV (Maxio), default values)				
Resolution	FOV (h x w)	kV	mA	Scanning time
Scout	50 x 50 mm	90	13	0,02 s
Scout	61 x 78 mm	90	13	0,02 s
Scout	78 x 78 mm	90	13	0,02 s
Scout	78 x 150 mm	90	13	0,04 s
Scout	130 x 150 mm	90	13	0,08 s

4 Touch screen display

4.1 Main control panel



1. Modality / imaging program section
2. Status of the unit
3. Settings
4. End examination
5. Exposure settings

4.2 Modality section

Select the modality tab PAN, CEPH or 3D.

When panoramic modality is selected, a program specific dental arch is shown. This can be used for partial panoramic imaging.

Cephalometric programs have their own, program specific model heads and setting buttons for the start position of lateral scanning.

OP300 3D SFOV (two FOV sizes) has buttons for selecting standard resolution, high resolution and scout image mode.

OP300 MFOV (Maxio) (five FOV sizes) has buttons for selecting Low DoseMinidose, standard or high resolution and scout image mode.

The FOV for 3D imaging can be positioned on the XY-plane by selecting the center point of the FOV on the dental arch of the touch screen display. The FOV is positioned in the Z-direction by using the chin rest movement and positioning lights.

4.2.1 Exposure indicators and settings



kV value



mA value



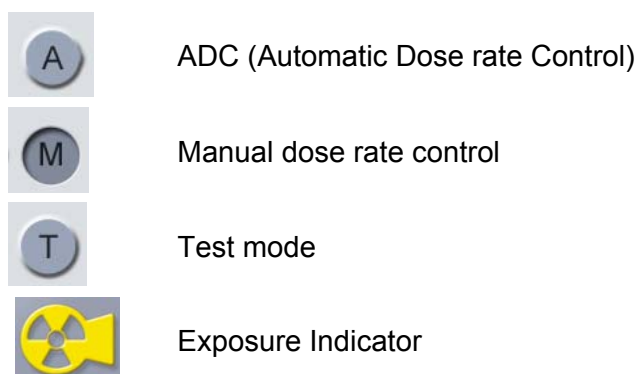
Exposure time



Dose value



Patient size settings
(Child, juvenile,
small adult, large adult)



4.3 Automatic dose control (ADC)

With ORTHOPANTOMOGRAPH® OP300 it is possible to take panoramic exposure with Automatic Dose Control (P1 through P5).

The software will monitor the amount of radiation the CMOS sensor is receiving and automatically set the exposure factors for proper dose. After the exposure the adjusted values are shown on the display. The ADC will stay engaged with all the panoramic procedures (P1 through P5) unless set to manual mode. Any of sectional panoramic programs cannot utilize ADC.

The signal to noise ratio can be adjusted while keeping ADC engaged. Adjustment is done from the GUI.

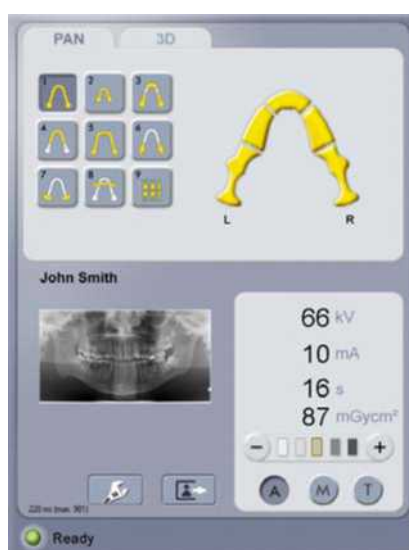


Fig 1.8. ADC signal to noise ratio scale

4.4 Status section

Status field shows when the unit is ready for capturing or when any trouble occurs. Green, yellow and blue color indicate the status in question.

4.5 Other sections



General settings



- **Retrieve Last Image**
- Use this to retrieve the last image from the device memory e.g. after a system error
- **Retrieve With MAR**
- After taking a with MAR either on or off, image can be re-reconstructed with different setting by selecting either “Retrieve last image” or “Retrieve With MAR” from the general settings on the touch screen. If reconstruction with MAR is unavailable for the last captured image, e.g. last taken image is cephalometric, “Retrieve With MAR”-button will not be visible.

NOTE! Only the last taken x-ray image is saved in the unit until the power is switched off. This image data is used in the retrieve procedure.

- Languages
 - Use this to select language on the touchscreen
- Service
 - Use this to reach the programs for periodical maintenance

5 Using the unit

5.1 Attaching and removing the sensor

Note! The pixel calibration results are sensor specific. If the x-ray unit is equipped with separate panoramic and cephalometric sensors, the cephalometric sensor cannot be used for panoramic imaging without re-calibration (and vice versa).

Re-do panoramic pixel calibration, if cephalostat sensor is moved to panoramic side or the sensor is changed.

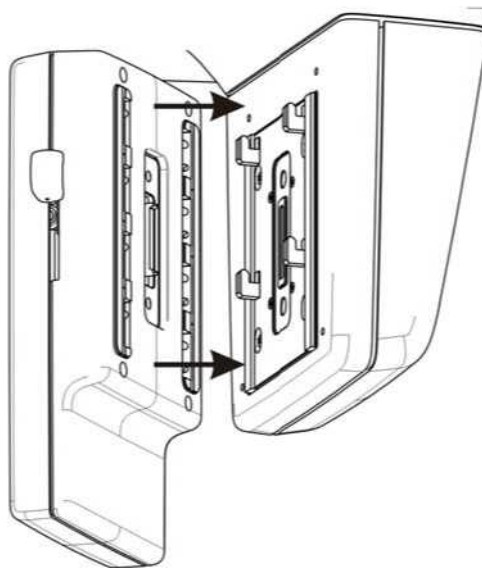


WARNING!

Handle the sensor with care as instructed in this manual. The sensor must not be dropped or exposed to impacts. A shock indicator inside the sensor shows if the sensor has been exposed to excess impact.

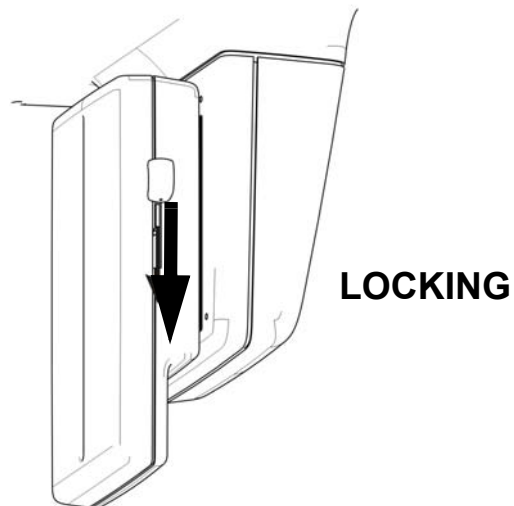
5.1.1 Attaching the sensor

1. Insert the four slots on the rear of the sensor, into the four hooks in the sensor holder.



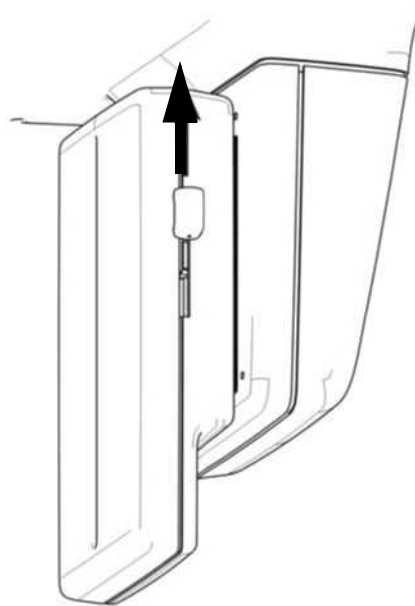
2. Pull the sensor downwards firmly until it stops and then slide the locking knob down on the side of the sensor to lock the sensor in position.

Note! Make sure that the sensor is seated properly before sliding the locking knob down. Forcing the locking knob down when the sensor is not in correct position may damage the sensor connectors!



5.1.2 Removing the sensor

1. Slide the locking knob upwards on the side of the sensor to unlock the sensor.



2. Slide the sensor up and remove it.

5.2 Preparing the system

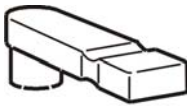
1. Switch on the unit and the PC.
2. **PC:** Start CLINIVIEW™ software (or 3rd party application).
3. **PC:** Open a new or existing patient or select a patient from the worklist. See the user's guide supplied with the dental imaging program.

5.3 Panoramic exposures

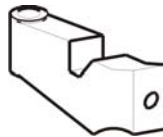
- Standard
- Pediatric
- Ortho Zone
- Orthogonal
- Wide arch
- Bitewing
- Ortho TMJ axially corrected lateral projection
- TMJ PA projection
- Maxillary sinus view

5.3.1 Positioning devices

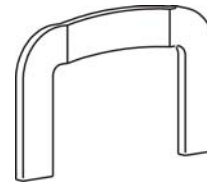
**Bite fork with
the bite block**



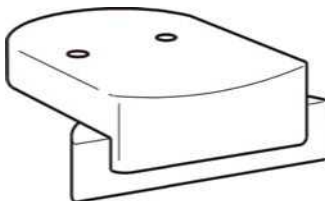
**Bite fork with
the edentulous bite
positioner**



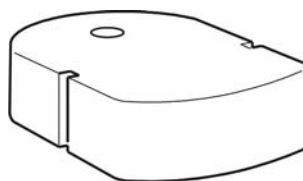
**Chin support
for edentulous
patients**



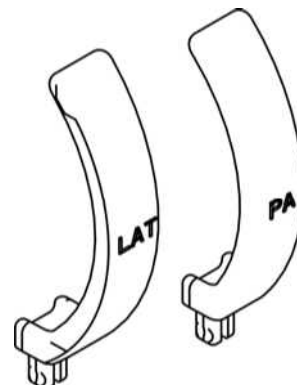
Sinus rest



Chin rest



TMJ nose support





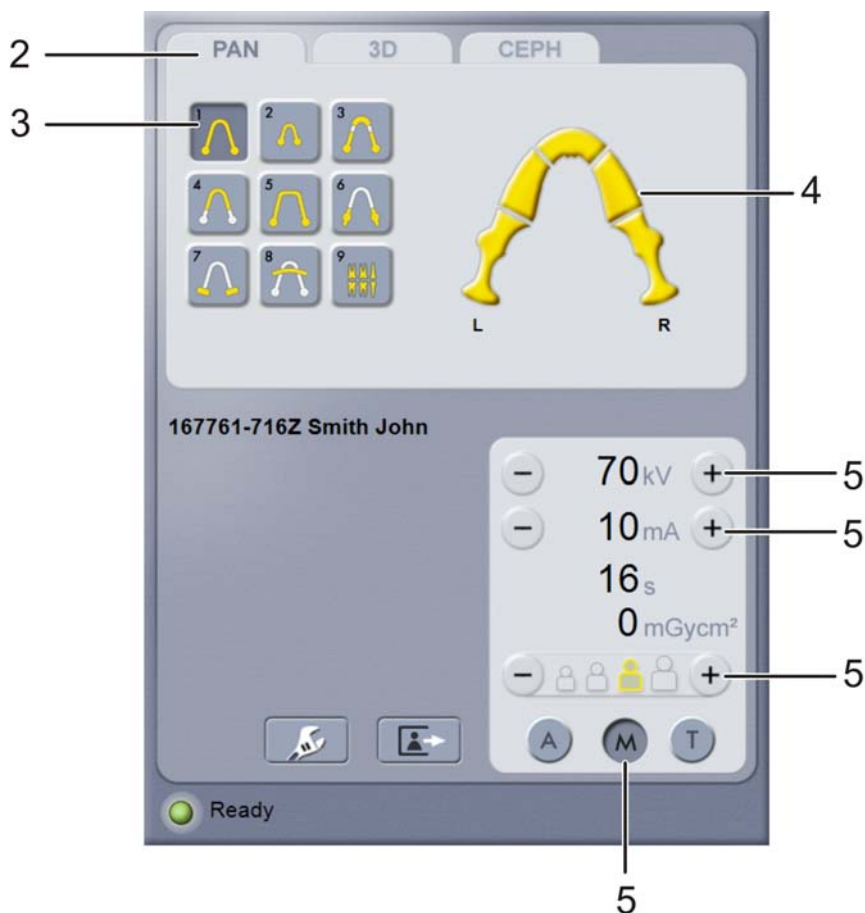
5.3.2 Sectional imaging

Dental arch on the touch panel shows the enabled and disabled arch sections from the result point of view. Select the image area from the dental arch.

5.3.3 General instructions



1. **PC:** Click **Image Capture**



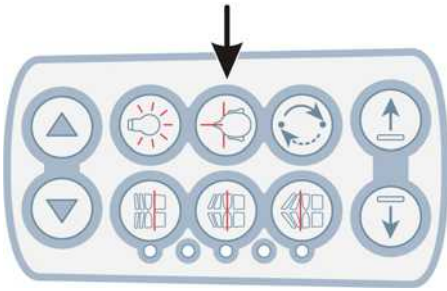
2. Select PAN tab.

3. Select the imaging program.

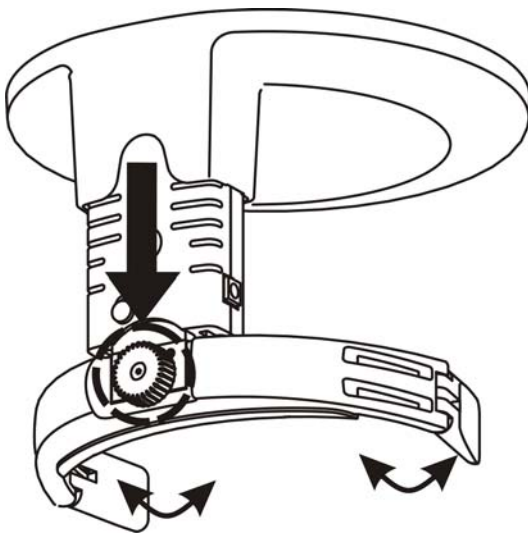
4. Any section of the tooth arch can be selected for the partial panoramic imaging to reduce the radiation.

5. Select the Manual mode.
Set the kV and mA, or select the patient size (child, juvenile, adult, large adult).

6. Press the patient positioning button to rotate the unit to 'patient in' position.



7. Open the temple supports.



8. Ask the patient to remove any spectacles, hearing aids, removable dentures, jewellery and hair clips and pins. Place a protective lead apron on the patient.

Note! Local country regulations may set different standard for lead apron usage needs.

5.3.4 Default exposure settings

User Configurable Panoramic mA Level

Default mA level for panoramic programs can be set using the touch screen display. To change the setting:

1. Start an exam and select a panoramic program.



2. Press **Settings** on the touch screen.

Imaging program defaults

3. Select *Imaging program defaults*.

Set current PAN mA as default

4. Select *Set current PAN mA level* as default.

The selected mA value is used as default value for the current program. The default mA level is adjusted by an equal amount also for other panoramic programs.

5.3.5 User Configurable Default Program

Default imaging program (pan/ceph/3D) can be set using the touch screen display. To change default program:

1. Start an exam and select the desired program.
2. Press the Settings button on the touch screen.



3. Select Imaging program defaults.

Imaging program defaults

4. Select Set current program as default.

Set current program as default

The selected program is automatically activated when an exam is started for a new patient and at startup.

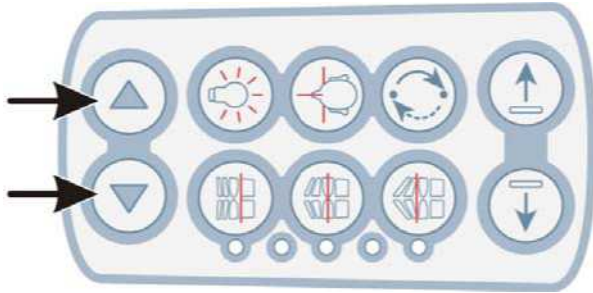
5.3.6 Patient positioning

5.3.6.1 Panoramic exposure

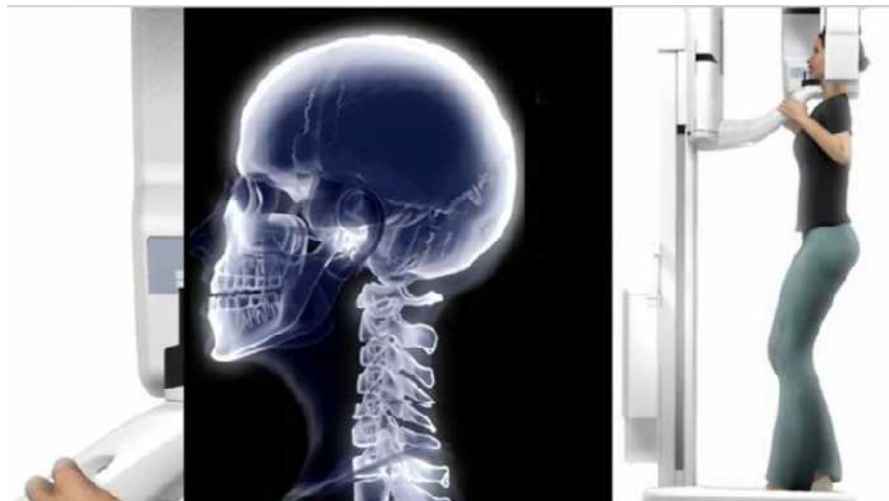
1. Insert the sinus rest, chin rest and bite fork with the bite block. Place the disposable covers.

Note! Use a new disposable cover for every patient.

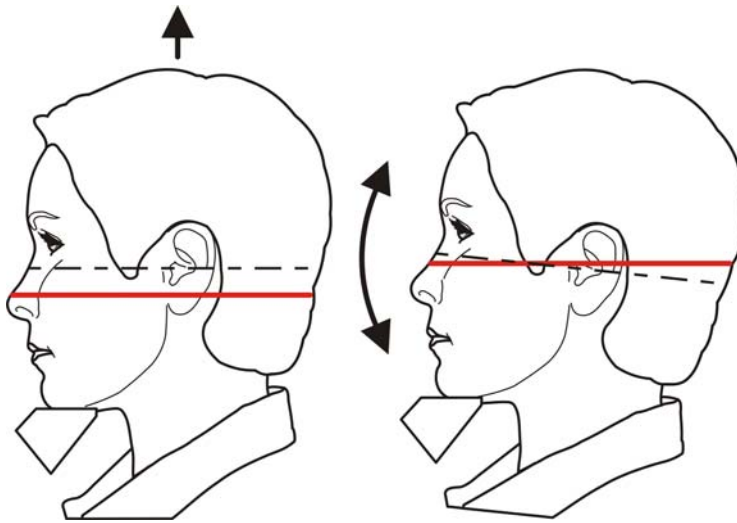
2. Adjust the unit height



3. Guide the patient to the unit and instruct to stand as straight and tall as possible. Exposure can be taken also in sitting position. Ask the patient to take grip on the handles and bite on the bite block. Use the edentulous bite positioner or the chin support for an edentulous patient.
4. Ask the patient to take one step forward to straighten the spinal column. Patient is slightly leaning backwards during the imaging.

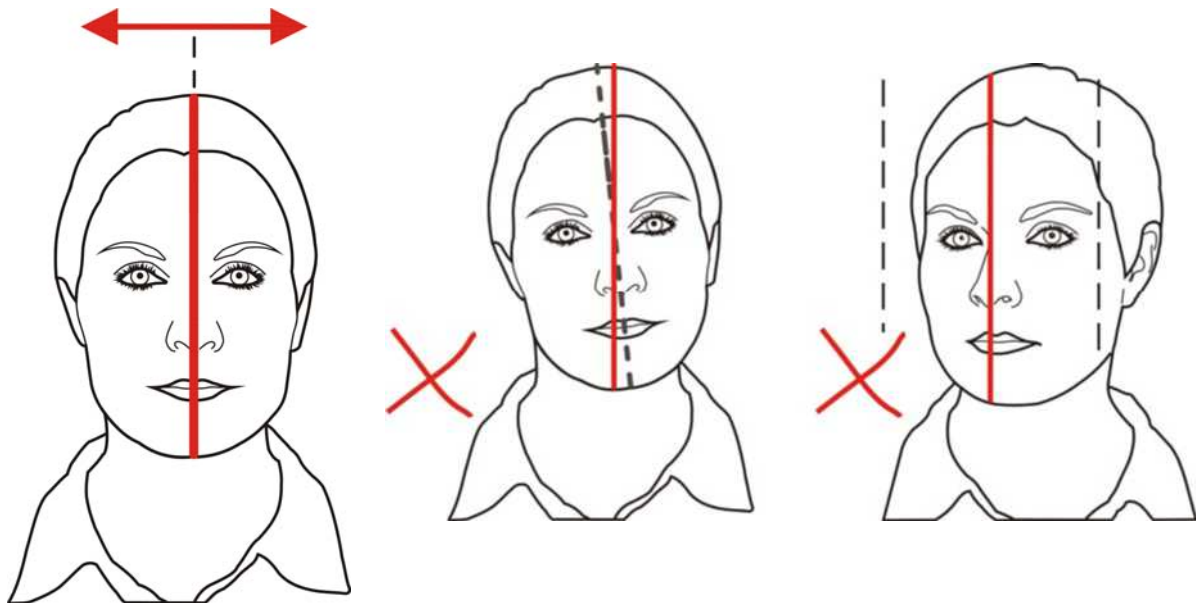


5. Adjust the height of the Frankfort-Horizontal plane (FH) laser to get the laser light over the orbita porion. Straighten the patient's head if needed.

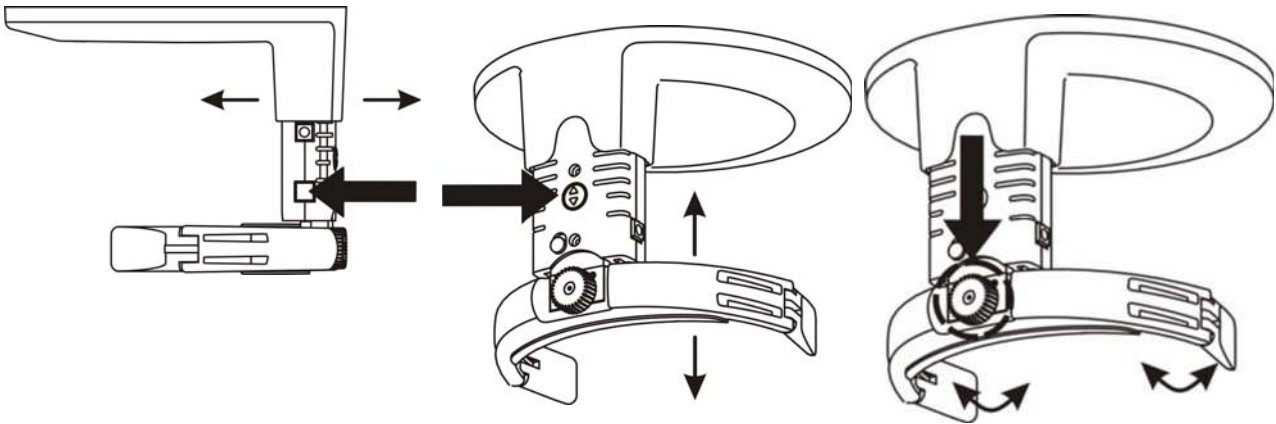


6. Check the position of the midsagittal light. If it is not on the midsagittal plane of the patient, adjust the patient's head.

Make sure the patient's head is not turned or tilted.



7. Move the head support against the patient's forehead. Adjust the height. Close the temple supports.



8. Check the position of the image layer light. If it is not on middle of the maxillary canine (or base of the nose, if edentulous), adjust the image layer.

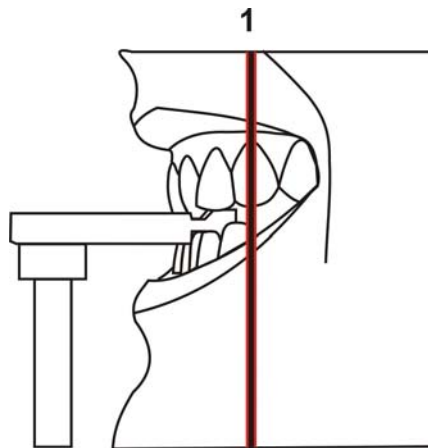
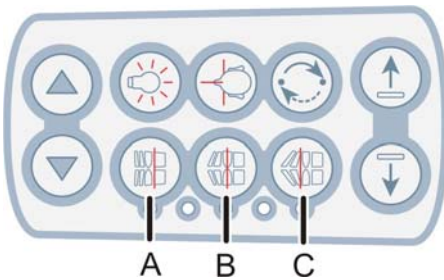


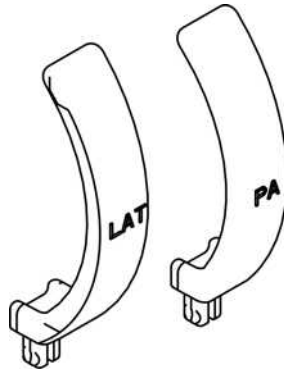
Image layer adjustment buttons:

- A)** Retrusion, 10 mm anterior
- B)** Normal occlusion (default), center
- C)** Protrusion, 10 mm posterior

9. Ask the patient to press their tongue against the roof of their mouth, **swallow** and remain still for the duration of the exposure.

5.3.6.2 TMJ exposure

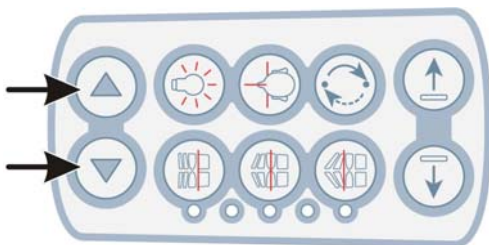
Nose support



1. Remove the chin rest. Insert the required positioning devices, including the TMJ nose support. Place the disposable covers.

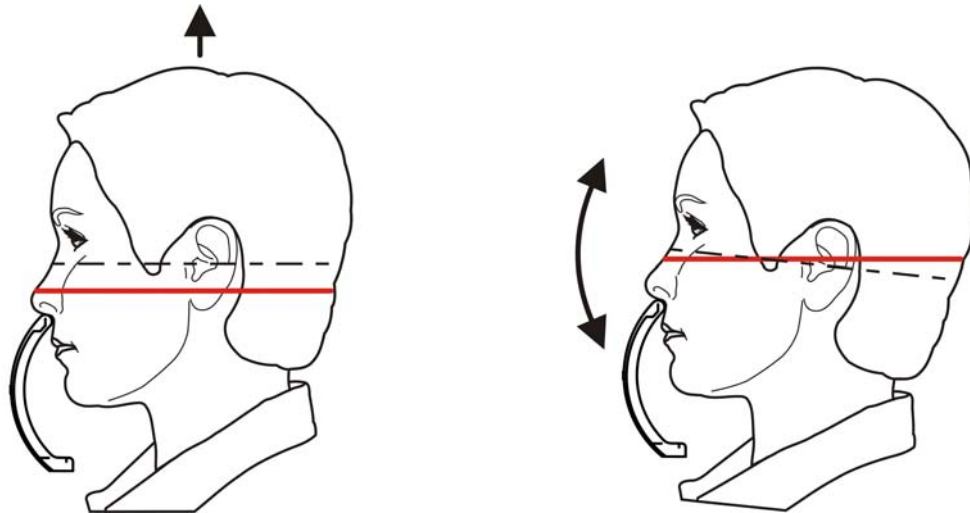
Note! Use a new disposable cover for every patient.

2. Adjust the unit height

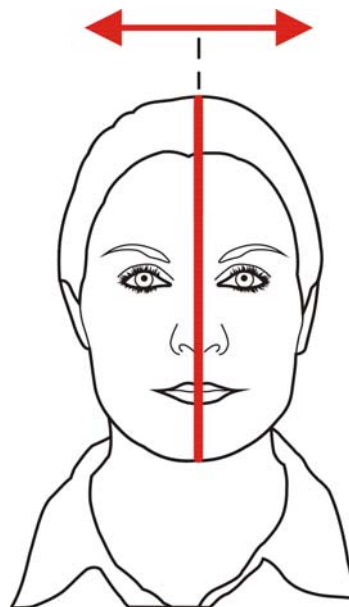


3. Guide the patient to the unit and instruct to stand as straight and tall as possible. Ask the patient to take grip on the handles and set the nose against the TMJ nose support.

4. Adjust the height of the Frankfort-Horizontal plane (FH) laser to get the laser light over the orbita porion. Straighten the patient's head if needed.



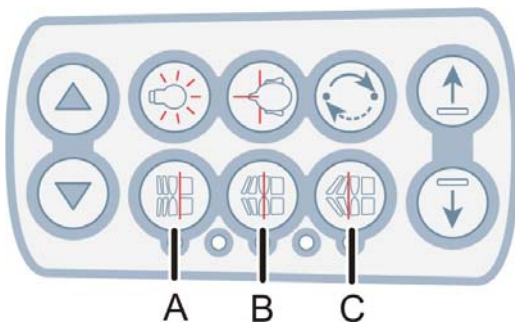
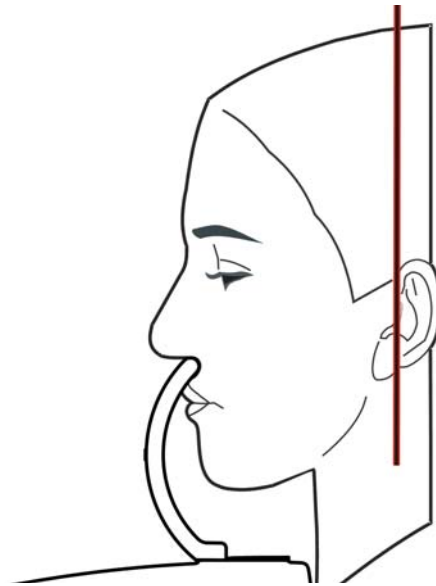
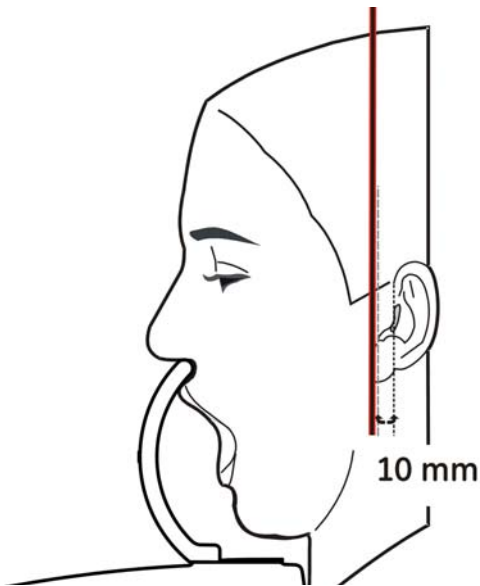
5. Check the position of the midsagittal light. If it is not on the midsagittal plane of the patient, adjust the patient's head.



6. Move the head support against the patient's forehead. Adjust the height. Close the temple supports.

7. Adjust the position of the TMJ light until it aligns in the middle of condyle.

Note! *The condyle moves forward by approximately 10 mm when the mouth is opened.*



TMJ light adjustment buttons:

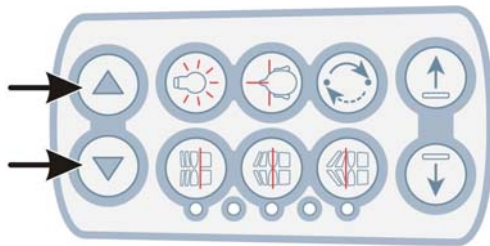
- A) Forward (towards the mirror)
- B) Reset
- C) Backward (away from the mirror)

5.3.6.3 Maxillary Sinus exposure

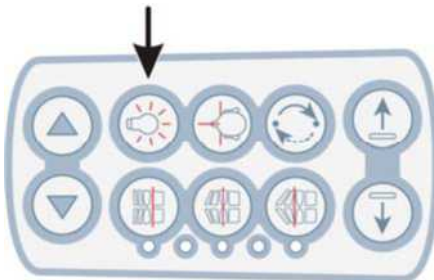
1. Insert the required positioning devices, bite fork with the bite block on the sinus rest. Place the disposable covers.

Note! Use a new disposable cover for every patient.

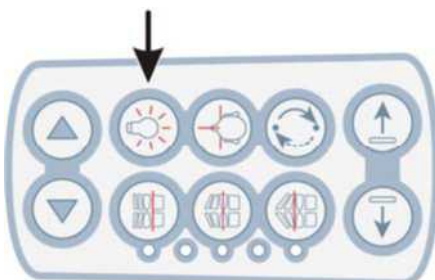
2. Adjust the unit height



3. Guide the patient to the unit and instruct to stand as straight and tall as possible. Ask the patient to take grip on the handles and bite on the bite block.
4. Adjust the height of the Frankfort-Horizontal plane (FH) laser to get the laser light over the orbita porion. Straighten the patient's head if needed.



5. Check the position of the midsagittal light. If it is not on the midsagittal plane of the patient, adjust the patient's head.



6. Move the head support against the patient's forehead. Adjust the height. Close the temple supports.
7. Adjust the position of the image layer as necessary. The image layer is 18 mm posterior compared to standard panoramic procedure.

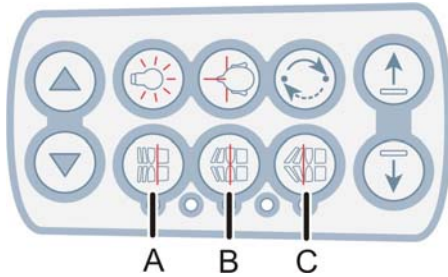


Image layer adjustment buttons:

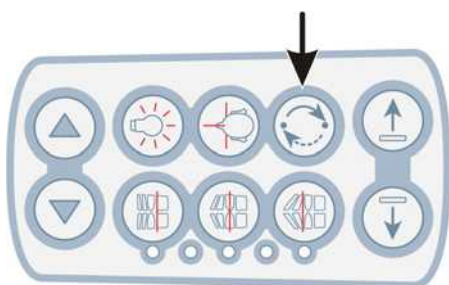
- A** 10 mm anterior
- B** Center
- C** 10 mm posterior

8. Ask the patient to press their tongue against the roof of their mouth and remain still for the duration of the exposure.

5.3.7 Taking the exposure

1. Press **Start position**. Check the patient positioning.

Protect yourself from radiation by standing behind a suitable x-ray radiation shield. Make sure that you can see and hear the patient during the exposure.



Note! In all examinations the user of the x-ray equipment should wear protective clothing. The operator does not need to be close to the patient during normal use. The protection against stray radiation can be achieved by using the hand switch not less than 2 m (7 ft) from the focal spot and the xray beam. Operator should maintain visible contact with the patient and technique factors. This allows immediate termination of radiation by the release of the exposure button in the event of a malfunction or disturbance.

Note! If the patient is nervous, or a child, you can demonstrate how the unit works to reassure them. Press the T (Test mode) button and then press and hold the exposure button. The unit will complete an exposure cycle without generating x-rays.



2. Press and hold down the exposure button. During the exposure you hear an audible signal and the exposure warning symbol on the touch screen display appears.

The unit rotates around the patient's head and stops. When the rotating unit stops, the exposure has been taken.

3. After the exposure the rotating unit is in 'patient out' position, if the exposure switch has been pressed until all



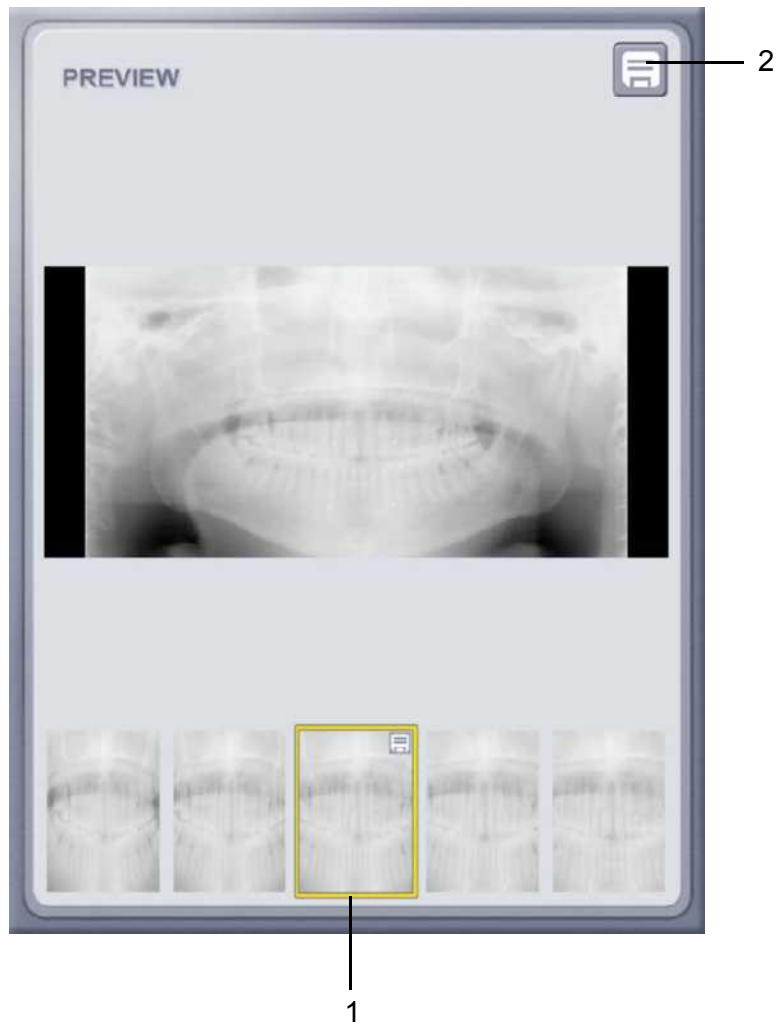
movements have stopped. Release temple supports. Guide the patient out. Remove disposable covers and disinfect the unit.

5.3.8 Multilayer Selection

For maximum image clarity and sharpness, the position of the focal trough may be adjusted after exposure. Five pre-adjusted images are calculated and displayed on the touch screen. In the midmost image no adjustment is applied, i.e. the focal through is located exactly at the layer laser position. To the left of this are images where the focal through is adjusted towards the patients neck (posterior). To the right are images where the focal through is adjusted towards the patients lips (anterior). The difference in adjustment between the images is 3 mm.

1. The touch screen shows one of the thumbnail images as magnified. To select which image is shown, press the corresponding thumbnail image in the lower part of the screen. Once shown as magnified, an image may be marked for saving by again pressing the thumbnail. An save indication icon is shown in the upper right of the thumbnail image. Repeat this process for all images of interest.
2. Press the save button in the upper right of the screen. The image(s) marked for saving will be sent to the workstation.
3. **PC:** The image(s) can be examined using the Cliview software. See CLINIVIEW™ user manual for details.

Note! *The X-ray unit may be configured to automatically select either the unadjusted (midmost) image or send all images to the workstation. In such cases, the selection process described above is bypassed.*



1. Preview selection

2. Save button

5.4 Cephalometric exposures

- Pediatric lateral projection
- Lateral projection
- PA projection
- Reverse towne projection
- Waters view
- Carpus view (Not available in USA and Canada)

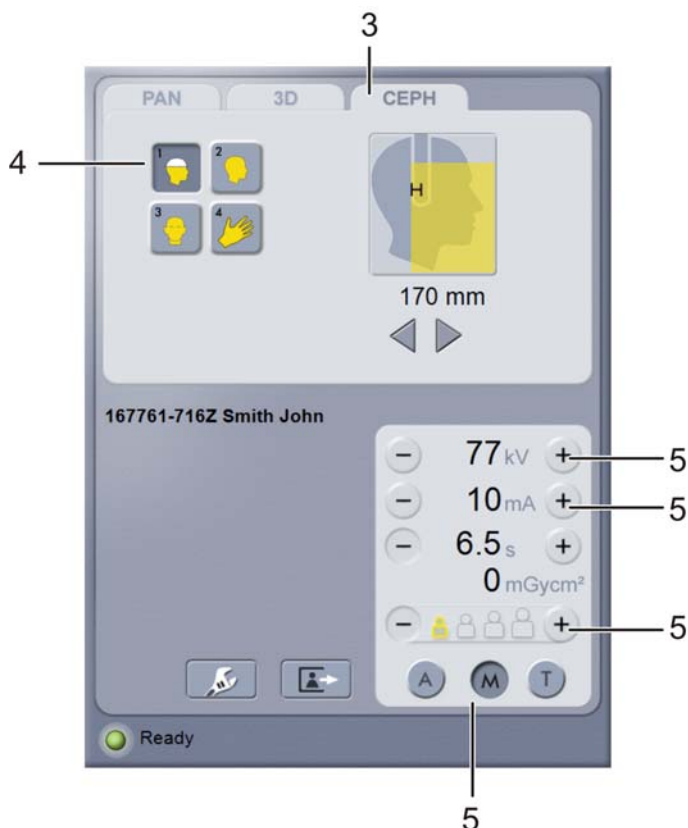


WARNING!

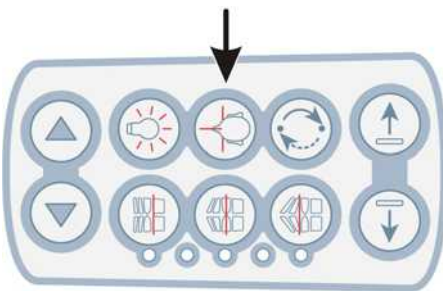
Remove all calibration tools, Pan and 3D patient positioning accessories before taking any cephalometric exposures!

5.4.1 General instructions

1. Move the ceph sensor to the ceph sensor holder.
2. **PC:** Click **Image Capture**.
3. Select CEPH tab.



4. Select the imaging program.
5. Select the Manual mode (Default).
Set the kV and mA or
select the patient size
(child, juvenile, adult, large adult).
6. Press the patient positioning button to drive the unit to
'patient in' position.



7. Ask the patient to remove any spectacles, hearing aids, removable dentures, jewellery and hair clips and pins. Place a protective lead apron on the patient.

5.4.2 Patient positioning

5.4.2.1 Pediatric lateral and Lateral projection

1. Unlock the lever and turn the ear rods to the lateral projection position. Lock the position. Tilt the nasion support aside. Place the disposable covers.

Figure 1.1 Unlock first the lever, turn the ear rods and lock the lever again.

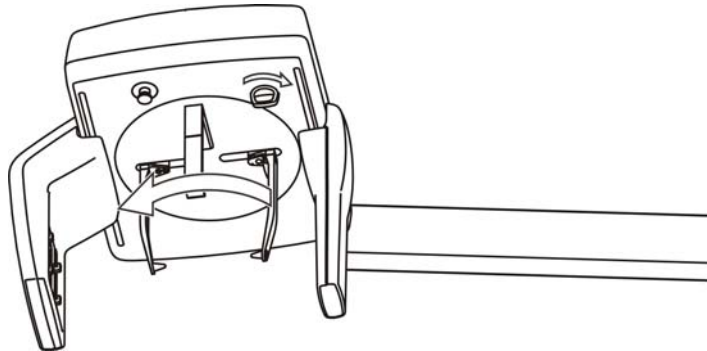
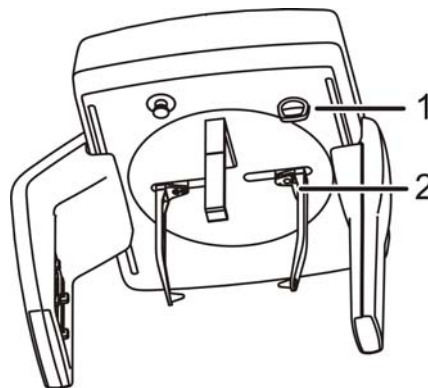
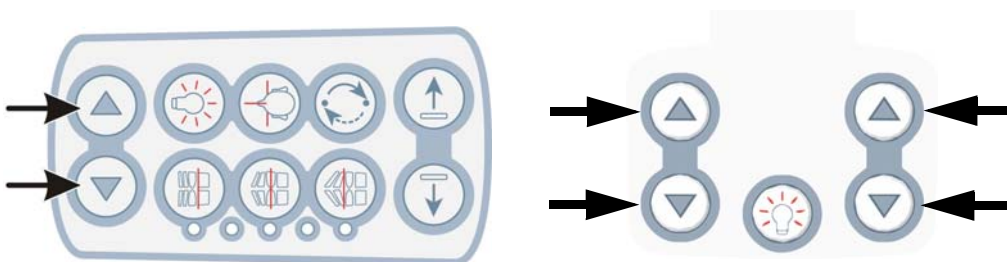


Figure 1.2 Locking lever (1), ear holder brake (2)

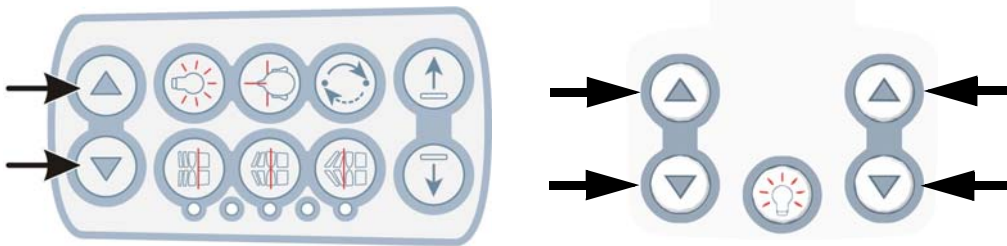


Note! Use a new disposable cover for every patient.

2. Adjust the unit height.

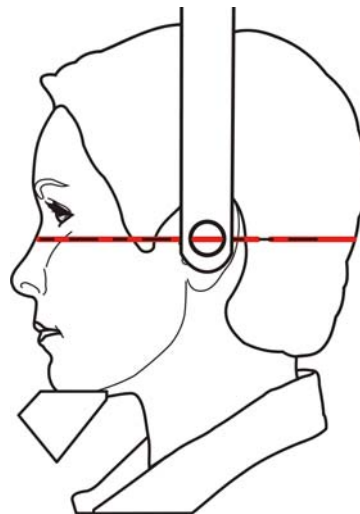
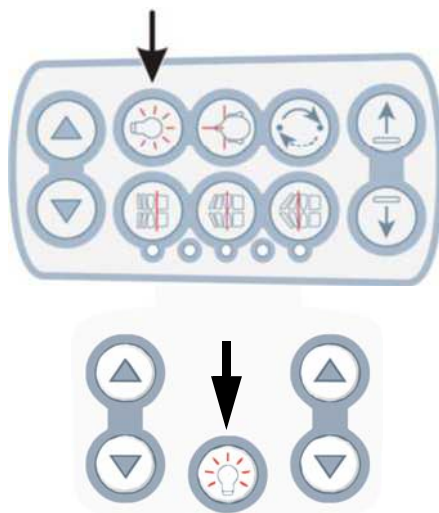


3. Guide the patient to the unit. Instruct the patient to stand as straight and tall as possible under the cephalostat head. Slide the ear rods towards to patient's ears. Tall patients can also sit on a chair.
4. Adjust the unit height to get the Frankfort-Horizontal plane (FH) laser light passing over the orbita porion.



Note!

The shown laser line is a horizontal reference line.



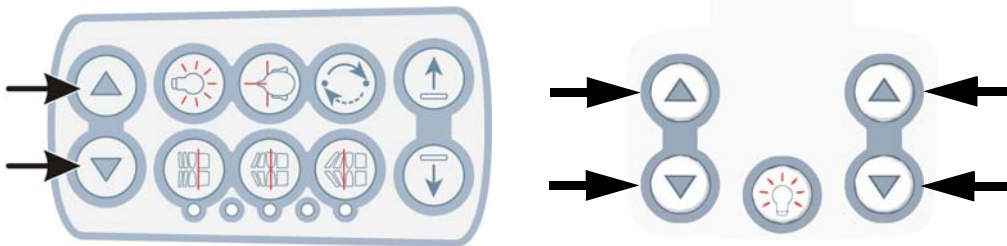
5. Tilt the nasion support down and slide it towards patient's nasion.

5.4.2.2 PA projection

1. Unlock the lever and turn the ear rods to the PA projection position. Lock the position. Tilt the nasion support aside. Place the disposable covers.

Note! Use a new disposable cover for every patient.

2. Adjust the unit height.



3. Guide the patient to the unit facing the sensor. Instruct the patient to stand as straight and tall as possible under the cephalostat head. Slide the ear rods towards patient's ears. Tall patients can also sit on a chair.

Figure 1.3 Unlock first the lever, turn the ear rods and lock the lever again.

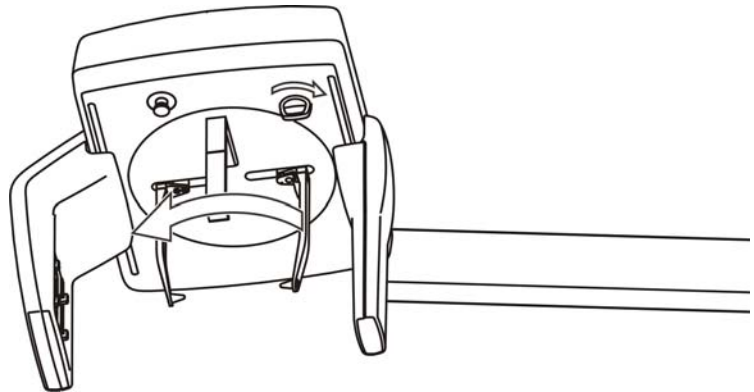
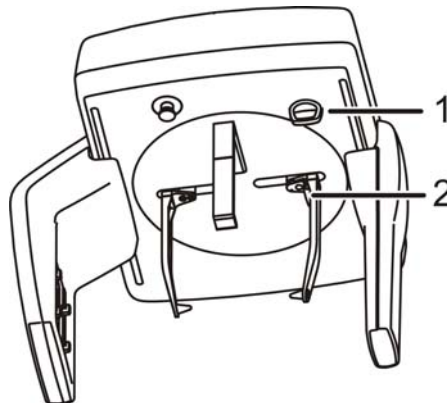


Figure 1.4 Locking lever (1), ear holder brake (2)



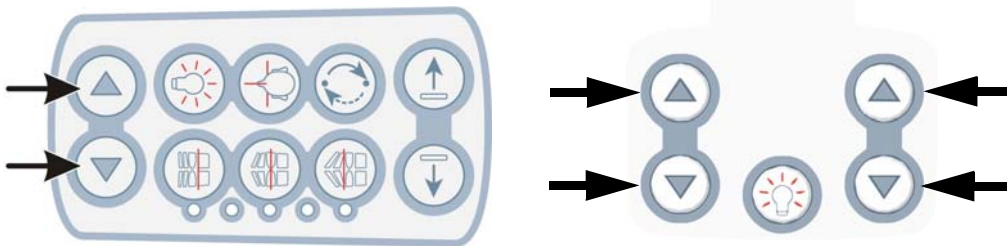
Note! Use a new disposable cover for every patient.

5.4.2.3 Reverse towne projection

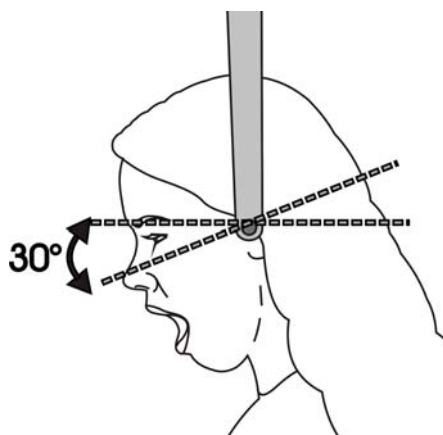
1. Unlock the lever and turn the ear rods to the PA projection position. Lock the position. Tilt the nasion support aside. Place the disposable covers.

Note! Use a new disposable cover for every patient.

2. Adjust the unit height.



3. Guide the patient to the unit. Instruct the patient to stand as straight and tall as possible under the cephalostat head.



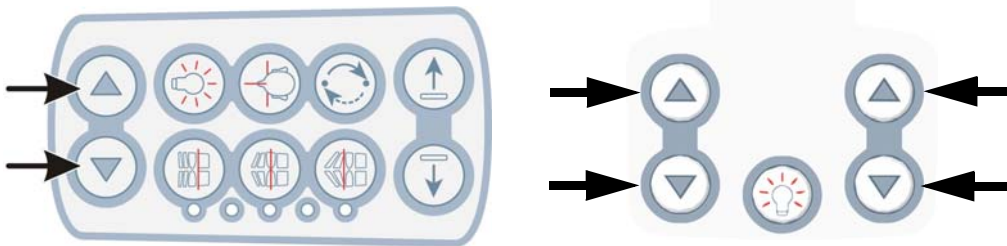
4. Turn the head ventral as reference to the canthomeatal line about 30° below the horizontal plane.
5. Slide the ear rods towards patient's ears.
6. Ask the patient open mouth maximally.

5.4.2.4 Waters view

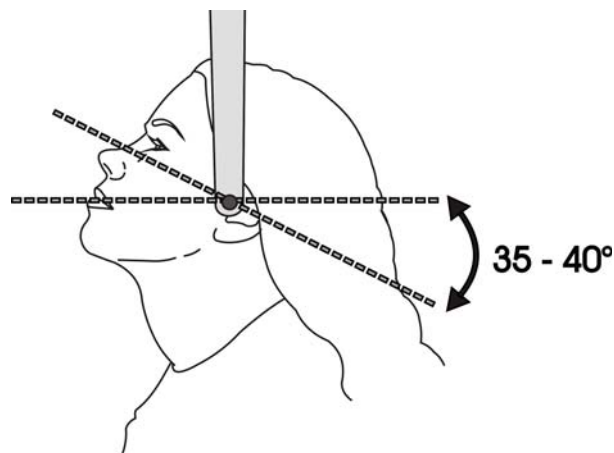
1. Unlock the lever and turn the ear rods to the PA projection position. Lock the position. Tilt the nasion support aside. Place the disposable covers.

Note! Use a new disposable cover for every patient.

2. Adjust the unit height.



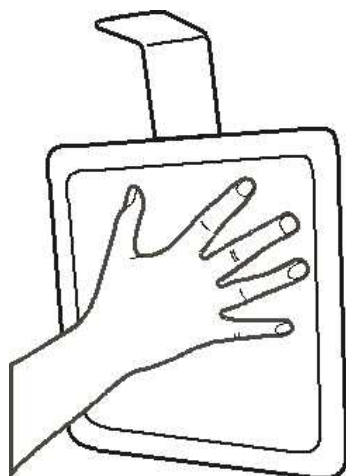
3. Guide the patient to the unit. Instruct the patient to stand as straight and tall as possible under the cephalostat head.
4. Turn the head dorsal as reference to the canthomeatal line about 35-40° above the horizontal plane.
5. Slide the ear rods towards patient's ears.
6. Ask the patient open or close mouth.



5.4.2.5 Carpus view (Not available in USA and Canada)

Before taking Carpus image make sure this imaging method is approved by local authorities of your country.

Note! If Carpus program button is not displayed in Cephalometric imaging modality tab, ask local distributor to activate the button. (Not in USA or Canada).



1. Unlock the lever and turn the ear rods to the PA projection position. Lock the position. Tilt the nasion support aside. Place the carpus holder to the nasion support holder.
2. Adjust the unit height if needed.
3. Ask the patient to remove rings and metal objects and to place hand on the carpus holder.

5.4.2.6 Taking the exposure



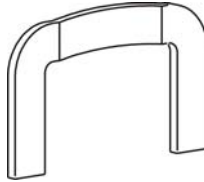
1. Protect yourself from radiation by standing behind a suitable x-ray radiation shield. Make sure that you can see and hear the patient during the exposure.
2. Press and hold down the exposure button. During the exposure you hear an audible signal and the exposure warning symbol on the touch screen display appears.
3. Release the ear rods and guide the patient out. Remove disposable covers and disinfect the unit.
4. **PC:** The image can be examined using the CLINIVIEW™ software. See CLINIVIEW™ user manual.

Note! In all examinations the user of the x-ray equipment should wear protective clothing. The operator does not need to be close to the patient during normal use. The protection against stray radiation can be achieved by using the hand switch not less than 2 m (7 ft) from the focal spot and the xray beam. Operator should maintain visible contact with the patient and technique factors. This allows immediate termination of radiation by the release of the exposure button in the event of a malfunction or disturbance.

5.5 3D exposures

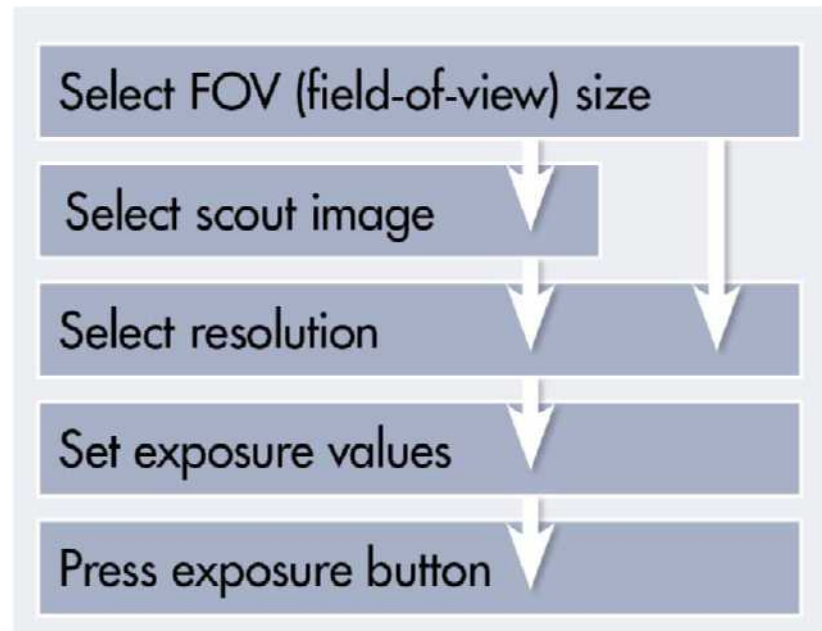
5.5.1 Positioning devices

Chin support

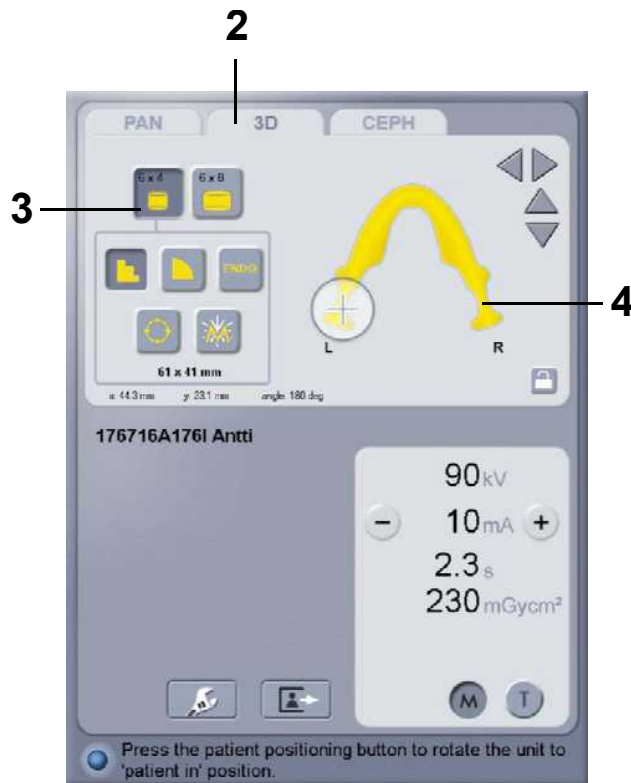


5.5.2 General instructions

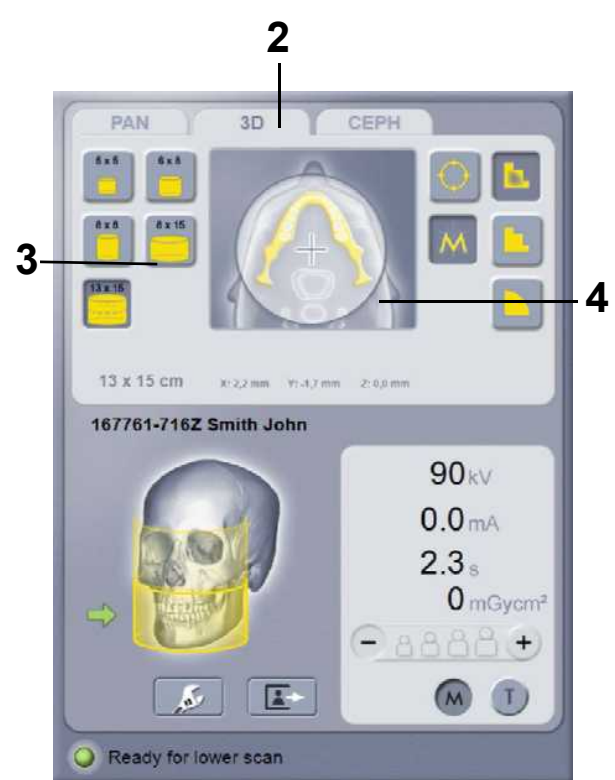
Workflow



1. PC: Click **Image Capture**.
2. Select the 3D modality tab.
3. Select the Field Of View (FOV).
4. With 3D SFOV unit move the FOV cursor on the area of interest. For precise adjustment the arrow keys can be used. The 3D FOV is positioned more accurately by using the scout image mode. The area of interest can be adjusted on the touch screen display after the scout image has been taken.



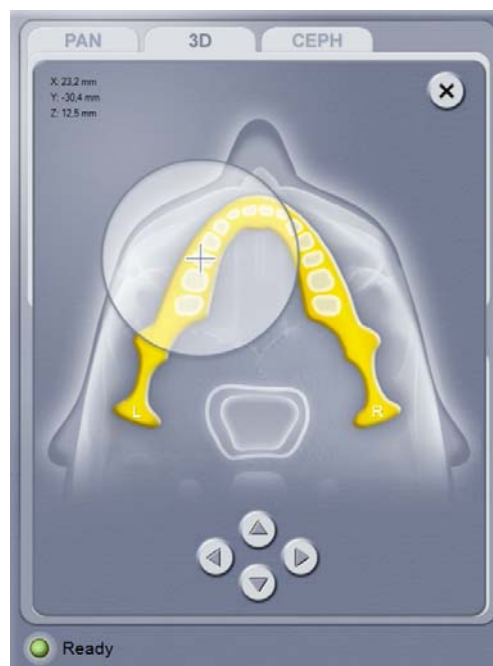
3D SFOV touch panel



MFOV touch panel



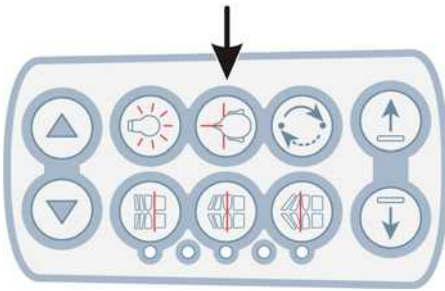
With MFOV (Maxio) touch panel touch somewhere in the dentition. Pop up screen appears. Move the FOV cursor on the area of interest. For precise adjustment use the arrow keys.





5. With 3D SFOV lock the position by pressing the lock key. Use the same key for unlocking if the position needs to be changed. Lock key is not in the MFOV (Maxio) touch panel.

6. Press the patient positioning button to rotate the unit to 'patient in' position



7. Open the temple supports. Remove head support in case FOV 130 x 150 mm is selected.
8. Ask the patient to remove any spectacles, hearing aids, removable dentures, jewellery and hair clips and pins.

5.5.3 Patient positioning

1. Insert the chin support. In case of FOV 130 x 150 mm insert lower head support and chin support.

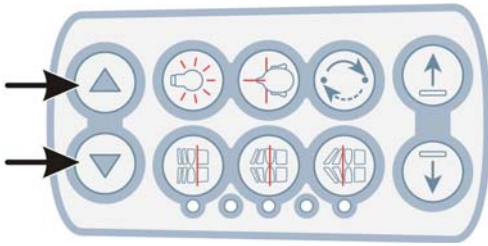


Fig 1.9. Chin rest.

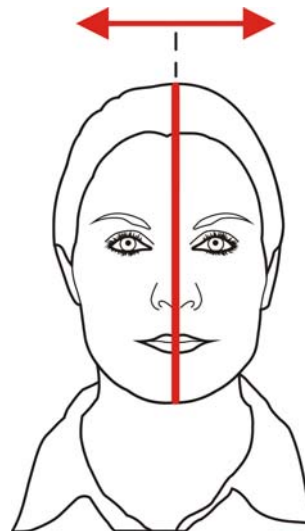
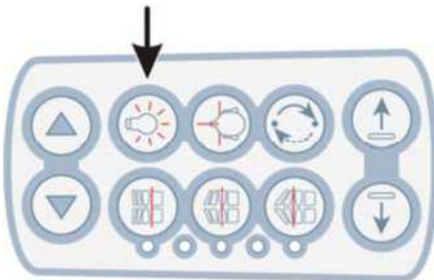


Fig 1.9. Lower head support and chin support for FOV 13 x15.

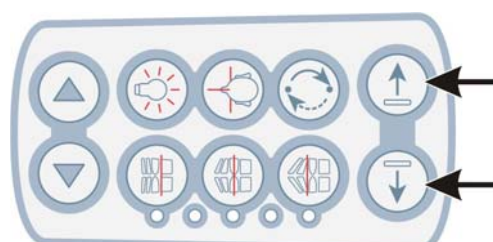
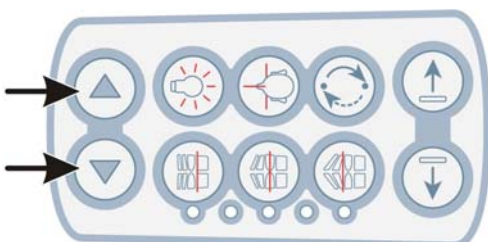
2. Adjust the unit height.

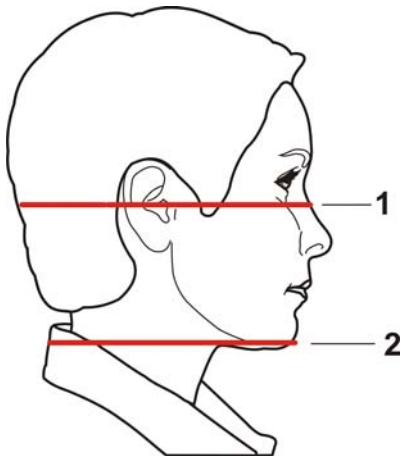


3. Guide patient to the unit. Instruct patient to stand as straight and tall as possible next to the unit. Patient can also be imaged in sitting position. Ask the patient to take grip on the handles and place chin on the chin rest.
4. Check the position of the midsagittal light. If it is not on the midsagittal plane of the patient, adjust the patient's head.



5. Adjust the unit height and chin rest height to get the area of interest between the top and bottom FOV lights. Position the patient so that the occlusal plane is horizontal.

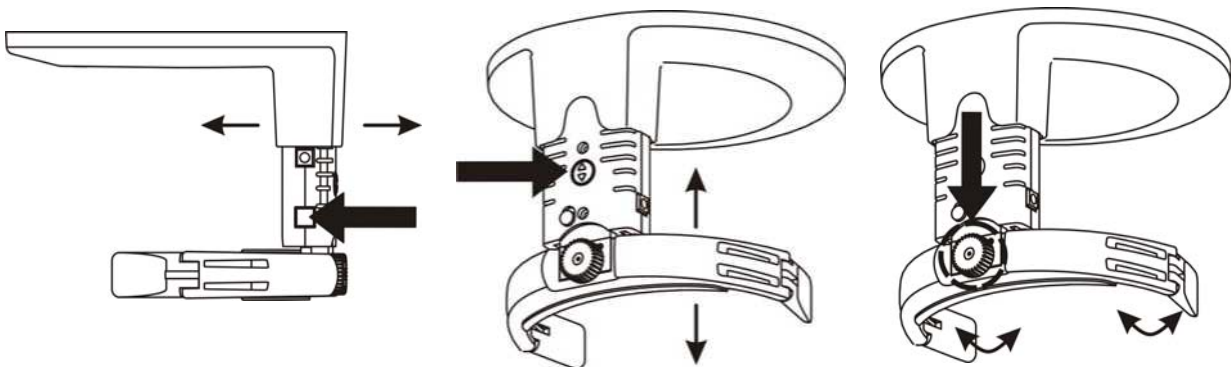




1. H light, top of FOV
2. H light, bottom of FOV

Note! In case of 130 x 150 mm Field of View, chin rest movement cannot be used for FOV height adjustment. To adjust FOV higher, remove the chin support. The FOV height (130 mm) is indicated with FH light. Move FH light to 130 mm position (up).

6. Move the head support against the patient's forehead. Adjust the height. Close the temple supports. In case of FOV 13 x 15 adjust height of the temple support to correct level and close head support strap.



7. Select the FOV size based on the indication.

SFOV	61 x 41 mm	
	61 x 78 mm	
MFOV (MAXIO)	50 x 50 mm	
	61 x 78 mm	
	78 x 78 mm	
	78 x 150 mm	
	130 x 150 mm	

8. Select either the scout for FOV positioning or 3D resolution for direct image capture.

SCOUT

Scout image

**3D SFOV****A** Standard resolution**B** High resolution**C** Endo resolution**MFOV (Maxio)****A** Low Dose**B** Standard resolution

C High resolution



D Endo resolution



5.5.4 Scout image

Note! When scout image has been selected, follow these instructions.

Note! For 130 x 150 FOV, the scout image is showing only the first (lower) scan.



1. Press and hold down the exposure button. During the exposure you hear an audible signal and the exposure warning symbol on the touch screen display appears.
2. Scout preview image appears to the touch screen display.

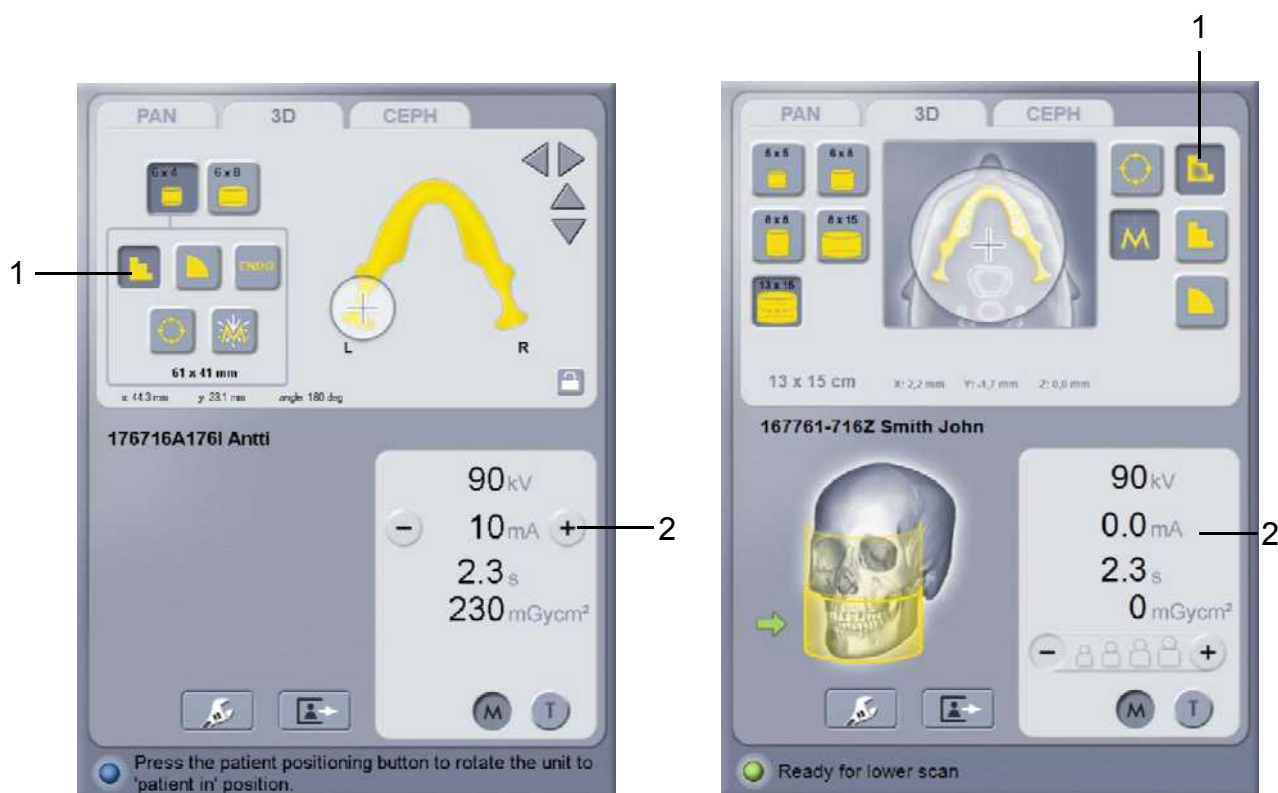
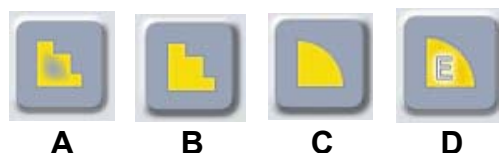


3. Fine adjust scout position using the side arrow keys. Press the right upper corner icon to continue.
4. Take a new scout or save current and continue to 3D image.

5.5.5 3D image

Note! After selecting FOV size and taking the scout image.

1. Select the Low Dose (A), standard (B), high resolution (C) or endo program (D). Endo program is only for 61 x 41 mm 3D SFOV and 50 x 50 mm MFOV (Maxio). Low Dose resolution is only available for MFOV (Maxio).



2. Select mA.



3. Press and hold down the exposure button. During the exposure you hear an audible signal and the exposure warning symbol on the touch screen display appears.
4. Select **MAR** ON or OFF based on your estimation of the need. See chapter 3.4.1. Instructions for using MAR.

5.5.5.1 Stone model and radiographic guide scan

For scanning of stone models and radiographic guides, a positioning plate is available for the system.

1. Scan the patient with an open bite by securing the bite with cotton pads.
2. Install positioning plate. Position stone model.



Note! It is recommended to use a sponge or a foam under radiographic guide during the scan.

3. Take scout image with default values. Correct position if needed.



4. Select same resolution and parameters as in patient scan.

Note! More detailed instructions in *OP300 Quick Guide Stone model and radiographic guide scan protocol*.

5.6 Warnings and error messages

The unit responds to error situations by showing a dialog box containing an error code and descriptive text on the touch screen.

When an error code appears on the display the unit will stop working and cannot be operated while the error code is on the display. In less severe cases a warning message will be displayed, leaving the unit operable.

5.6.1 Acknowledging errors

Most errors may be acknowledged by closing the dialog box the error is reported in. Some errors require the unit to be rebooted. If such an error occurs, or if the unit fails to operate as described in the user's manual, switch the unit off, wait a few seconds and switch the unit on again.

5.6.2 Image transfer errors

If an image is not transferred successfully to the PC, close and then reopen the dental imaging software and/or restart the PC. DO NOT restart the unit as this will erase any image that is stored in the unit memory and this retrievable image will be lost. If restarting the PC and/or restarting the dental imaging software does not allow you to retrieve the images, contact technical support without restarting the unit.

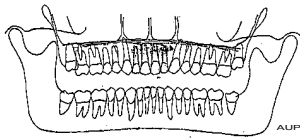
6 Troubleshooting

High quality images with sharp contrast and good detail provide optimum diagnostic information. Images with less quality are usually the result of one or more common problems.

6.1 Patient positioning

Problem

Incisors and canines narrow and unsharp. Overshadow in molar and premolar areas. Rows of teeth are compressed.



Possible cause

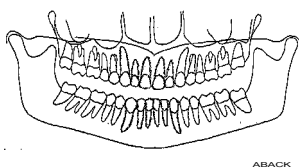
1. Occlusal correction of focal trough set too far posterior
2. Image layer laser light not obeyed
3. Bite block was not used

Remedy

1. Check patient positioning with laser light lines and occlusion correction buttons
2. Check patient positioning with laser light lines and occlusion correction buttons
3. Insert bite block

Problem

Incisors and canines wide and unsharp. Rows of teeth widened.



Possible cause

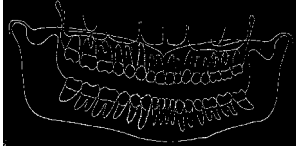
1. Occlusal correction of focal trough set too far anterior
2. Image layer laser light not obeyed
3. Bite block was not used

Remedy

1. Check patient positioning with laser light lines and occlusion correction buttons
2. Check patient positioning with laser light lines and occlusion correction buttons
3. Insert bite block

Problem

Teeth appear wider on one side and narrower on the opposite. Ramus widths are different on opposite sides.

**Possible cause**

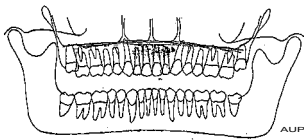
1. Midsagittal line not obeyed
2. Patient's head not in center position

Remedy

1. Check patient's mid sagittal plane with laser light line
2. Check that patient's head is centered, and that the head support side clamps were closed to keep the head straight.

Problem

The shadow of hard palate is exposed over maxillary molars. Row of teeth has a wavy appearance. TM joints are exposed outward. Image is not "smiling". Mandible is imaged sharper than maxilla.

**Possible cause**

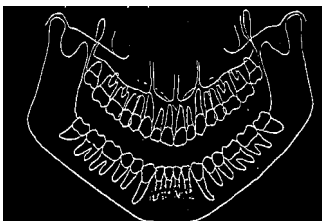
Patient head tilted back

Remedy

Check FH plane

Problem

Rows of teeth curved upwards. Mandibular incisors are unsharp. TMJ joints exposed high and are often cut off from the image. Image is "smiling" too much.

**Possible cause**

Patient head tilted forward

Remedy

Check FH plane

Problem

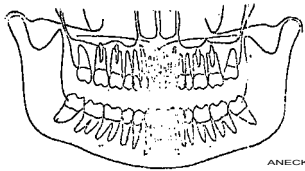
Middle area of the image too bright and unsharp. Spine shadow.

Possible cause

Patient's neck was not stretched

Remedy

Stretch patient's neck

**Problem**

Black shadow over maxillary teeth apex area.

Possible cause

Tongue was not against the roof of palate.

Remedy

Ask patient to swallow and place tongue against the roof of palate during the exposure.

Problem

TMJ's exposed on different heights on image. Bilateral distortion in molar and premolar regions.

Possible cause

1. Patient tilted to one side
2. Midsagittal laser light line not obeyed.

Remedy

1. Check midsagittal plane and center patient's head.
2. Check midsagittal plane and center patient's head.

Problem

Rows of teeth exposed too high. TMJ's cut off.

Possible cause

1. Chin was not resting on chin support
2. Patient positioned too high

Remedy

1. Check patient positioning and type of bite fork rod.
2. Check patient positioning and type of bite fork rod.

Problem

Rows of teeth exposed too low. Mandible not exposed completely to the image.

Possible cause

Chin rest was not used with bite fork.

Remedy

Install chin rest.

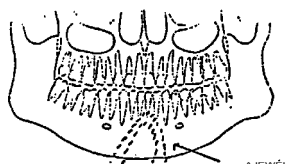
6.2 Image appearance

Problem	Possible cause	Remedy
Images are too light	1. CLINIVIEW™: Contrast and brightness not optimum 2. CLINIVIEW™: Gamma not set correctly	1. Adjust contrast and brightness. 2. Select a more fitting histogram type and check gamma setting.
Images are too dark	1. CLINIVIEW™: Contrast and brightness not optimum. 2. Manual technique factors used too high.	1. Adjust contrast and density. 2. Decrease technique factors.
Lack of image contrast	1. CLINIVIEW™: Contrast and brightness not optimum. 2. kV used is too high. 3. Gamma value is not correct for the monitor being used.	1. Adjust contrast and brightness. 2. Lower the kV setting. 3. Adjust Gamma value

6.3 Artefacts

Problem

Irregular, bright shadows or artefacts



Possible cause

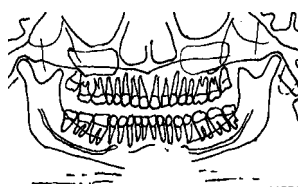
Patient is wearing metal objects, such as earrings, necklace etc.

Remedy

Ask patient to remove objects.

Problem

An unexposed area is shown down in the lower middle section of the image.



Possible cause

Lead apron misplaced.

Remedy

Check the lead apron positioning.

Problem

Partial lack of detail and motion artefacts. Irregular vertical bright lines on image.

Possible cause

Patient has moved during the exposure.

Remedy

Retake the image.

Problem

Vertical dark lines on image.

Possible cause

Patient's shoulder in touch with machine parts.

Remedy

Check patient positioning.

Problem

Patient's right side tooth are not exposed.

Possible cause

Exposure button released prematurely.

Remedy

Retake the image.

Problem

Right and left image sides are uncomplete. TMJ's are not shown.

Possible cause

TMJ areas in sectional images where deselected.

Remedy

Select all sections in panoramic image.

Problem	Possible cause	Remedy
A light horizontal line on QA image.	Bite block was left on place.	Remove the bite block and retake QA image.

Problem	Possible cause	Remedy
Horizontal lines on image.	Sensor problem.	Consult the dealer.

Problem	Possible cause	Remedy
CEPH: Lateral view has 2 ear holder pins.	1. Cephalostat lock not locked 2. Ear holders misaligned	1. Lock it 2. Call service

6.4 Unit operation

Problem	Possible cause	Remedy
Back of the patient's head is touching the x-ray tube during the exposure.	1. Patient's head inclination not correct 2. Patient is too big for the unit. 3. Patient has slumped.	If the image is not acceptable then 1. Check the head position and retake the image. 2. Check the patient positioning. Make the exposure even though the head may touch the tube head. 3. Check the patient positioning. Make the exposure even though the head may touch the tube head.
Patient's shoulders are touching the x-ray tube or sensor.	Patient is too big for the unit. Wide and high shoulders.	Reverse patient's hands on handles: left to right side handle and vice versa.

7 Maintenance

7.1 Maintenance procedure

The maintenance procedure described below shall be seen as a minimum requirement and can be made more stringent to comply with regulations regarding the use and maintenance of dental x-ray devices that are in force in the country in which the unit is installed.

7.1.1 Annual maintenance

An annual maintenance procedure must be carried out at least once a year by qualified service personnel. Contact your local distributor for details.

7.1.2 Calibration intervals

To keep the image quality at best possible level, calibrations and quality checks shall be carried out at regular intervals according to the table below.

Modality	Minimum requirement	Recommendation
3D	Two (2) times annually	Four (4) times annually
Panoramic	Annually during normal maintenance	Two (2) times annually
Cephalometric	Annually during normal maintenance	Two (2) times annually

Note! The calibrations mentioned in this manual can be done by the user or qualified service personnel.

7.2 Changing the fuses

Main fuses are located next to the on/off power switch. Push inward on the fuse base and twist it counterclockwise with a screwdriver. The fuse with the base comes out.

Remove the fuse from the base and replace it with the new one. Repeat this with each blown fuse. Fasten both fuses by pushing the base in and twisting it clockwise with a screwdriver.

Use only appropriate fuses:

- Line voltage 220-240 Vac: 326 Littelfuse 10A (slow blow) or Cooper Bussman MDA-10 (time delay)
- Line voltage 100-120 Vac: 326 Littelfuse 15A (slow blow) or Cooper Bussman MDA-15 (time delay)

7.3 Cleaning and decontaminating the unit



CAUTION!

Switch the unit off or disconnect it from mains before cleaning the unit. If you use a spray cleaner do not spray into any ventilation grills. Do not allow water or other cleaning liquids to enter the unit interior since these may cause short-circuits or corrosion. The unit should be cleaned after every usage.



CAUTION!

Clean the dust off the unit regularly. The unit might overheat if excess dust is gathered on the cooling grilles.

Unit surfaces

All surfaces can be wiped clean with a soft cloth dampened with a mild detergent, e.g. soapy water. DO NOT use abrasive cleaning agents or polishes on this equipment.

Positioning light covers

The positioning light covers are made of clear plastic. Use a soft cloth dampened with a mild detergent, e.g. soapy water. NEVER use abrasive cleaning agents or polishes to clean the covers.

Surfaces that the patient touches

All surfaces and parts that the patient touches or comes into contact with must be decontaminated after each patient. Use a disinfectant that is formulated specifically for decontaminating dental equipment and use the disinfectant in accordance with the instructions supplied with the disinfectant. All items and surfaces should be dried before next usage.

Note! *Wear gloves and other protective equipment during decontamination process. In accordance with the instructions supplied with the cleaner.*



WARNING!

Do not use any disinfecting sprays since the vapor could ignite causing injury.

Decontamination techniques for both the unit and the room must comply with all laws and regulations within the local jurisdiction.

Examples of cleaning agents that can be found in disinfectant products which are allowed or prohibited **when cleaning the unit:**

Allowed: Methanol (metyl alcohol), Soap, Isopropyl alcohol, distilled water.

Not allowed: Bentzene, Chlorine bentzene, Acetone, Acetic ether, agents containing phenol, paracetic acid, peroxide and other oxygen-cleaving agents, sodium hypochlorite and iodine-cleaving agents.

Autoclave

Some removable parts in contact with the patient may be autoclaved. These parts are: bite rods, bite guides and chin supports.

If autoclaving is performed for these items, disinfection by alternate methods is not needed.

Steam sterilization

Recommended parameters for sterilizable parts are:

- Gravity-displacement steam sterilization
 - "Flash" sterilization:
 - Temperature:* 270 F (132°C)
 - Exposure time:* 3 minutes
- Prevacuum steam sterilization
 - "Flash" sterilization:
 - Temperature:* 270 F (132°C)
 - Exposure time:* 3 minutes
- Steam-flush pressure-pulse steam sterilization
 - Temperature:* 270 F to 275 F (132°C to 135°C)
 - Exposure time:* 3 to 4 minutes

8 Calibration and adjustment

8.1 Introduction

Calibrations and quality checks are performed by taking exposures of calibration tools. The system does needed adjustments according to the image data captured. For panoramic and cephalometric quality checks the quality is visually evaluated by the operator.

Resulting from the each calibration is an image containing calibration results, telling the operator how to proceed with the calibration and adjustment procedure. In addition to the calibration name (e.g. Adjustment panCol) the images contain image data sampled during the calibration, adjustment instructions and a "Passed / Not Passed / Failed" calibration status.

- **Passed** means that the calibration program is successfully done. Move on to next calibration.
- **Not passed** means that adjustment is still needed. Follow the instructions the image (if any) and take another exposure. Some calibration programs are iterative and demand a few repetitions.
- **Failed** means that the system could not decide what adjustment should be done in order for the calibration to succeed. This calibration status is always the result of some error condition. Taking another exposure will not help. The image may give a hint on what the problem is (e.g. no radiation, collimator severely tilted, image data corrupted...). Contact service if the problem persists after restarting the unit and PC.

8.2 Preparing for calibration

1. Close the head support and lock it in its upmost position.
2. Switch the PC and unit on.
3. **PC:** Open the dental imaging software and then open a patient (card) and give it an identifiable name, for example: calibration (refer to the user's manual supplied with the dental imaging software for more information).
4. **PC:** Click the image acquisition button to activate image capture.



5. Touch the **settings** button on the touch screen display.
6. Select the **Quality assurance** button.
The calibration display appears.



8.3 Panoramic calibration

8.3.1 Panoramic geometry calibration



1. Select the program.



2. Press **Patient Positioning**.

3. Install the double cone calibration tool.



4. Take an exposure.

5. Repeat the calibration until calibration result "passed" is achieved.



8.3.2 Panoramic pixel calibration

Note! The pixel calibration results are sensor specific. If the x-ray unit is equipped with separate panoramic and cephalometric sensors, the cephalometric sensor cannot be used for panoramic imaging without re-calibration (and vice versa).

Note! Re-do panoramic pixel calibration, if cephalostat sensor is moved to panoramic side or the sensor is changed.

1. Remove the double cone calibration tool.



2. Select the program.



3. Press **Patient Positioning**.

4. Take an exposure.

FFPan detector calibration Passed

CTQ [>3.0] = 9.0

Signal level [>5461] = 10559

Dark level [<819] = 291

8.3.3 Panoramic Quality Check (optional)

Note! Use the same tool for cephalostat Quality Check.

1. Attach a panoramic Quality Check Tool (optional) to the chin support.



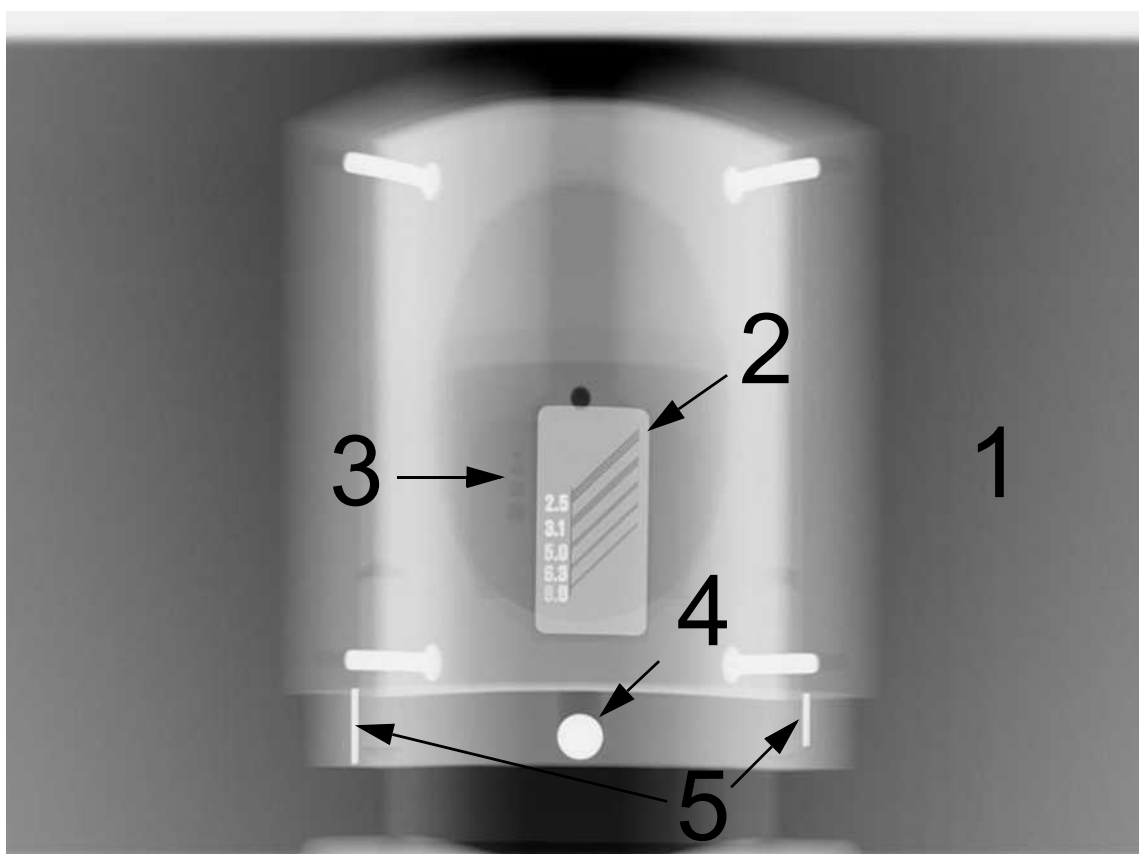
2. Select the Pan QC program.



3. Press **Patient Positioning**.

4. Take an exposure.

5. Visually evaluate the result using the installed imaging software.



Subjects to be evaluated:

1. Smoothness of the exposed area. Non-exposed area surrounds the whole image.
2. High contrast resolution; minimum 3.1LP/mm must be distinguishable.
3. All four low contrast holes must be visible.
4. Roundness of the ball.
5. The ball should be placed symmetrically between the two pins. The distance from both pins to center should be equal length.

Note! The panoramic QC collimator is equipped with a 0.8 mm copper filter. If more filtration is required, additional filtration may be attached to the tubehead cover. The unit may be configured to use higher exposure values to compensate for an additional 1 mm copper filter.

Ask Technical Support to adjust the copperthickness setting as required.

8.4 3D calibration

8.4.1 3D geometry calibration

1. Attach the base of the phantom to the lower shelf.
Level it with the bubble.

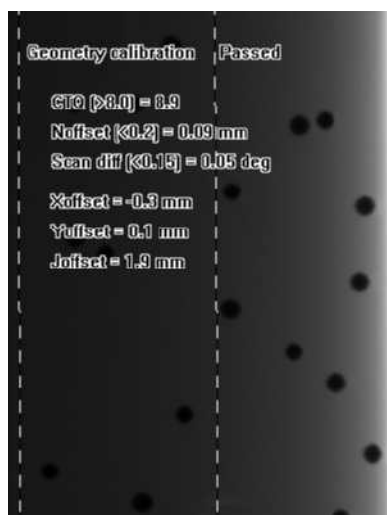


2. Select the program. There is a calibration procedure for both 3D imaging modes, standard and high resolution. Standard geometry calibration has to be done first.



3. Press **Patient Positioning**.

4. Install the 3D calibration phantom.



5. Take an exposure.
6. Repeat the calibration until calibration result "passed" is achieved. This calibration is only needed with 3D units.

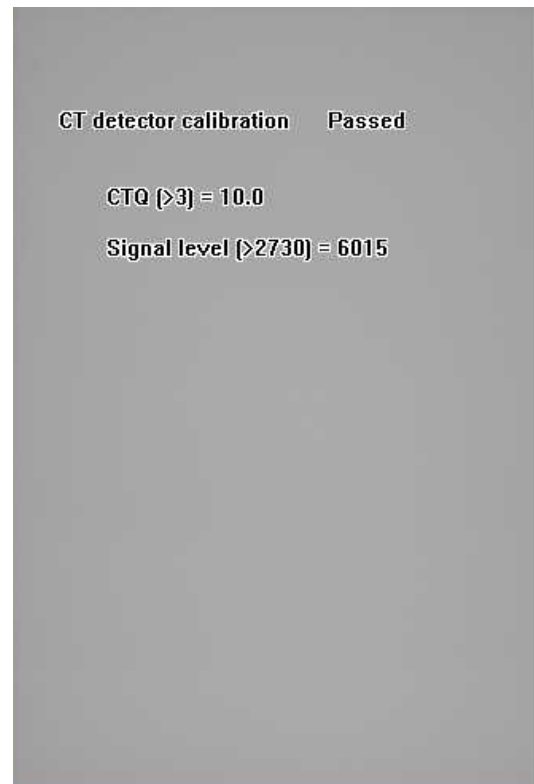
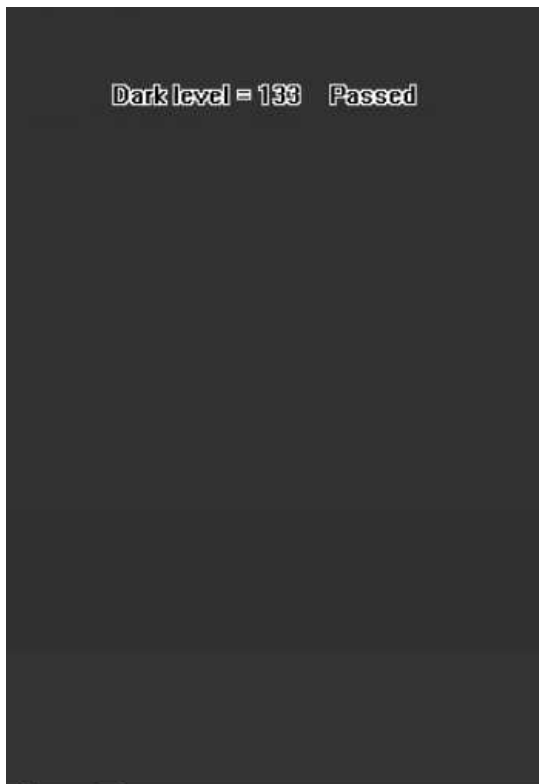
8.4.2 3D pixel calibration



1. Remove the 3D calibration phantom.
2. Select the program.



3. Press **Patient Positioning**.
4. Take an exposure. The result image informs when the calibration is passed.



8.4.3 3D Quality Check program

1. Attach the QC phantom to the unit.



2. Ensure that QC phantom is aligned with libel.



3. Select the 3D QC program.



4. Press **Patient Positioning**.

5. Take an exposure.

6. The resulting image contains information on whether the quality check was passed.

8.5 Cephalometric calibration

8.5.1 Ceph pixel calibration

Note! The pixel calibration results are sensor specific. If the x-ray unit is equipped with separate panoramic and cephalometric sensors, the cephalometric sensor cannot be used for panoramic imaging without re-calibration (and vice versa).

Note! Re-do panoramic pixel calibration, if cephalostat sensor is moved to panoramic side or the sensor is changed.

1. Rotate the ear holders into PA view position and move them completely apart. Turn nasion support up out of the way.



2. Select Ceph Pix program.



3. Press **Patient Positioning**.

4. Take an exposure.

5. This calibration should always be a pass.

8.5.2 Ceph Quality check program (Optional)

Note! Rotate the ear holders into PA view position. Turn nasion support up out of the way.

1. Attach the QC phantom to the ceph unit and ensure that it's leveled from the spirit level.





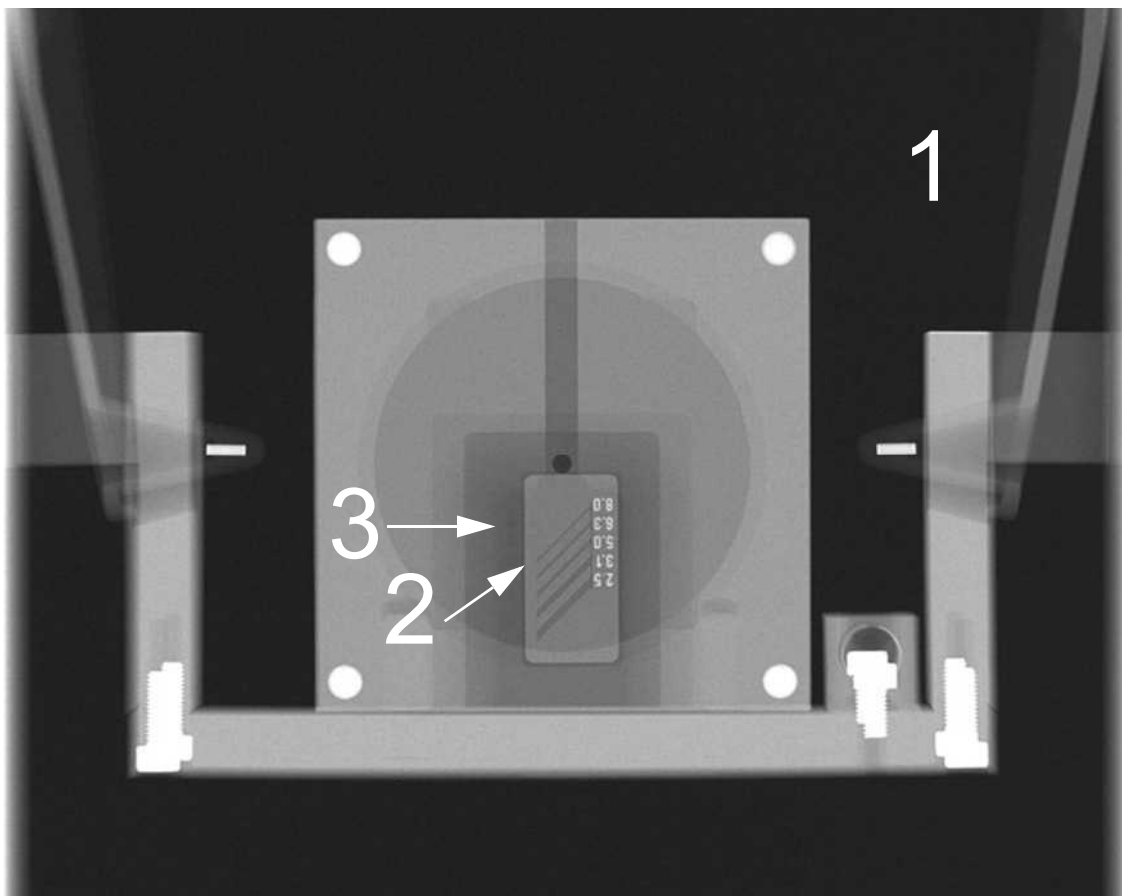
2. Select the Ceph QC program.



3. Press **Patient Positioning**.

4. Take an exposure.

5. *Visually evaluate the result using the installed imaging software.*



Subjects to be evaluated:

1. Smoothness of the exposed area. Non-exposed area surrounds the whole image.
2. High contrast resolution; minimum 3.1LP/mm must be distinguishable.
3. All four low contrast holes must be visible.

9 Technical data

9.1 Technical specifications

Manufacturer	Instrumentarium Dental, Nahkelantie 160 (P.O. Box 20) FIN-04300 Tuusula, FINLAND
Quality system	In accordance with ISO13485 and ISO9001 standard
Environmental management system	In accordance with ISO14001 standard
Conformity to standards:	IEC 60601-1: 1988 and A1+A2 IEC 60601-1-1: 2000 IEC 60601-1-4: 1996 and A1 IEC 60601-2-7: 1998 IEC 60601-2-28: 1993 IEC 60601-2-32: 1994 IEC 60601-1-2: 2001 and A1 IEC 60601-1-3: 1994 UL 60601-1: 2003 CAN/CSA –C22.2 No. 601-1-M90 and S1+A2 standards This product complies with DHHS 21 CFR Chapter I, Subchapter J at the date of manufacture. OP300 is in conformity with the provisions of Council Directive 93/42/EEC as amended by the Directive 2007/47/EC concerning medical devices. Performance Standards and European Union Directive 93/42/EEC (Medical Devices Directive).

Product name	ORTHOPANTOMOGRAPH® OP300
Model:	OP300
Product type:	Digital dental imaging system with panoramic cephalometric and Cone Beam 3D imaging programs.

Unit data	
Protection against electric shock	Class I
Degree of protection	Type B applied with no conductive connection to the patient
Protection against the ingress of liquids	IP20
Disinfection methods	- mild soapy water (non-abrasive) - non-alcohol based disinfectant for the chin rest - disposable plastic covers for bite block, chin rest and chin support
For use	In environments where no flammable anaesthetics nor flammable cleaning agents are present
Mode of operation	continuous operation/intermittent loading
Safety	IEC 60601-1
EMC Classification	Class B

Tube head assembly	
Tube head assembly type	THA 300
Tube type	Toshiba D-052SB, D-054SB-C Stationary anode
Tube voltage	57 - 90 kV
Max. tube current	16 mA
Max. electric output	1,44 kW
Target angle	5 degrees
Focal spot	0,5 x 0,5 mm (IEC 336/1982)
Nominal anode input	1750 W
Reference axis	In the middle of the panoramic sensor's active area
Max. anode heat content	35 kJ
Max. X-ray tube assembly heat content	385 kJ
Max. continuous heat dissipation of the X-ray tube assembly	38 W
Total filtration	>3,2 mm Al
Leakage Technique Factors	90 kV / 4 mA

Electrical connections	
Nominal mains voltage	220-240V / 100-120V (Selectable)
Input power frequency	50 / 60 Hz
Nominal current	10A @ 230 VAC, 15A @ 110 VAC
Fuses	230 Vac: Littelfuse 326 (slow blow) 10A Cooper Bussman (time delay) MDA-10 110 Vac: Littelfuse 326 (slow blow) 15A Cooper Bussman (time delay) MDA-15
Power consumption	2.3 kVA @ 230 VAC, 1.65 kVA @ 110 VAC
Maximum impedance of main	0,2 Ω

2D modalities

The following charts represent technique factors that can be used with the selected line voltage and continuous radiation. One of the three technique factors is always fixed.

Table 1: 100 VAC

mA											
16											
13	x	x	x								
10	x	x	x	x	x	x	x				
8	x	x	x	x	x	x	x	x	x	x	
6	x	x	x	x	x	x	x	x	x	x	
4	x	x	x	x	x	x	x	x	x	x	
	57	60	63	66	70	73	77	81	85	90	kV

Table 2: 120 VAC

mA											
16	x	x	x								
13	x	x	x	x	x	x	x	x			
10	x	x	x	x	x	x	x	x	x	x	
8	x	x	x	x	x	x	x	x	x	x	
6	x	x	x	x	x	x	x	x	x	x	
4	x	x	x	x	x	x	x	x	x	x	
	57	60	63	66	70	73	77	81	85	90	kV

Table 3: 240 VAC

mA											
16	x	x	x	x	x	x	x				
13	x	x	x	x	x	x	x	x	x	x	
10	x	x	x	x	x	x	x	x	x	x	
8	x	x	x	x	x	x	x	x	x	x	
6	x	x	x	x	x	x	x	x	x	x	
4	x	x	x	x	x	x	x	x	x	x	
	57	60	63	66	70	73	77	81	85	90	kV

3D modalities

The following charts represent technique factors that can be used with the selected line voltage in 3D imaging mode. 3D modality uses pulsed x-rays with fixed kV and exposure time.

Table 4:
100 VAC @ 90 kV
120 VAC @ 90 kV
240 VAC @ 90 kV

mA				
13	x	x	x	
10	x	x	x	x
8	x	x	x	x
6	x	x	x	x
4		x		x
	6 x 4 std res	6 x 4 high res	6 x 8 std res	6 x 8 high res

Table 5: (MFOV (Maxio) UNIT)
100 VAC @ 90 kV
120 VAC @ 90 kV
240 VAC @ 90 kV

mA													
12.5		x	x	x		x			x				
10		x	x	x		x	x		x	x		x	
8		x	x	x		x	x		x	x		x	x
6.3	x	x	x	x		x	x		x	x		x	x
5	x	x	x	x	x	x	x		x	x		x	x
4	x		x	x	x		x			x		x	x
3.2	x				x			x			x		x
	5 x 5 low dose	5 x 5 std res	5 x 5 high res	5 x 5 endo	6 x 8 low dose	6 x 8 std res	6 x 8 high res	8 x 8 low dose	8 x 8 std res	8 x 8 high res	8 x 15 low dose	8 x 15 std res	8 x 15 high res

Positioning laser lights	
Panoramic, TMJ & Maxillary Sinus Programs	laser light (CLASS 1 LASER PRODUCT) max output 100µW Warning symbols are placed next to the laser lights and the label describing the laser light classification is placed inside the carriage side cabinet. USA / Canada models have different types of laser light stickers according to local requirements. Caution - use of controls or adjustments or performance of procedures other than those specified herein may result in hazardous radiation exposure.
Cephalostat FH laser light	
3D imaging programs	
	IEC 60825-1:1993+A1:1997+A2:2001

X-ray generator	
Nominal power	1750 W nominal at 90 kV, 12 mA
Tube voltage	57 - 90 kV (+/- 5 kV)
Tube current	3,2 - 16 mA (+/- 1 mA)
Supply frequency	75 - 150 kHz
Spine compensation	kV / mA compensated
Spine compensation mode	Pre-programmed

User interface	
Program and technique factors selection, exposure control	Touch screen panel, optional remote exposure switch
Patient positioning	Positioning panel, integrated
Connection cable (OP300 - PC)	CAT6 UTP Ethernet cable

Panoramic programs & technique factors & magnification:		
Standard Adult Panoramic	57-90 kV/ 3.2-16 mA/16.4 s	30%
Pediatric Panoramic	57-90 kV/ 3.2-16 mA/14.4 s	30%
Ortho Zone	57-90 kV/ 3.2-16 mA/17.9 s	25%
Orthogonal Panoramic	57-90 kV/ 3.2-16 mA/12.9 s	30%
Wide Arch Panoramic	57-90 kV/ 3.2-16 mA/16.2 s	30%

Panoramic programs & technique factors & magnification:		
Ortho TMJ	57-90 kV/ 3.2-16 mA/10.6 s	23%
PA TMJ View	57-90 kV/ 3.2-16 mA/10.6 s	55%
Maxillary Sinus	57-90 kV/ 3.2-16 mA/12.5 s	30%
Bitewing	57-90 kV/ 3.2-16 mA/11.9 s	30%
Panoramic QC	57-90 kV/ 3.2-12.6 mA/16.4 s	30%

Exposure Control	Automatic Dose Rate Control (ADC) (P1-P5) Pre-programmed icons for all programs Automatic Spine Compensation	
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Cephalometric programs & technique factors:	
Pediatric lateral view	85-90 kV / 8-12.6 mA / 6.9-14 s
Lateral view	85-90 kV / 8-12.6 mA / 10-20 s
PA/AP, facial and oblique views	85-90 kV / 8-12.6 mA / 10-20 s
Carpus View (Not available in USA and Canada)	60-90 kV / 3.2-12.6 mA / 8-20 s
Exposure Control	Automatic Facial Contour (AFC), Pre-programmed icons for all programs.
Magnification factor	1.15 (15%)

3D MFOV (Maxio) imaging programs:	
50 x 50 mm FOV Low Dose	90 kV / 3.2 - 6.3 mA / 1.17 s
50 x 50 mm FOV standard resolution	90 kV / 5 - 12.5 mA / 2.34 s
50 x 50 mm FOV high resolution	90 kV / 4 - 12.5 mA / 17.4 s
50 x 50 mm FOV endo program	90 kV / 4 - 12.5 mA / 17.4 s
61 x 78 mm FOV Low Dose	90 kV / 3.2 - 5 mA / 1.17 s
61 x 78 mm FOV standard resolution	90 kV / 5 - 12.5 mA / 2.34 s
61 x 78 mm FOV high resolution	90 kV / 4 - 10 mA / 6.1 s
78 x 78 mm FOV Low Dose	90 kV / 3.2 mA / 1.17 s
78 x 78 mm FOV standard resolution	90 kV / 5 - 12.5 mA / 2.34 s
78 x 78 mm FOV high resolution	90 kV / 4 - 10 mA / 17.4 s

3D MFOV (Maxio) imaging programs:	
78 x 150 mm FOV Low Dose	90 kV / 3.2 mA / 2.25 s
78 x 150 mm FOV standard resolution	90 kV / 4 - 10 mA / 4.5 s
78 x 150 mm FOV high resolution	90 kV / 3.2 - 8 mA / 8.5 s
130 x 150 mm FOV Low Dose	90 kV / 3.2 mA / 4.5 s
130 x 150 mm FOV standard resolution	90 kV / 3.2 - 10 mA / 9 s
130 x 150 mm FOV high resolution	90 kV / 4 - 10 mA / 9 s
50 x 50 mm FOV scout	90 kV / 4 - 13 mA / 0.02 s
61 x 78 mm FOV scout	90 kV / 4 - 13 mA / 0.02 s
78 x 78 mm FOV scout	90 kV / 4 - 13 mA / 0.02 s
78 x 150 mm FOV scout	90 kV / 4 - 13 mA / 0.04 s
130 x 150 mm FOV scout	90 kV / 4 - 13 mA / 0.04 s

Image storing and retrieving:	
File formats	PNG (16-bit), JPG (12-bit)
File compression	PNG (lossless), JPG (100%-60% quality)
Panoramic file size	2-4 MB
Cephalometric file size	3-5 MB
3D file size	12-400 MB (DICOM)
Patient database	Standalone workstation Server on local area network (LAN)

Panoramic patient positioning	
Operation	Left or right side of unit Motorised carriage movement
Positioning aids	Chin rest, bite block, 3-point headrest Curved mirror, 3 positioning laser lights, Occlusion correction buttons

Cephalostat patient positioning	
Operation	Arm mounts on left or right side of the unit Interlocked pan/ceph sensor Motorised carriage buttons at cephalostat head assembly.

Cephalostat patient positioning	
Positioning aids	Ear holders, Nasion support with vertical mm scale, Frankfort horizontal plane laser light, Contact plate (Carpus program).

3D imaging patient positioning	
Operation	Left or right side of unit Motorised carriage movement
Positioning aids	Chin rest, chin support, 3-point headrest, Curved mirror, 3 positioning laser lights

Cephalostat scanning	
Scanning method	Horizontal scan, synchronized sensor and secondary slot motion
Scanning time	10 - 20 s.

Panoramic image receptor	
Sensor unit	Pan sensor or interchangeable Ceph sensor
Technology / Sensor type	CMOS
Image pixel size and depth	100 x 100 µm and 14 bits
Active area height	5.8 inches / 148 mm / 1480 pixels
Resolution	Pan: 5 LP/mm

Cephalometric image receptor	
Sensor unit	interchangeable Ceph sensor
Technology / Sensor type	CMOS
Image pixel size and depth	100 x 100 µm and 14 bits
Active area height	223,2 mm / 2232 pixels
Image field width in lateral view	10.2 inches / 260 mm, maximum 6.7 inches / 170 mm, minimum
Image field width in PA view	7.9 inches / 200 mm
Resolution	4 LP/mm (cephalometric)

receptor (3D SFOV unit)	
Sensor unit	3D sensor
Technology	CMOS
Image pixel size and depth	200 x 200 µm and 13 bits
Photodiode area:	100 x 68.2 mm

receptor (MFOV (Maxio) unit)	
Sensor unit	3D sensor
Technology / Sensor type	CMOS
Image pixel size and depth	200 x 200 µm and 13 bits
Photodiode area:	124,8 x 124,8 mm

Unit physical measures:	
source-image distance (SID)	500 mm (Panoramic) 570 mm (3D)
Installation	Standard wall mount with ±45° angled joint. Optional base for free standing unit (unit height is increased 25 mm).
Height x Width x Depth (inches/mm)	2410x830x1126mm (standard column) 94.9 x 32.7 x 44.3 inches -Max.
Weight	200 kg / 441 lbs. (Panoramic)

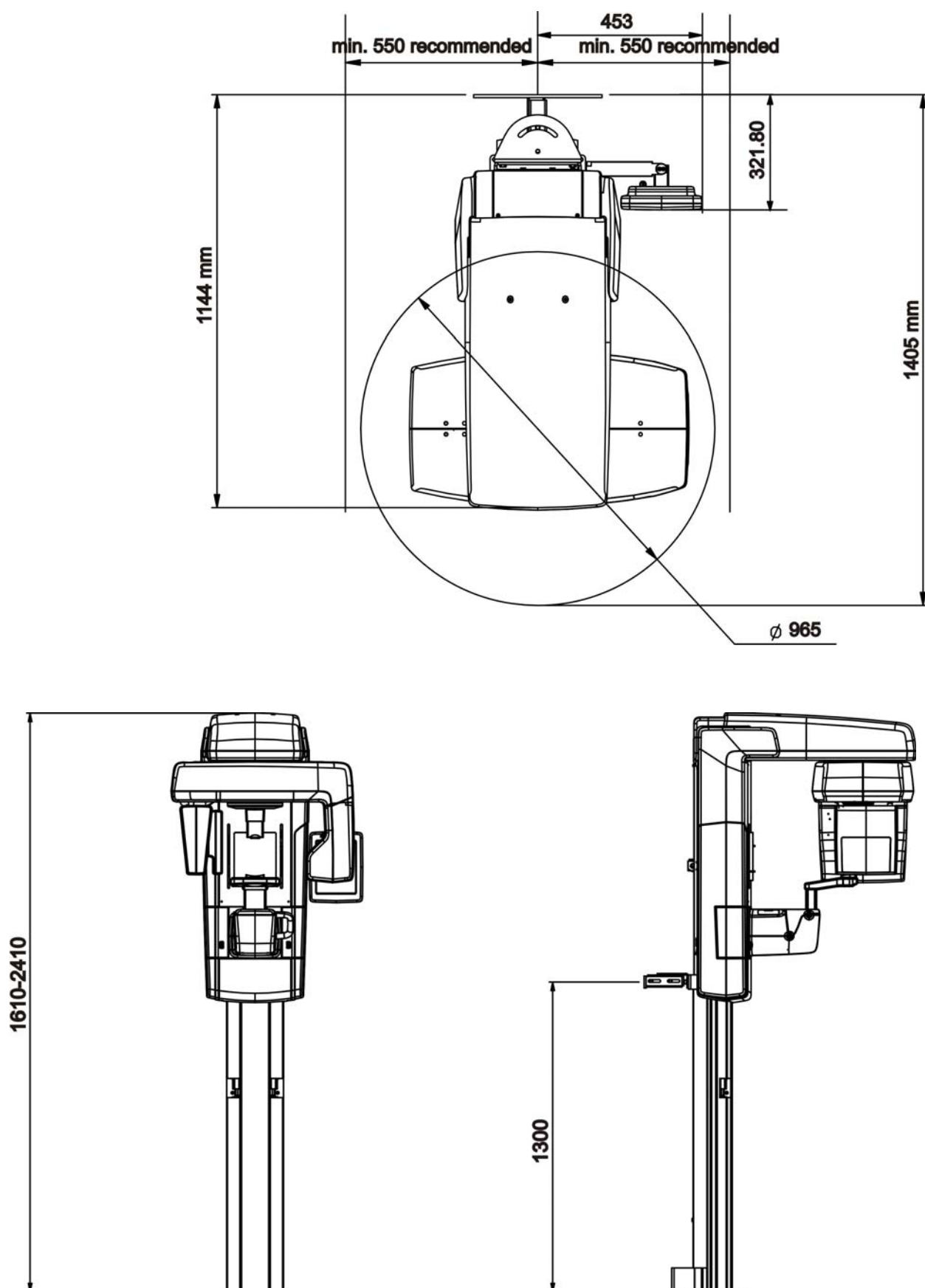
OP300 ceph physical measures:	
source-image distance (SID)	1745 mm / 68.7 inches
source-object distance (SOD)	1520 mm / 60 inches
Installation	Standard wall mount with 45° angled joint. Optional base for free standing unit (unit height is increased 25 mm)
Height x Width x Depth (inches/mm)	2410 x 1931 x 1193 mm 94.9 x 76 x 47 inches
Weight	240 kg / 529 lbs. (Cephalometric)

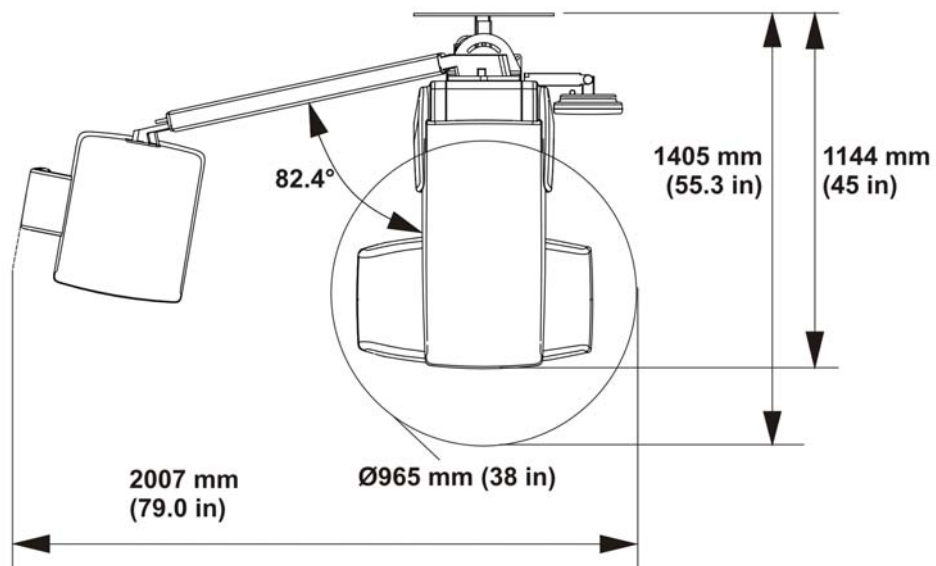
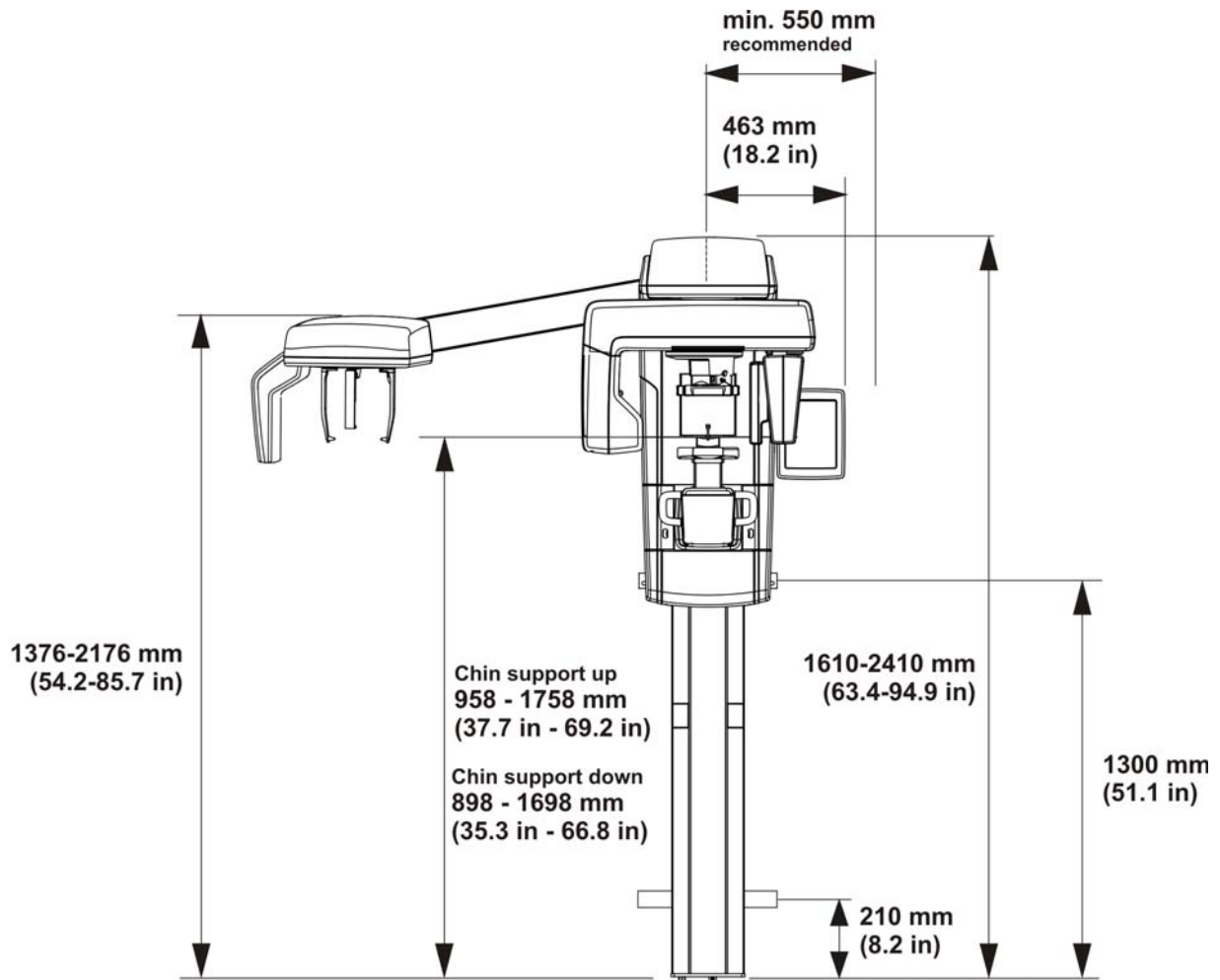
Ambient temperatures:	
Transportation and Storage	-10°...+60°C
Operation Temperature	+10°...+35°C, RH max. 85%

Ceph ready option (Ordered separately)	
Options	Description
Ceph Upgrade to OP300 pan	Unit has the same sensor as ceph unit. Cost saving with future digital ceph upgrade.

Configuration options for model OP300	
Upgrade	Description
Cephalostat Upgrade	Add ceph imaging to OP300 pan
Left / right	Change handedness to OP300 pan ceph

9.2 Unit dimensions





9.3 Symbols that appear in the unit



Radiation warning



Dangerous voltage



On or enabled



Off or disabled



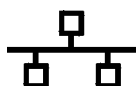
Exposure switch



Connector for exposure switch



CLASS II equipment
(Double insulated electrical appliance)



Ethernet connector



Operating instruction
(More details in operation instructions)



Laser radiation



Attention, consult accompanying documents



Ground (Functional)



Protective ground

CLASS 1 LASER PRODUCT

EN 60 825-1/A2:2001

Laser class label (Patient positioning lights)

Rx only

Caution: Federal law restricts this device to sale by or on the order of a licensed healthcare practitioner.



This symbol indicates that the waste of electrical and electronic equipment must not be disposed as unsorted municipal waste and must be collected separately. Please contact an authorized representative of the manufacturer for information concerning the decommissioning of your equipment.



Type B equipment



CE (0537) symbol
MDD 93/42/EEC



ETL Classified conforms to UL STD 60601-1.
Certified to CSA.

9.4 Labels on the unit

The main label of the unit is located on the vertical carriage next to the on/off power switch. The unit is class I, type B and with IP20 protection.



9.5 Electromagnetic Compatibility (EMC) tables

Note! Medical electrical equipment needs special precautions regarding EMC and needs to be installed according to EMC information.

Table 1.5 Electromagnetic emissions IEC 60601-1-2 Ed2

ORTHOPANTOMOGRAPH® OP300 is suitable for use in the specified electromagnetic environment. The purchaser or user of ORTHOPANTOMOGRAPH® OP300 should assure that it is used in an electromagnetic environment as described below:		
Emissions Test	Compliance	Electromagnetic Environment
Radio-Frequency Emissions CISPR11	Group 1	ORTHOPANTOMOGRAPH® OP300 uses RF energy only for its internal function. Therefore, the RF emission is very low and not likely to cause any interference in nearby electronic equipment.
Radio-Frequency Emissions CISPR11	Class B	ORTHOPANTOMOGRAPH® OP300 is suitable for use in all establishments, including domestic establishments and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.
Harmonic emissions IEC 61000-3-2	IEC 61000-3-2 Class A	ORTHOPANTOMOGRAPH® OP300 is suitable for use in all establishments, including domestic establishments and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.
Voltage fluctuations/ flicker emissions IEC 61000-3-3	Complies	ORTHOPANTOMOGRAPH® OP300 is suitable for use in all establishments, including domestic establishments and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.

Table 1.6 Electromagnetic immunity IEC 60601-1-2 Ed2

ORTHOPANTOMOGRAPH® OP300 is suitable for use in the specified electromagnetic environment. The purchaser or user of ORTHOPANTOMOGRAPH® OP300 should assure that it is used in an electromagnetic environment as described below:			
Immunity Test	IEC 60601-1-2 Test Level	Compliance Level	Electromagnetic Environment
Electrostatic discharge (ESD) IEC 61000-4-2	± 2, 4, 6 kV for contact discharge ± 2, 4, 8 kV for air discharge	± 2, 4, 6 kV for contact discharge ± 2, 4, 8 kV for air discharge	Floors are wood, concrete, or ceramic tile, or floors are covered with synthetic material and the relative humidity is at least 30 percent.
Electrical fast transient/burst IEC 61000-4-4	± 2 kV for power supply lines ± 1 kV for input/output lines	± 2 kV for power supply lines ± 1 kV for input/output lines	Mains power quality is that of a typical commercial and/or hospital environment
Surge IEC 61000-4-5	± 1 kV differential mode ± 2 kV common mode	± 1 kV differential mode ± 2 kV common mode	Mains power quality is that of a typical commercial and/or hospital environment.
Voltage dips, short interruptions and voltage variations on power supply input lines IEC 61000-4-11	< 5 % UT (> 95 % dip in UT) for 0,5 cycle 40 % UT (60 % dip in UT) for 5 cycles 70 % UT (30 % dip in UT) for 25 cycles < 5 % UT (> 95 % dip in UT)	< 5 % UT (> 95 % dip in UT) for 0,5 cycle 40 % UT (60 % dip in UT) for 5 cycles 70 % UT (30 % dip in UT) for 25 cycles < 5 % UT (> 95 % dip in UT)	Mains power quality is that of a typical commercial and/or hospital environment. If the user of ORTHOPANTOMOGRAPH® OP300 requires continued operation during power mains interruptions, it is recommended that ORTHOPANTOMOGRAPH® OP300 be powered from an uninterruptible power supply or a battery.
Power frequency (50/60 Hz) magnetic field IEC 61000-4-8	3 A/m	3 A/m	Power frequency magnetic fields are at levels characteristic of a typical location in a typical commercial and/or hospital environment.
NOTE: U_T is the a.c. mains voltage prior to application of the test level.			

Table 1.7 RF immunity of non-life-support equipment or system IEC 60601-1-2

ORTHOPANTOMOGRAPH® OP300 is suitable for use in the specified electromagnetic environment. The purchaser or user of ORTHOPANTOMOGRAPH® OP300 should assure that it is used in an electromagnetic environment as described below:			
Immunity Test	IEC 60601-1-2 Test Level	Compliance Level	Electromagnetic Environment
Conducted RF IEC 61000-4-6 Radiated RF IEC 61000-4-3	3 V 150 kHz to 80 MHz 3 V/m 80 MHz to 2,5 GHz	[V1] 3 V [E1] 3 V/m	Portable and mobile RF communications equipment are used no closer to any part of ORTHOPANTOMOGRAPH® OP300, including cables, than the recommended separation distance calculated from the equation appropriate for the frequency of the transmitter. Recommended Separation Distance: $d = \left[\frac{3,5}{V_1} \right] \sqrt{P}$ $d = \left[\frac{3,5}{E_1} \right] \sqrt{P} \quad 80 \text{ MHz to } 800 \text{ MHz}$ $d = \left[\frac{7}{E_1} \right] \sqrt{P} \quad 800 \text{ MHz to } 2,5 \text{ GHz}$ Where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and d is the recommended separation distance in meters (m). Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey,* are less than the compliance level in each frequency range.** Interference may occur in the vicinity of equipment marked with the following symbol: 

*Field strengths from fixed transmitters, such as base stations for cellular telephones and land mobile radios, amateur radio, AM and FM radio broadcast, and TV broadcast cannot be estimated accurately. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be performed. If the measured field strength exceeds the RF compliance level above, observe ORTHOPANTOMOGRAPH® OP300 to verify normal operation in each use location. If abnormal performance is observed, additional measures may be necessary, such as re-orienting or relocating ORTHOPANTOMOGRAPH® OP300.

Over the frequency range 150 kHz to 80 MHz, field strengths are less than [V1] V/m. **The Recommended Separation Distances are listed in the next table. **Note:** These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects, and people.

Note! RF communications equipment can effect medical electrical equipment.

Note! This equipment generates, uses and can radiate radio frequency energy. If not installed and used in accordance with this manual, it may cause harmful interference to radio communications. Portable and mobile RF communications equipment can also affect the performance of OP300.

Table 1.8 Table 4

Recommended Separation Distances for Portable and Mobile RF Communications Equipment IEC 60601-1-2			
Frequency of Transmitter	150KHz to 80 MHz	80 MHz to 800 MHz	800 MHz to 2,5 GHz
Equation	$d = [\frac{3,5}{V_1}] \sqrt{P}$	$d = [\frac{3,5}{E_1}] \sqrt{P}$	$d = [\frac{7}{E_1}] \sqrt{P}$
Rated Maximum Output Power of Transmitter (watts)	Separation Distance (meters)	Separation Distance (meters)	Separation Distance (meters)
0,01	0,12	0,12	0,23
0,1	0,37	0,37	0,74
1	1,17	1,17	2,34
10	3,69	3,69	7,38
100	11,67	11,67	23,34

USE LIMITATION:

External components

The use of accessories, transducers, and cables other than those specified may result in degraded ELECTROMAGNETIC COMPATIBILITY of the EQUIPMENT and/or SYSTEM

INSTALLATIONS REQUIREMENTS & ENVIRONMENT CONTROL:

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

Cables shielding & grounding

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

Electrostatic discharges environment & recommendations

In order to reduce electrostatic discharge interference, a charge dissipative floor should be installed to prevent charge accumulation.

- The dissipative floor material must be connected to the system reference ground, if applicable.
- Relative humidity must be maintained above 30 percent.

Stacked components & equipment

The ORTHOPANTOMOGRAPH® OP300 should not be used adjacent to or stacked with other equipment; if adjacent or stacked use is necessary, the ORTHOPANTOMOGRAPH® OP300 should be observed to verify normal operation in the configuration in which it will be used.

Interference may occur in the vicinity of equipment marked with the following symbol:



No portable or mobile RF communications equipment may be used closer to any part of the ORTHOPANTOMOGRAPH® OP300 including cables, than the recommended separation distance calculated from the equation appropriate to the frequency of the transmitter. See Table 4.

9.6 X-ray tube assemblies

Duty cycle 1:8

Rectification type: Constant potential x-ray generator

Generator rating: Generator nominal power 1750W

Figure 1.5

Anode Thermal Characteristics

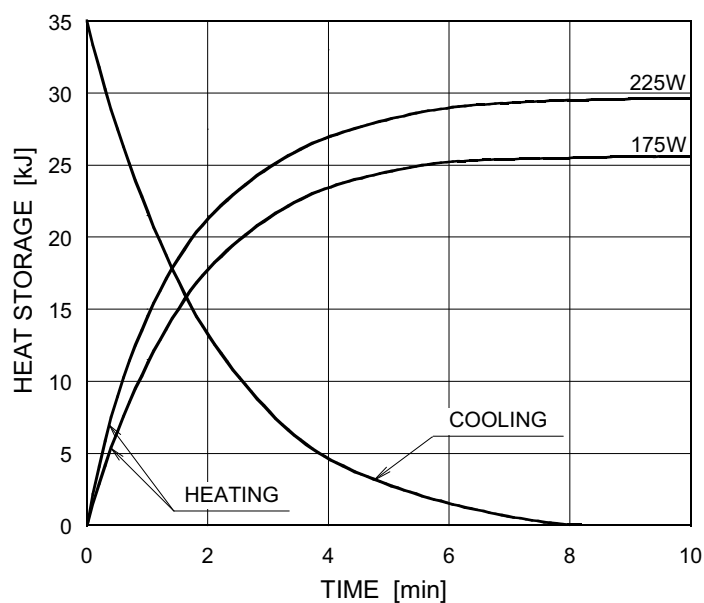


Figure 1.6

Maximum Rating Charts (Absolute maximum rating charts)

DC (Center-Grounded)

Focal Spot : 0.5 mm

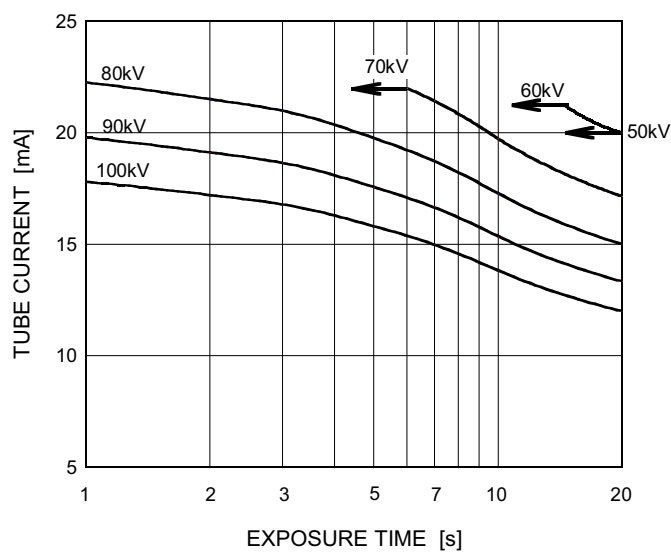
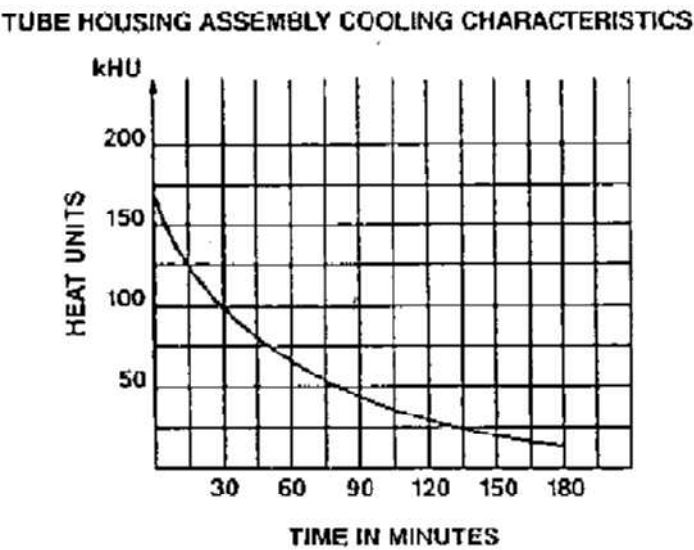


Figure 1.7



10 PC requirements

10.1 Minimum PC requirements

Minimum PC requirements for 3D acquisition workstation	
Processor	2.5 GHz dual core, or better
Memory	8 Gigabytes RAM, or more
Hard disk	500 GB, or more
Power supply	500 watt minimum
Network	Gigabit Ethernet 1000Base-T
Operating system	Windows 7, Windows Vista or Windows 8 (64-bit)
Display	20" LCD display, 1600 x 1200 or 22" LCD widescreen display, 1680 x 1050, or better
Standard	The PC must meet the IEC 60950 standard (minimum requirements)
Graphics board	The system is highly dependent on the used graphics board and graphics driver of the reconstruction workstation. Consult technical support for information on the supported boards and drivers. Examples of supported boards: NVidia Quadro 4000 NVidia GeForce GTX 660 The compatible graphics driver versions are found on the driver update DVD. For MFOV (Maxio) 2 GB dedicated display memory is required.
PCI board connection	Full-length PCIe x16 slot (for GPU board)
USB	USB ports (for HASP Dongle keys) • 1 for reconstruction system • 1 for 3D viewing SW (if needed)
Mouse	Mouse with scroll wheel

Minimum PC requirements for 2D acquisition workstation	
Processor	2.5 GHz dual core, or better
Memory	3 Gigabytes RAM, or more
Hard disk	500 GB, or more
Graphics board	NVidia GeForce 6600 or ATI Radeon X700 or better; 256MB or more memory (integrated graphics are not supported)
Power supply	500 watt minimum
Network	Gigabit Ethernet 1000Base-T
Operating system	Windows 7, Windows Vista or Windows 8 (32 or 64-bit)
Display	20" LCD display, 1600 x 1200 or 22" LCD wide screen display, 1680 x 1050, or better
Standard	The PC must meet the IEC 60950 standard (minimum requirements)
Mouse	Mouse with scroll wheel
OpenCL support	OpenCL 1.1 supported by either processor or graphics adapter

Note! This is an abbreviated list of requirements.
Please refer to the software installation manual or contact
your local dealer for detailed installation requirements.

Minimum PC requirements for 2D/3D Viewing workstation *	
Processor	2.0 GHz dual core, or better
Memory	3 Gigabytes RAM, or more
Graphics board	1GB or more memory (integrated graphics are not supported)
Hard disk	3 GB free space, or more
Network	Gigabit Ethernet 1000Base-T (recommended) or Fast Ethernet 100Base-TX
Operating system	Windows 7, Windows Vista or Windows 8 (32 or 64-bit)
Display	19" LCD display, 1280 x 1024, or better

* For the PC requirements of the 3D viewing software consult the 3D software manuals.

System requirements and connections

- The PC and any other external device(s) connected to the system must meet the IEC 60950 standard (minimum requirements). Devices that do not meet the IEC 60950 standard must not be connected to the system as they may pose a threat to operational safety.
- The PC and any other external devices must be connected in accordance with IEC 60601-1-1.
- The x-ray unit must be connected to its own separate power supply. The PC and any other external devices must NOT be connected to the same power supply as the x-ray unit.
- The unit shall be connected directly to the acquisition PC with an Ethernet cable. Connection through the LAN-network of the site is not allowed. Two network ports are needed in the PC in order to connect also to the site network.
- Position the PC and any other external device at least 1.5 m (60") from the xray unit so that the patient cannot touch the PC or any other external device while being x-rayed.
- The PC and any other external devices shall not be connected to an extension cable.
- Multiple extension cables shall not be used.
- Do not position the PC where it could be splashed

with liquids.

- Clean the PC in accordance with the manufacturer's instructions.
- The PC to be used with the unit must be installed in a location that meets all local and national safety requirements with regards the connection of a PC to an x-ray device.
- The connection of the unit to the PC must meet IEC 60601-1 requirements.
- The use of accessory equipment not complying with the equivalent safety requirements of this equipment may lead to a reduced level of safety of the resulting system.

Consideration relating to the choice shall include:

- Use of the accessory in the PATIENT VICINITY.
- The safety certification of the accessory has been performed in accordance to the appropriate IEC 601-1 and/or IEC 601-1-1 harmonized national standard.

10.2 The dental imaging software

The dental imaging software installed in the PC that is used with the unit must be MDD approved, for example CLINIVIEW™.