EzRay Air[™] Portable

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EzRay Air[™] Portable

User Manual

Model : VEX-P300 Version : 1.48

• English

Full Version





Notice

The EzRay Air Portable (Model: VEX-P300) is a portable dental X-ray system.

This manual contains descriptions, operational instructions, imaging procedures for the EzRay Air Portable (Model: VEX-P300) dental X-ray system. It is recommended that you thoroughly familiarize yourself with this manual to make the most effective use of this device. Observe all cautions, safety messages, and warnings that appear in this manual.

Always keep this manual with the device and review the operating procedures and safety instructions if needed.

The illustrations/photos of the device in this manual are only for illustration purposes. The actual device may differ.

Due to continuous technological improvements, the manual may not contain the most updated information. For further information not covered in this manual, please contact us at:

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This document is originally written in English.

The EzRay Air Portable is referred to as a device in this manual.

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The EzRay Air Portable (Model: VEX-P300) unit is intended to be used by a **certified professional**, exclusively for dental radiography.

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1. General and Regulatory Information

1.1 Manufacturer's Liability

The manufacturers and retailers of this device assume responsibility for the safe and normal operation only when:

- Always using genuine VATECH approved devices and parts.
- Performing all maintenance and repairs at a VATECH authorized distributor.
- Using this device properly following the user manual.
- The device damage or malfunction is not the result of an error on the owner or operator.

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EzRay Air Portable (Model: VEX-P300) is sold with the understanding that the user assumes sole responsibility for radiation safety (as well as any state, provincial, or local regulatory compliance) and that **VATECH**, its agents or representatives, do not accept responsibility for:

- 1) any injury or danger to personnel from X-ray exposure,
- 2) image over/underexposure due to poor operating techniques or procedures,
- 3) device which has been damaged, modified, or tampered with in any way.

1.2 Owner and Operator's Obligations

- This device owner must perform constancy tests at regular intervals to ensure patient and operator safety. Local X-ray safety regulations must perform these tests.
- This device owner must perform regular inspection and maintenance of the mechanical and electrical components to ensure safe and consistent operation (IEC 60601-1).
- This device owner must ensure inspection and cleaning work is performed by the maintenance schedule outlined in **Chapter 7 Cleaning and Maintenance**.



DO NOT operate this device until reading this manual and reviewed the related materials.

1.3 Conventions Used in this Manual

The following symbols are used throughout this manual. Make sure that fully understand each symbol and follow the instructions which accompany it.

Please observe all warnings and safety information included in this document to prevent personal injury and damage to the device.

	Indicates a potentially hazardous situation and improper handling may result in:
	Serious bodily injury (User or veterinary patient)
	Substantial property damage
	Indicates a potentially hazardous situation and improper handling may result in:
	Light injury
	Property damage
NOTICE	Indicates a potentially harmful situation, and improper handling may result in:
	Property damage
IMPORTANT	Indicates a usage and other valuable information.
RADIATION HAZARD	Indicates a possible danger from radiation exposure.

1.4 Marks and Symbols

The following table describes the purpose and location of safety symbols and other valuable information provided on the device.

Mark/Symbol	Description	Location
\sim	Alternate current	Battery Charger Label
	Direct current	Main Label
\triangle	Attention: Consult accompanying documents	Main Label
4	Dangerous voltage	Powerboard, X-ray Generator, Generator Label
OFF	Off (power: disconnect from the main switch)	Battery bay access door (outside)
ON	On (power: connect to the main switch)	Battery bay access door (outside)
*	IEC60601-1 Degree of Protection from Electric Shock TYPE B Equipment	Main Label
	Radiation hazard	Generator Label
EC REP	Authorized European Representative address	Main Label
CE 2460	The CE symbol indicates that this device complies with the European Directive for Medical Devices 93/42/EEC as amended by 2007/47/EC as a class IIb device.	Main Label
C U US E476672	UL mark No. E476672	Main Label

Dy Only	Caution: Federal law restricts this	
Rx Only	device to sale by or about a licensed healthcare practitioner.	Main Label
	Manufacturer's name and address	Main Label, Generator Label
\sim	Date of manufacture	Main Label, Generator Label
	This symbol indicates that electrical and electronic devices must not be disposed of as unsorted municipal waste and must be collected separately.	Main Label
	ESD susceptibility symbols indicate that an item is susceptible to damage from electrostatic discharges.	Board package
	Refer to the User Manual.	Main Label
	Using a Torx wrench, unscrew the battery bay access door.	Main Label
	Lift the door after the removal of the screw.	Main Label
OPEN CLOSE	This symbol indicates the direction of cover attachment/detachment.	Cone's upper part

1.4.1 Label Locations





1. Generator Label



2. Main Label



3. UDI Label





The labels' design and contents may be different in some countries based on local regulations and standards.

1.5 Standards and Regulations

Standards:

The device is designed and manufactured to meet the following standards:

MEDICAL - APPLIED ELECTROMAGNETIC RADIATION EQUIPMENT

AS TO ELECTRICAL SHOCK, FIRE, AND MECHANICAL HAZARDS ONLY IN ACCORDANCE WITH

ANSI/AAMI ES60601-1 (2005) + AMD 1 (2012),

CAN/CSA-C22.2 No. 60601-1 (2014), IEC 60601-1-3 (2008), IEC 60601-2-65(2012)

- ISO 13485
- 21 CFR 1020.30 & 1020.31

C E 2460	This is Class IIb equipment and obtained CE marking in April 2007 for regulation compliance by the revised European Union's MDD (Medical Devices Directive) 93/42 EEC.
CULUS E476672	This equipment received the UL certification mark in accordance with ANSI/AAMI, CAN/CSA-C22.2 No. 60601-1 regulations.

Classifications (IEC60601-1 6.1):

Protection against the ingress of water: Ordinary Equipment (IPX0)

Protection against electric shock: Class II equipment,

Type B Applied Parts: Backscatter Shield, Round Cover, Rectangular Cover, Rotating Rectangular Cover



2. Safety Instructions

2.1 General Safety Guidelines

	 This device is designed and manufactured to ensure the most significant safety of operation. Operate and maintain it in strict compliance with the safety precautions and operating instructions contained in this manual. Only a legally qualified person, i.e., a radiographer, diagnostic radiographers, medical radiation technologists, or healthcare professionals who specialize in imaging medical for the diagnosis and treatment of pathology, can operate this equipment in a controlled environment.
	 Observe all local fire regulations. Always keep a fire extinguisher near the device.
	• The device must be installed, maintained, and serviced by qualified service personnel according to the procedures and preventive maintenance schedules. Users can perform an only battery replacement.
	• Ensure that the on/off switch is set to off when the device is not in use.
	 Always disconnect the power supply before cleaning the device.
	 DO NOT keep the device or its parts in a humid place or a liquid substance.
	 Avoid placing the device near chemical storage and gas- filled storage facilities.
	• The backscatter shield protects users from backscatter radiation that they might be exposed to during X-ray exposure. Operating the device with the backscatter shield allows the users to be exposed to less radiation than when operating without. For details on the scattering data of the device configured both with and without the backscatter shield, please review 10.4.3 Scattered Dose .
	 Wireless communication equipment is used no closer than 30 cm (12 inches) to any part of the EzRay Air Portable(Model: VEX-P300), including cables specified by VATECH.
IMPORTANT	This device is shipped with the backscatter shield attached (firmly fixed), so users can not replace the backscatter shield. If it is damaged or defective, contact your Service Representative for replacement.
NOTICE	Mode of operation: N on -continuous operation with cyclic loading—This device needs a rest time of at least 60 times the exposure time before starting the next exposure.

2.2 Warnings and Safety Instructions

	 This X-ray unit may be dangerous to patients and operators unless safe exposure factors, operating instructions, and maintenance schedules are observed. It is essential to read this user manual carefully and strictly abide by all warnings and cautions stated. To avoid the risk of electric shock, this device must only be connected to the supply main with protective earth. Since rules and regulations concerning radiation safety differ between countries, it is the responsibility of the owner and operator of this device to comply with all applicable rules and regulations concerning radiation safety and protection in their area. DO NOT open or remove the cover panels on this device. Never expose this device to liquids, mists, or sprays. Exposing this device to liquids may cause an electrical shock or otherwise damage the system. DO NOT use spray cleaners on this device, as this could cause a fire. Never use this device in an environment that is susceptible to explosion. DO NOT place flammable materials near this device. Never touch the patient while also touching the SIP/SOP connectors. The medical electrical device is subject to special EMC preventive measures. For more details, refer to 10.5
RADIATION HAZARD	 Patients and operators are encouraged to wear leaded protective lead-lined aprons unless other Radiation Protection Protocols are applied locally. Children and pregnant women must consult with a doctor before X-ray exposure.
IMPORTANT	 Never try to modify this device, including the wires or cables. Modifying this device may damage it beyond repair. Serious dangers may occur from electromagnetic interference (i.e., noise) between other devices in the area during specific examinations or medical treatment.
NOTICE	Battery chargers must be in an accessible area where they can be easily unplugged from the power source.

	 Make sure to charge the battery in the external environment from the patient.
	 Make sure to use the battery only provided or approved by VATECH. If non-standard or damaged batteries are used, there is a risk of fire and explosion.
	 Make sure to use the battery charger only provided or approved by VATECH. Using an unauthorized charger may result in battery damage.
	 DO NOT expose batteries to heat or fire. Avoid storage in direct sunlight.
	 DO NOT short-circuit, crush, puncture, mutilate, or disassemble the battery.
A	 DO NOT store batteries haphazardly in a box or drawer where they may short-circuit each other or be short- circuited by other metal objects.
	 Observe the plus (+) and minus (-) marks on the battery and device and ensure correct use.
	 DO NOT subject batteries to mechanical shock.
	 In a cell leaking event, do not allow the liquid to meet the skin or eyes. If the contact has been made, wash the affected area with copious amounts of water, and seek medical advice.
	 Keep the battery away from children and pets.
	 DO NOT make the battery wet or let it be in the water. Keep batteries clean and dry.
	 Seek medical advice immediately if a battery has been swallowed.
	 Make sure to turn off the device before replacing the battery.
	 DO NOT remove a battery from its original packaging until required for use.
	 DO NOT dispose of batteries with ordinary trash. Turn in discharged batteries to local supply or discard or recycle batteries according to your local government regulations.
	 DO NOT leave a battery on a prolonged charge when not in use.
IMPORTANT	 If the device has not been used for long periods, it is recommended to charge the battery before use.
	 After extended periods of storage, it may be necessary to charge and discharge the cells or batteries several times to obtain maximum performance.

IMPORTANT	 If the device not in use has been turned on for long periods, the battery may be fully discharged. Depending upon the battery discharge status, it takes about 1 day for charging the battery. If the device is not turned on after charging the battery for about 1 day, it indicates that the battery has been fully discharged. Contact your Service Representative for battery replacement. DO NOT charge a fully discharged battery, as this may cause fire or explosion. Be sure to replace the battery (provided by VATECH).
NOTICE	 Users can replace batteries. When charging the battery, the exposure function is locked. Be sure to turn off the device when not in use. This helps to ensure the life of the battery. Be sure to charge the battery frequently. This helps to ensure the life of the battery.

Radiation Safety

	 When using the device, it is recommended that all users comply with the following radiation safety guidelines for the users and the patients' safety.
	 All users and patients should wear a protective device, such as a lead apron, thyroid collar, etc. This device should be operated in an area that is more than 6 feet away from other personnel, such as assistants or other patients. If they should stay closer than 6 feet, it is recommended that they wear a lead apron, thyroid collar, or stay behind a lead shield.
RADIATION	 Pregnant women should not be exposed to X-rays unless it is strictly necessary.
HAZARD	 All users should comply with the Radiation Protection Policies established by the government.
	 When selecting a Position Indicating Device, it should be considered if the PID can be used with the backscatter shield attached at the cone's outer end for most operator protection.

3. System Overview

3. System Overview

The device, a portable dental X-ray system, operates on 21.6 V_{dc} supplied by a rechargeable Li-ion battery pack. The portable X-ray system is an X-ray generating device designed for dental examination (teeth and jaw). The portable X-ray system is composed of an X-ray generating part with an X-ray tube including a device controller, a power controller, a user interface, a beam limiting part, a backscatter shield, and an optional Remote Exposure Switch. The device is designed to diagnose teeth and jaw through X-ray exposure using intra-oral image receptors.

3.1 Indications for Use

EzRay Air Portable (Model: VEX-P300) is an extraoral diagnostic dental X-ray source to produce X-ray images using intraoral image receptors. It is indicated for use by a dentist or a dental technician for both adult and pediatric patients.

3.2 Principles of Operation

X-rays are emitted when a high voltage is supplied to the X-ray tube assembly, which frees electrons from the cathode. They hit anode to produce X-rays. The device acquires images by emitting X-rays continuously on the human tooth.

Considerations	Requirement Description
Education	A licensed dentist or dental hygiene, radiologist, and graduates of relevant bachelor's degree (national qualifications)
Knowledge	 The operator must have understood: treatment and diagnosis of dental disease terms and guidance of diagnostic medical radiation devices device connection, installation, and operating conditions.
Language understanding	The operator must have understood:the English or Korean manuals (or other languages provided).
Experience	 The operator must have understood: objectives and effects of treatment and diagnosis of dental disease using diagnostic medical radiation devices normal operation of diagnostic medical radiation devices the contents of the user manual.

3.3 Intended User Profile

ENGLISH

3.4 Components



No.	Item	Standard	Option	Qty.
1	Main Body (included the Backscatter Shield and the Round Cover)	•		1
2	Hand/Neck Strap	•		1
3	Rectangular Cover (4x3)	•		1
4	power cord	•		1
5	Battery Charger	•		1
6	User Manual	•		1
7	Cradle		•	1
8	Rectangular Cover (2x3)		•	1
9	Rotating Rectangular Cover (4x3)		•	1
10	Rotating Rectangular Cover (2x3)		•	1
11	Remote Exposure Switch		•*	1
12	Base Holder		•*	1
13	Tripod		•*	1

**Read the notice on the next page for the options with the symbol (*)

The remote exposure switch, base holder, and tripod are provided with **EzRay Air Portable (Model: VEX-P300)** for users in the countries where a hand-held X-ray device is prohibited in the clinic. For more information, please contact a VATECH representative in your area.

3.5 Features

The device is an intra-oral portable X-ray system that offers safety, reliability, and greater functionality:

- Lightweight and ergonomic design
- The convenience of cordless design by using battery-pack
- Micro-computer and a specialized circuit that monitors and precisely regulates the exposure technique factors (kV, mA, and exposure time)
- Pre-programmed exposure time makes the operation fast and easy.
- A selection of phosphor plates or digital sensors

3.6 General View of the Device

Main Body



No.	ltem	Description
1	Backscatter Shield	Shields from the backscattering radiation.
2	X-ray Beam Limiting Device	Limits the X-ray exposure area. Default type: Round Cone + Round Cover (FOV: Ø 6 cm)
3	X-ray Exposure Button	Press the button for X-ray exposure.
4	Handle	Grip the handle securely when using the system.
5	Power Button	Power On/Off button
6	Battery	Rechargeable Lithium-ion battery
7	Adaptor Connector	Connect the charging adaptor.
8	Remote X-ray Exposure Switch Port	Connect the X-ray exposure cable switch. Alternatively, it can be used as a service port
9	Strap Loop	Connect the strap.
10	Control panel	Display for the X-ray exposure settings and operation conditions
11	X-ray Generator	Includes the X-ray tube and the high-voltage generator.

Control panel



No.	ŀ	tem	Description
1	65kVp / 2.5mA	Tube Voltage/Current Indicator	Indicates the tube voltage and tube current of the system.
2	≈8.88 °	Angle/Time Display	Displays the X-ray exposure time, error code, cooling time, and exposure angle.
3	🖞 សិ	Adult/Child Selection	Indicates a patient type (adult or child).
4		Remaining Battery Indicator	Indicates remaining battery level.
	4	Battery Charging Indicator	 1) Indicates that the battery charger is connected to the device. 2) Indicates the battery needs to be charged when it flickers.
5	SET	SET Button	Resets the X-ray exposure angle.
6		Speaker	Sound alarm for the X-ray exposure
7		Tooth Type Selection	Selects the tooth type.

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8		X-ray Exposure Indicator	Indicates the X-ray exposure status. (Green: Ready / Yellow: X-ray On)
9		Jog Dial	Turn the jog dial left (-) or right (+) to select an X-ray exposure setting, press the jog dial to confirm the operating setting.
			Available only in countries that allow the auto mode for use.
10	Αυτο	AUTO Button	The mode is not available in the United Kingdom, the United States, and some countries. (Consult the sales representative in your country for details)
			In these countries, only "NA" is displayed on the panel when this button is pressed.

Available Option Items

No.	Figure	Option name
1		Rectangular Cover 2x3 (3x2) FOV: 2x3 cm, 3x2 cm (This cover can be used as both 2x3 and 3x2.)
2		Remote Exposure Switch
3		Rotating Rectangular Cover 4x3 (3x4) FOV: 4x3cm, 3x4cm (This adaptor can be used as both 4x3 and 3x4.)
4		Rotating Rectangular Cover 2x3 (3x2) FOV: 2x3cm, 3x2cm (This adaptor can be used as both 2x3 and 3x2.)
5		Base Holder*
6		Cradle
7		Tripod

* In case a tripod is used with the Base Holder, refer to the specifications below.

- Fixing bolt size: 3/8 inch
- Maximum supportable weight: about 5 kg
- Minimum height: > 130 cm
- · More than 3 columns are required.
- When using a tripod with 3 columns, make sure to have space of at least 1 min-width on the bottom.



Refer to **Appendix 10.2 Using the Rotating Rectangular Cover** for instructions for use.

4. Operation

4.1 Power On/Off

1. Turn on the system referring to the following figure.



- 2. The following displays and indicators light up:
 - Current Angle/Time display
 - Tooth type selection display
 - Adult/Child selection display
 - Remaining battery indicator
 - X-ray exposure indicator
- 3. Make sure that at least one battery indicator light comes on.



Battery level 1



4.2 Enter Password



To turn on or off the password mode, refer to **5.3.4. Password mode On/Off**. To change the password, refer to **5.3.11. Password Setting**.

3

1. Enter the 3 digits numeric password using the jog dial. (default password: 000)



2. Turn the jog dial to choose a number and press the dial to move to the next digit.



3. When all three digits are set, press the jog dial again to save the setting.

4.3 **Operation Mode**

This system can be operated in Manual Mode.

Manual Mode

1. When the tooth type selection area flickers, turn the jog dial to select the tooth type. To see the Control panel before and after selection, refer to the figures below.



Before tooth type selection

Tooth Type

Symbol	Туре
\heartsuit	Incisor
\heartsuit	Canine
\square	Molar/Premolar
	Bitewing

2. After tooth type selection, a patient type should be selected. When the Adult/Child selection area flickers, turn the jog dial to select the patient type. To see the Control panel after selection, refer to the figure below.



After patient type selection

Patient Type





After the tooth type and patient type are selected, the exposure time is automatically displayed.

3. When changing the exposure time, turn the jog dial to adjust the exposure time from 0.05 to 1.0 s. (increments: 0.01 s)

NOTICE	When pressing the jog dial after adjusting the exposure time in Manual Mode, the exposure time is returned to the default setting.
	To save the exposure time as default in Manual Mode, press and hold the jog dial for about 3 seconds.

4.4 Positioning

Positioning the Patient

To obtain high-quality intra-oral radiography with maximum details, take extra care in all steps of the radiography process: positioning the patient and the X-ray imaging system; exposing the phosphor plate or the intra-oral sensor.

- 1. Place a protective lead apron on the patient's chest.
- 2. Have the patient sit on the chair with the sagittal plane vertical.
 - For radiography of the upper maxillary, the Frankfort plane must be horizontal.



• For radiography of the lower maxillary, the Occlusal plane must be horizontal.

Occlusal plane	J
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- 3. Place the tube head cone in the area want to take an image.
 - 1) When using the device as a hand-held

When holding the device, it is recommended to grip the handle by one hand and place the other on the underside of the device, as shown in the following figure.



2) When using the device with a tripod

If you are in a country that does not allow using a hand-held X-ray in the clinic, use the device with a tripod, base holder, and remote exposure switch. To learn the proper use of each of these components, see **4.5.1 Remote Exposure Switch** and **10.1 Combining the Holder and the device using a tripod**.



Depending on the imaging angles, exposure times vary. Since it is necessary to keep the patient with low X-ray doses and the user in the protected area, have the patient's head slightly tilted, and raise or lower the chin if needed. Please refer to **2.2 Warnings and Safety Instructions**.

For further information about the patient positioning and beam angle for each mode, refer to the following **Positioning Instructions**.

Positioning Instructions

Paralleling technique: The phosphor plate or sensor is placed in a holder used to align the phosphor plate or sensor parallel to the tooth' long axis.



Bisected angle technique: The patient holds the phosphor plate or sensor in place with his/her finger. The X-ray beam is directed perpendicularly towards an imaginary line, which bisects the angle between the phosphor plate or sensor plane and the tooth's long axis.



Position the tube head to the patient using the accepted standard positioning procedures.

Here are the specific angulations and directions for the tube head to take the best images of a tooth (i.e., **Bisected angle technique**).

CAUTION Position the receptor carefully not to damage the soft tissue of the patient's intra-oral area.

Maxillary Incisor

The x-ray beam is directed downward at 45°.



Teeth		Angle of inclination
Incisor	Maxilla	+45°

Mandibular Incisor

The x-ray beam is directed upward at 25°.



Teeth		Angle of inclination
Incisor	Mandible	-25°

Maxillary Canine

The x-ray beam is directed downward at 45°.



Тее	th	Angle of inclination
Canine	Maxilla	+45°

Mandibular Canine

The x-ray beam is directed upward at 20°.



Teeth		Angle of inclination
Canine	Mandible	-20°

Maxillary Molar and Premolar

The x-ray beam is directed downward at 30°.



Teeth		Angle of inclination
Molar and Premolar	Maxilla	+30°

Mandibular Molar and Premolar

The x-ray beam is directed upward at 5°.



Teeth		Angle of inclination
Molar and Premolar	Mandible	-5°
Bitewing

For a bitewing exposure, the patient closes their teeth during exposure on the phosphor plate/sensor holder.

The x-ray beam is directed downward at $5^{\circ} \sim 8^{\circ}$



Teeth	Angle of inclination
Bitewing exposure	+5°~ +8°

Positioning the Imaging Sensor

Using **EzRay Air Portable(Model: VEX-P300)**, a portable dental X-ray system, you may create an X-ray image on different types of imaging receptors:

- · Digital sensors
- Phosphor plate

To ensure image quality, the digital imaging sensor must be appropriately positioned (for information about the imaging sensor's proper placement, please refer to 'Positioning Instructions.'

 Failure to position the imaging sensor correctly can result in errors on the radiograph, such as distorted teeth and roots, elongation, magnification, and overlapping contacts.



The paralleling technique generally reduces the risk of such errors, but if the sensor's position improperly, angulation errors may occur (angulation of the sensor to the tooth itself).

 Failure to align the imaging sensor with the X-ray beam's exit pattern can result in cone-cuts on the radiograph. The cone-cuts are bright areas shown on the radiograph when part of the radiograph is not exposed to radiation. Please refer to the following figure as an example of cone-cuts.



The following figure indicates how the cone-cut occurred by showing the imaging sensor's position and the radiation area.



To ensure proper alignment between the imaging sensor and the X-ray beam, it is recommended to use a PID (Position Indicating Device), as shown in the following figure.



When using the PID, the X-ray device's exit pattern should be aligned perpendicular to the target receptor, as shown in the following figure.





Once the PID is appropriately aligned, instruct the patient not to move.

4.5 Exposure

IMPORTANT

The operator **MUST** instruct the patient to refrain from moving during the entire exposure.

- 1. Instruct the patient not to move.
- 2. Press the Exposure Button for exposure duration.



IMPORTANT	The operator in the countries where a hand-held X-ray device is prohibited must use a remote exposure switch.
	See 4.5.1 Remote Exposure Switch for details.

- 3. While X-ray is being exposed,
 - The X-ray Exposure Indicator lights up, and an audible sound is produced.
 - Keep pressing until the X-ray Exposure Indicator light goes out, and the audible sound stops.



4.5.1 Remote Exposure Switch

The **Remote Exposure Switch** allows the operator to control image acquisition from outside of the X-ray room.

Press and hold the **Remote Exposure Switch** until the acquisition is completed. Premature release of the **Remote Exposure Switch** will abort image acquisition.

Pressing the **Remote Exposure Switch** activates the X-ray Exposure Indicator to turn yellow. This color indicates that the X-ray is being emitted.





4.6 Using the Battery

The battery level indicator with a residual quantity is shown on the left side of the Control panel. When the battery indicator has one flickering light (level 1), charge the battery immediately. The figure below shows each battery indicator from level 1 to 3

Level 3



When the battery reaches Level 1, charge the battery at least for an hour to avoid a low voltage. All displays are turned off under the low voltage except for the error code **A.10**, as shown (see the figure below).

After the device is connected to the battery charger, and the battery level becomes 'Level 1', all functions are returned to normal operation.



System Status depending on Battery Levels

ltem		System Status			
		Battery Level 3, 2	Battery Level 1		Low Battery
	Operation	Normal	Normal	Normal	Not operated
When	Battery Level Indicator	Normal	Normal	Flickers	Not displayed
turning on the system	Battery Charging Indicator	Not displayed	Not displayed	Flickers	Not displayed
	Control panel Brightness	Normal	Normal	Dark	Normal (Error code A.10 is only displayed)
	Operation	Normal	Normal	Normal	Not operated
When	Battery Level Indicator	Normal	Normal	Flickers	Not displayed
operating the system	Battery Charging Indicator	Not displayed	Not displayed	Flickers	Not displayed
	Control panel Brightness	Normal	Normal	Normal	Normal (Error code A.10 is only displayed)

	Make sure to charge the battery if the Battery Charging Indicator flickers (or the error code A.10 is displayed). If the device has been turned on for long periods with the error code A.10, the battery may be discharged.
	DO NOT charge a fully discharged battery, as this may cause fire or explosion. Be sure to replace the battery (provided by VATECH).
NOTICE	When the device is connected to the battery charger, the Battery Charging Indicator is always displayed except that the battery is fully discharged.

Charging the Battery

1. Connect the battery charger to the battery charger connector, as shown in the following figure.



2. When the battery charger is connected, the battery charging LED indicator light comes on. Charge the battery until all the three LED indicators are filled up.





Usually, it takes about 3 hours to charge the battery after a complete discharge fully.

3. When the battery charge is completed, remove the battery charger from the device.



Battery Use Cycle

The battery is a consumable part. It is expected to degrade gradually, so it should be recharged more frequently. When the battery duration decreases to half or less than half compared to when the battery was new, contact your Service Representative to get a new battery.

To check how to replace the battery, see 'Battery Replacement.'

Battery Replacement



Users can replace batteries. To replace the battery, contact your Service Representative to get a battery kit (including a new battery and a Phillips screwdriver).

1. Workforce

One person, 3 minutes

2. Tools required

Phillips screwdriver (size: T20)

- 3. Removal Procedure
 - 1) Using a Phillips screwdriver, unscrew the battery bay access door.



2) Lift the door and remove the battery.



3) Disconnect the device's battery cable by pressing the battery cable connector, as shown in the following figure.



After disconnecting the battery cable





DO NOT pull excessively on the battery cable.

4) Install the new battery in the reverse order of removal.

Sleep Mode 1

To minimize battery consumption, Sleep Mode 1 starts when the device stays inactive for one minute (To learn how to change the default waiting time for Sleep Mode 1, see **5.3.6 Waiting Time Setting for the Sleep Mode 1**).

1. When Sleep Mode 1 starts, the Control panel becomes lightly dark, as shown in the right figure.



Brightness: Normal

Brightness: Lightly dark

 To return to normal operation, press any button on the device (except for the Xray Exposure Button).



Sleep Mode 2

Sleep Mode 2 starts when the device stays inactive for 5 minutes after entering Sleep Mode 1 (To change the default waiting time for Sleep Mode 2, see **5.3.7 Waiting Time Setting for the Sleep Mode 2**).

If the battery charger is connected to the device, Sleep Mode 1 is maintained.

1. When the Sleep Mode 2 starts, all displays are turned off on the Control panel, as shown in the right figure.



Brightness: Lightly dark

Brightness: Dark

2. To return to normal operation, press any button on the device (except for the X-ray Exposure Button).



Power Down Mode

When the device stays inactive for 5 hours after entering Sleep Mode 2, the Power Down Mode starts. (To change the default waiting time for the Power Down Mode, see **5.3.8 Waiting Time Setting for the Power Down Mode**).

IMPORTANT

If the battery charger is connected to the device, Sleep Mode 1 is maintained.

1. All displays are turned off on the Control panel.



2. To return to normal operation in the Power Down Mode, **MUST** turn off the system and turn it back on.





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5. Service Mode

5.1 Overview

In the Service Mode, users can check and change the following settings:

- 5.3.1 Factory Default Settings
- 5.3.2 Exposure Time Settings
- 5.3.3 User Default Settings (for each patient and tooth type))
- 5.3.4 Password Mode On/Off
- 5.3.5 Angle Increments Setting
- 5.3.6 Waiting Time Setting for the Sleep Mode 1
- 5.3.7 Waiting Time Setting for the Sleep Mode 2
- 5.3.8 Waiting Time Setting for the Power Down Mode
- 5.3.9 Buzzer On/Off
- 5.3.10 Power Down Mode On/Off
- 5.3.11 Password Setting

5.2 Changing System Parameters

To change system parameters:

1. Press and hold the **SET button** and **jog dial** simultaneously (for about 3 seconds).



2. Enter the 3 digits numeric password using the jog dial. (default password: 000)



3. Turn the dial to choose a number and press it to move to the next digit.



- 4. When all three digits are set, press the jog dial again to save the setting.
- 5. After entering the password is completed, the service mode starts, as shown in the figure below.



 The system provides 24 service modes. To change each mode's settings from n.01 to n.24, press the AUTO button on the left. Upon pressing the button, n.01 is displayed on the control panel.



- To move to the next mode, press the AUTO button again. Each time you press the button, the service mode number goes up by one. To return to the previous move, press the SET button on the right. (See 5.3 Service Mode Menu to learn more information on each mode)
- 8. Each time you finish changing the system parameter, press the **jog dial** and hold it until you hear a beep.



9. To resume the operation, press the **SET button** and the jog dial simultaneously and hold them for about 3 seconds.



10. Restart the device to check if the change is saved correctly

5.3 Service Mode Menu

See 5.2 Changing System Parameters for entering each service mode.

5.3.1 Factory Default Settings

Factory default settings restore all system mode parameters to their default values except for **n.24**. Follow the procedures below to activate the settings.

Service Mode No.	Item
n.01	Factory default settings

To restore the system to the factory default settings,

1. Select **n.01** and press the jog dial. When the message "**YES**" is displayed on the control panel, the system is restored to its initial settings.



- 2. To save the setting, press the jog dial and hold it until you hear a beep.
- 3. Restart the device to check if the factory default setting is on.

5.3.2 Exposure Time Settings (for each patient and tooth type)

Service Mode No.	Item
n.02	Adult Incisor
n.03	Adult Canine
n.04	Adult Molar/ Premolar
n.05	Adult Bitewing
n.06	Child Incisor
n.07	Child Canine
n.08	Child Molar/ Premolar
n.09	Child Bitewing

Select a service mode from **n.02** to **n.09** to set up an exposure time for each patient and tooth type. See the table below for more information.

To set the exposure time for a specific patient and tooth type as described above,

1. Select the Service Mode number according to the patient and tooth type (i.e., **n.06** for Child Incisor)



- 2. After entering the mode, turn the jog dial to adjust the exposure time.
- 3. Press the jog dial again and hold it until you hear a beep.
- 4. Restart the device to check if the new setting is saved.

5.3.3. User Default Settings (for each patient and tooth type)

Service Mode No.	Item
n.10	Adult
n.11	Child
n.12	Incisor
n.13	Canine
n.14	Molar/Premolar
n.15	Bitewing

User default settings are activated as the device starts. See the table to change each service mode number aligned with the patient or tooth type.

To set the default setting for a specific patient type or a tooth type,

 Select a service mode number between n.10 (Adult) and n.11 (Child) for a specific patient type and from n.12 to n.15 for a tooth type. For instance, when select n.10 by pressing the jog dial to set the patient type 'adult' as default, then "YES" is displayed on the Control panel.



- 2. Press the jog dial again and hold it until you hear a beep.
- 3. Restart the device to check if the new setting is saved correctly.

5.3.4 Password Mode On/Off

Follow the procedure below to turn on and off the password mode.

Service Mode No.	Item
n.16	Password On
n.17	Password Off

To activate the password mode,

1. Select **n.16** and press the jog dial. The message "**YES**" is displayed on the Control panel.



- 2. Press the dial again and hold it until you hear a beep.
- 3. Restart the device to check if the setting is saved correctly.
- 4. To deactivate the password mode, select **n.17**, and repeat the procedure.

5.3.5 Angle Increments Setting

	,
IMPORTANT	Available only in countries that allow the auto mode for use. The mode is not available in the United Kingdom, the United States, and some countries (Consult the sales representative in your country for details). When the auto button is pressed, only "NA" is displayed on the panel.

Follow the procedure below to set up the angle increments for the Auto Mode.

Service Mode No.	Item
n.18	Angle increments setting

When using the Auto Mode, the angle value increases and decreases according to the increments setting (default: 1 degree).

The angle increments can be set from 1 to 5 degrees.

To change the angle increments setting,

1. Select **n.18** and press the jog dial. The default angle increments "**001**°" is displayed on the control panel, as shown in the right figure.



- 2. Turn the jog dial to adjust the angle increments you want.
- 3. Press the dial again and hold it until you hear a beep.
- 4. Restart the device to check if the setting is saved correctly.

5.3.6 Waiting Time Setting for the Sleep Mode 1

Follow the procedure below to set up the waiting time for Sleep Mode 1.

Service Mode No.	Item
n.19	Waiting time setting for the Sleep Mode 1

To change the waiting time for Sleep Mode 1

1. Select **n.19** and press the jog dial. The default time **"001**" (1 minute) is displayed on the control panel, as shown in the right figure.



- 2. Turn the jog dial to the right to change the time (The selection is available from **1 to 999** minutes).
- 3. Press the dial again until you hear a beep.
- 4. Restart the device to check if the new setting is saved correctly.

5.3.7 Waiting Time Setting for the Sleep Mode 2

Follow the procedure below to set up the waiting time for Sleep Mode 2.

Service Mode No.	Item
n.20	Waiting time setting for the Sleep Mode 2

To change the waiting time for Sleep Mode 2,

1. Select **n.20** and press the jog dial. The default time "**005**" (5 minutes) is displayed on the control panel, as shown in the right figure.



- 2. Turn the dial to change the default time (The selection is available from **1 to 999** minutes).
- 3. Press the dial again and hold it until you hear a beep
- 4. Restart the device to check if the new setting is correctly saved.

5.3.8 Waiting Time Setting for the Power Down Mode

Follow the procedure below to set up the waiting time for the Power Down Mode.

Service Mode No.	Item
n.21	Waiting time setting for the Power Down Mode

To change the waiting time for the Power Down Mode,

1. Select **n.21** and press the jog dial. The default time "**300**" (300 minutes) is displayed on the control panel, as shown in the right figure.



- 2. Turn the jog dial to change the default time (The selection is available from **5 to 999** minutes).
- 3. Press the dial again and hold it until you hear a beep.
- 4. Restart the device to check if the new setting is saved correctly.

5.3.9 Buzzer On/Off

Follow the procedure below to turn on/off the buzzer. **Buzzer On** mode has three options: Battery Level 1 (flickering) warning only, Power Down Mode warning only, and both.

Service Mode No.	Item
n.22	Buzzer On/Off

To change the setting,

1. Select **n.22** and press the jog dial. The default setting **"002**" (Buzzer On for Power Down Mode only) is displayed on the control panel, as shown in the right figure.



- 2. Turn the jog dial to select an option out of the four options, as described below.
 - "000" = Buzzer Off
 - "001" = Buzzer On for Battery Level 1 (flickering) warning only
 - "002" = Buzzer On for Power Down Mode warning only (default)
 - "003" = Buzzer On for both
- 3. Press the dial again and hold it until a beep is heard.
- 4. Restart the device to check if the new setting is saved correctly (optional).

5.3.10 Power Down Mode On/Off

Follow the procedure below to turn on/off the Power Down Mode.

Service Mode No.	Item
n.23	Power Down Mode On/Off

To change the Power Down Mode On/Off setting,

1. Select **n.23** and press the job dial. The default setting "**001**" is displayed on the Control panel. ("**001**" = On, "**000**" = Off)



2. To turn off the Power Down Mode, turn the dial to change the setting from "**001**" to "**000**".



- 3. Press the dial again and hold it until a beep is heard.
- 4. Restart the device to check if the new setting is saved correctly.
- 5. To turn on the Power Down Mode again, select **n.23** and repeat the procedure above.

5.3.11 Password Setting

Follow the procedure below to change the password.

Service Mode No.	Item
n.24	Password setting

To change the password,

1. Select **n.24** and press the jog dial. The default password "**000**" is displayed on the control panel, as shown in the right figure.



- 2. When the first digit flashes, turn the jog dial to change the password and then save it by pressing the jog dial.
- 3. Follow the same procedure for the next two digits.
- 4. Press the dial until you hear a beep.
- 5. Check if the new password has been correctly saved.

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6. Troubleshooting

In instances of abnormal operation, error messages will be displayed on the Control panel. If a problem persists, please request assistance from the customer support information services.

Alarm/Error Messages

Error Code	Check Parameter	Description
E.02		When error codes "E.02", "E.03", "E.04," and "E.05" appear, X-ray exposure is impossible
E.03		even as the device's power remains steady.
E.04	X-ray Generator	Restart the device to resolve those error codes. The device resumes to normal operations after
E.05		a re-boot. (If the problem persists, please contact your Vatech representative.)
A.06		Appears when the system needs cooling time due to continuous operation. This alarm clears when the system temperature goes down to normal.
A.07	System	Appears when the power is turned on while the user is pressing the Exposure Button or Remote Exposure Switch. Release the switch.
A.08	User	Appears when the exposure button is released before the due time. Hold the exposure button until the end of the selected exposure time.
A.09	Battery	Appears when the battery voltage is higher than the reference value. Check the battery.
A.10		Appears when the battery voltage is lower than the reference value during X-ray exposure. Charge the battery.
A.11		Appears when the user attempts imaging while the device is still connected to the charger. Always disconnect the device from the charger before the start.
E.12		Appears when the battery level is below the reference value during X-ray exposure.

Troubleshooting

Problem	Cause	Solution
The device is not turned on.	The power switch is not turned on properly.	Turn the device power switch off and turn it back on.
	Battery discharged	Recheck after charging the battery with a charger.
	Battery cable is not correctly connected.	Contact your Service Representative.
	Defective battery	Contact your Service Representative.
The control panel is not turned on.	Defective mainboard	Contact your Service Representative.
	Internal cable disconnected	Contact your Service Representative.
No X-ray emission	The generator is cooling.	Wait for the cooling time (refer to 'Duty Cycle').
	Defective Remote Exposure Switch	Contact your Service Representative.
	Internal cable disconnected	Contact your Service Representative.
	Defective generator	Contact your Service Representative.
	Tube lifecycle termination	Contact your Service Representative.
X-ray emission works, but exposure is too light or completely white.	The device has been positioned incorrectly.	Adjust the position of the device.
	Exposure time is too long.	Decrease the exposure time.
	The receptor is facing the wrong way.	Reposition the receptor.
X-ray emission works, but exposure is too dark.	Exposure time is too short.	Increase the exposure time.

6. Troubleshooting

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7. Cleaning and Maintenance

7.1 Cleaning

WARNING Before cleaning the device, make sure to turn off the device.

- The device surfaces can be cleaned with a soft cloth damped in an alcohol-based, non-corrosive cleaning solution. If necessary, wipe off surfaces with disinfectant.
- If necessary, wipe off surfaces with disinfectant.
- Please observe the hygiene instructions of the phosphor plate scanner manufacturer.

	When cleaning the surfaces, make sure that the device is not connected to the battery charger.
	 DO NOT expose the device to any liquids. DO NOT use spray cleaner or disinfectant directly into the device as this could cause a fire.
NOTICE	The soft cloth should be damp but not dripping wet.
NOTICE	The cloths or wipes cannot be re-used.

7.2 Maintenance

VATECH requires periodic constancy tests to ensure image quality and the safety of the patient and operator.

Only **VATECH** authorized technicians can perform the inspection and service of this device. For technical assistance, contact the **VATECH** service center or your local **VATECH** representative.

Please observe the maintenance instructions of the phosphor plate scanner manufacturer.

Cautions and Notes

	DO NOT keep the device or its parts in a humid place or near a liquid substance.
	Avoid placing the device near chemical storage and gas-filled storage facilities.
NOTICE	When the device is not in use for a long time, fully charge the battery and remove it from the device before storage.
7.2.1 Maintenance Task Checklist

WARNING Always turn off the device before performing any maintenance.

7. Cleaning and Maintenance

Tasks	Period
Before the operation, ensure that the device is clean and ready for use.	Daily
After using the device, make sure that the device has been turned off.	Daily
Wipe the device's outer covers with a dry cloth at the end of each day's operation. DO NOT use detergents or solvents to clean the outer covers of the device.	Daily
Ensure that the signal is audible and the X-ray emission light is visible when making an exposure.	Daily
Ensure that the yellow (exposure) indicator light turns on when the Exposure Button is pressed.	Daily
Ensure that the battery charging LED indicator comes on when charging the battery.	Daily
Ensure that the battery level indicator displays at least two levels (Battery Level 2). For more information on the battery levels, see '4.5 Using the Battery'.	Daily
Ensure that all visible labels are intact and legible.	Monthly



If any defects are found, do not operate the device since the problem must be handled only by qualified personnel.

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8. **Disposing of the Unit**

This device is designed to be as safe as possible to use and dispose of to reduce environmental contamination. Many components of this device are environmentfriendly and can be recycled.

All parts and components that contain hazardous materials must be disposed of by disposal regulations. (IEC 60601-1 Clause 7.9.2.15)

Part	Material	Recyclable	Waste Disposal Site	Hazardous waste; Needs Separate Collection
Covers	Plastics	•		
Boards		•		
Cables and transformer	Copper	•		
	Polystyrene	•		
Packing	Cardboard	•		
	Paper	•		
X-ray tube				•
Battery				•
Other parts			•	

IMPORTANT

Observe all regulations relevant to the disposal of waste in your country.



This symbol on the device and accompanying documents means that used electrical and electronic devices (WEEE) should not be mixed with general household waste.



For professional users in the European Union:

When discarding electrical and electronic devices (EEE), please contact your dealer or supplier for further information.

For disposal in countries outside of the European Union:

This symbol is only valid in the European Union (EU). When discarding this device, please contact your local authorities or dealer and ask for the correct disposal method.

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9. Device Specifications

9.1 Mechanical Specifications

Dimensions



Item		Description		
Main	Dimension (mm)		280(L) x 296(H) x ø 165	
Body	Weight (kg)		2.14 (± 10 %)	
X-ray	X-ray Beam	Round Type	FOV: < ø 60	
Beam Limiting	Area (mm)	Rectangular Type	FOV: 20 x 30, 40 x 30	
Device	SSD(Source	to Skin Distance) (mm)	200	

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9.2 Technical Specifications

X-ray Generator

	Item	Description
	Model	DG- S0101V1
	Rated output power	Max. 0.2 kW
	Duty Cycle	1:60 or more
	Duty Cycle	(Exposure time: Interval time)
High Voltage Generator	Cooling Protection	Thermistor ≥ 65 °C
(Assembly)	Inherent Filtration	1.8 mm Al / 65 kV
	Total Filtration	Min. 1.5 mm Al
	Туре	Inverter Type
	Tube Voltage	55-65 kV
	Tube Current	1.0-3.0 mA
	Manufacturer	VATECH Co., Ltd.
	Model	V1-650304 (Stationary Anode type)
	Focal spot size	0.4 mm (IEC 60336)
	Anode heat contents	Max. 2.7 kJ
	Maximum Anode Heat Dissipation	200 W
X-ray Tube	Target Material	Tungsten
	Target Angle	12.5°
	Inherent Filtration	Min. 1.5 mm Al
	X-ray Coverage	70 mm at SID 200 mm
	Tube Voltage	Max. 65 kV
	Tube Current	Max. 3.0 mA

X-ray Tube Characteristics

1) Maximum rating chart



2) Emission characteristics



3) Heating and cooling curves of X-ray tube housing assembly



4) Tube Dimensions [mm]



Item	Description		
Model	1MET-D801 VT0601-2500		
Туре	Lithium-Ion Rechargeable Battery		
Nominal Capacity	2500 mAh		
Nominal Voltage	21.6 V _{dc}		
Charging Voltage	25.2 V _{dc} (4.2 V _{dc} /Cell)		
Discharge Voltage	19.8 - 25.2 Vdc		

IMPORTANT	This equipment uses either 1MET-D801 battery or VT0601- 2500 battery. The two battery specifications are identical.
	Make sure to use the battery only provided or approved by VATECH . Using an unauthorized battery may result in severe injury and device damage. For details on using the battery, see 'Battery Use.'

Battery Charger

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Item	Description
Model	XVE-2520200
Manufacturer	JIN XIN YU POWER(SHENZHEN)SUPPLY CO., LTD.
Dating	Input: 100-240 V~, 50/60 Hz, 1.5A
Rating	Output: 25.2 Vdc, 2.0 A
Frequency	50/60 Hz
Standard	IEC 60950-1 (UL)
Power Cord	300 V, 2.5 A

	Make sure to use the battery charger only provided or approved by VATECH . Using an unauthorized charger may result in severe injury and device damage. For details on using the battery charger, see 'Battery Use.'
NOTICE	 Power Supply is specified as a part of the ME EQUIPMENT. Power plugs may have various specifications for each country.

NO

9.3 Electrical Specifications

Item	Description
Tube Valtere	Option 1: 60 kV fixed (± 5 %)
Tube Voltage	Option 2: 65 kV fixed (± 5 %)
Tube Current	2.5 mA (± 10 %)
Exposure Time	0.05-1.0 s (± 3 % or 10 ms)
Rated Voltage	21.6 V _{dc}
	e system will be available with a fixed tube voltage ecification based on the user selection.



9.4 Environmental Specifications

Item		Description
During operating	Temperature	10 ~ 35 ℃
	Relative humidity	30 ~ 75 %
	Atmospheric pressure	860 ~ 1060 hPa
Transport and storage	Temperature	-10 ~ 60 °C
	Relative humidity	10 ~ 75 % non-condensing
	Atmospheric pressure	860 ~ 1060 hPa



Failure to follow the specifications above can result in severe injury and device damage.

10. Appendix

10.1 Combining the Holder and the devices using a tripod



1. Using a Phillips screwdriver, connect the EzRay Air Portable and Gender cable with the supplied screws.





2. Attach the holder to a tripod. Refer to the Tripod manual for how to attach to a tripod.



3. Attach the EzRay Air Portable unit to the holder with the tripod, as shown in the image below.



4. Connect one end of the Gender cable to the EzRay Air Portable device.



5. Connect the other end of the Gender cable connected to the holder to the supplied exposure cable.



10.2 Using the Rotating Rectangular Cover



1. Assemble the Rotating Rectangular Cover to the Cone of the Main body.





10.3 Tables of Exposure Times (Default)

The following exposure timetables were established with a unit equipped with a cone that corresponds to a focus-to-skin distance of 200 mm (8 inches), respectively.

DIGITAL SENSOR

1) Option 1: 60 kV, 2.5 mA, 0.05~1.0 sec

Patient	Teeth	Angle of inclination	SSD: 200 mm (8 inches)		
Patient	Teeth		kV	mA	S
	Incisor 🕅	Maxilla: +45° Mandible: -25°	60	2.5	0.14
Adult	Canine Ø	Maxilla: +45° Mandible: -20°	60	2.5	0.16
Ÿ	Molar/ Premolar Mandible: -5°	60	2.5	0.18	
	Bitewing	+5°~ +8°	60	2.5	0.20
Child	Incisor	Maxilla: +45° Mandible: -25°	60	2.5	0.12
	Canine Ø	Maxilla: +45° Mandible: -20°	60	2.5	0.14
	Molar/ Premolar	Maxilla: +30° Mandible: -5°	60	2.5	0.16
	Bitewing	+5°~ +8°	60	2.5	0.18

2)	Option 2: 65 kV, 2.5 mA, 0.05~1.0 sec
----	---------------------------------------

Patient	Teeth	Angle of	SSD: 200 mm (8 inches)		
Patient	inclination		kV	mA	S
	Incisor 🕅	Maxilla: +45° Mandible: -25°	65	2.5	0.12
Adult	Canine Ø	Maxilla: +45° Mandible: -20°	65	2.5	0.14
Ţ	Molar/ Premolar	Maxilla: +30° Mandible: -5°	65	2.5	0.16
	Bitewing	+5°~ +8°	65	2.5	0.18
	Incisor	Maxilla: +45° Mandible: -25°	65	2.5	0.10
Child	Canine Ø	Maxilla: +45° Mandible: -20°	65	2.5	0.12
ð	Molar/ Premolar	Maxilla: +30° Mandible: -5°	65	2.5	0.14
	Bitewing	+5°∼ +8°	65	2.5	0.16

PSP Scanner (VistaScan Nano Easy)

1) Option 1: 60 kV, 2.5 mA, 0.05~1.0 sec

Patient	Teeth Angle of inclination	Angle of	SSD: 200 mm (8 inches)		
		inclination	kV	mA	S
Adult	Incisor 🕅	Maxilla: +45° Mandible: -25°	60	2.5	0.30
	Canine Ø	Maxilla: +45° Mandible: -20°	60	2.5	0.42
	Molar/ Premolar	Maxilla: +30° Mandible: -5°	60	2.5	0.60
	Bitewing	+5°~ +8°	60	2.5	0.65
Child	Incisor	Maxilla: +45° Mandible: -25°	60	2.5	0.20
	Canine Ø	Maxilla: +45° Mandible: -20°	60	2.5	0.25
	Molar/ Premolar	Maxilla: +30° Mandible: -5°	60	2.5	0.40
	Bitewing	+5°~ +8°	60	2.5	0.40

2)	Option 2: 65 kV, 2.5 mA, 0.05~1.0 sec
----	---------------------------------------

Patient	Teeth	Angle of	SSD: 200 mm (8 inches)		
Patient	inclination		kV	mA	S
	Incisor	Maxilla: +45° Mandible: -25°	65	2.5	0.28
Adult	Canine Ø	Maxilla: +45° Mandible: -20°	65	2.5	0.40
Ŷ	Molar/ Premolar	Maxilla: +30° Mandible: -5°	65	2.5	0.58
-	Bitewing	+5°~ +8°	65	2.5	0.63
	Incisor	Maxilla: +45° Mandible: -25°	65	2.5	0.18
Child	Canine Ø	Maxilla: +45° Mandible: -20°	65	2.5	0.23
ð	Molar/ Premolar	Maxilla: +30° Mandible: -5°	65	2.5	0.38
	Bitewing	+5°~ +8°	65	2.5	0.38

10.4 X-ray Dose Data

The X-ray dose data is extracted from the X-ray Dose Test Report for the device. The IEC collateral standards measured the X-ray doses of the device in the test report. The device was designed by Part 1. General Requirements for Safety, IEC 60601-1-3.

Test Condition			
Model Name	VEX-P300		
Tube Model Name	V1-650304		
Generator Model Name	DG-S0101V1 (Inverter type)		
Looding Easter	Option 1: 60 kV, 2.5 mA		
Loading Factor	Option 2: 65 kV, 2.5 mA		

10.4.1 X-ray Dose Table

	Test Equipment				
Instrument	Instrument Manufacturer Model S/N				
Multi-Dose Meter	Raysafe	Unfors Xi mAs / Unfors Xi R/F & MAM	163288 / 161834		

Digital Sensor

Dose Area Product (DAP) Table (60 kVp, 2.5 mA, SSD 200 mm)					
	FOV: Ø 6 cm	FOV : 3 x 4 cm	FOV : 2 x 3 cm		
t (s)	Dose (mGy.cm ²)				
0.14	6.5	2.8	1.4		
0.16	7.5	3.2	1.6		
0.18	8.5	3.6	1.8		
0.20	9.8	4.2	2.1		

Dose Area Product (DAP) Table (65 kVp, 2.5 mA, SSD 200 mm)					
	FOV: Ø 6 cm	FOV : 3 x 4 cm	FOV : 2 x 3 cm		
t (s)	Dose (mGy.cm ²)				
0.12	6.7	2.9	1.4		
0.14	7.9	3.4	1.7		
0.16	9.1	3.9	1.9		
0.18	10.2	4.3	2.2		

Phosphor Plate

Dose Area Product (DAP) Table (60 kVp, 2.5 mA, SSD 200 mm)					
	FOV: Ø 6 cm	FOV : 3 x 4 cm	FOV : 2 x 3 cm		
t (s)	t (s) Dose (mGy.cm ²)				
0.30	14.7	6.2	3.1		
0.42	20.1	8.5	4.3		
0.60	29.2	12.4	6.2		
0.65	31.6	13.4	6.7		

Dose Area Product (DAP) Table (65 kVp, 2.5 mA, SSD 200 mm)					
	FOV: Ø 6 cm	FOV : 3 x 4 cm	FOV : 2 x 3 cm		
t (s)	t (s) Dose (mGy.cm ²)				
0.28	15.3	6.5	3.2		
0.40	23.1	9.8	4.9		
0.58	32.9	14.0	7.0		
0.63	36.4	15.4	7.7		

10.4.2 Leakage Dose

Scope

IEC 60601-2-65 203.12.4

Requirements

In the LOADING STATE, the AIR KERMA due to LEAKAGE RADIATION from X-RAY SOURCE ASSEMBLIES, 1 m from the FOCAL SPOT, average over an area of 100 cm² of which no principal linear dimension exceeds 20 cm when operated at the NOMINAL X-RAY TUBE VOLTAGE under the condition of LOADING corresponding to the reference LOADING conditions, shall not exceed 0.25 mGy in one hour.

Leakage Dose	Permissive Range
65 kVp, 2.5 mA, 1.0 s (Max. Exposure Condition)	
At Focal Spot to Distance 1 m	< 0.25 mGy/h
1 : 60 Duty Cycle	

Test Equipment				
Instrument	Manufacturer	Model	S/N	
X / Gamma Survey Meter	Radcal Co.	9015/10X5-180	91-1470/19069	

Results

The following exposure timetables were established with a unit equipped with a cone that corresponds to a focus-to-skin distance of 200 mm (8 inches), respectively. When the leakage doses have been measured with each cover type (default, rectangular 2x3, and rectangular 4x3), all the results have been ND (Not Detected). The raw data about the results are shown in the table below.



Direction	Default type [mGy/h]	Rectangular 2x3 [mGy/h]	Rectangular 4x3 [mGy/h]
0°	ND	ND	ND
20°	ND	ND	ND
45°	ND	ND	ND
65°	ND	ND	ND
90°	ND	ND	ND
110°	ND	ND	ND
135°	ND	ND	ND
155°	ND	ND	ND
180°	ND	ND	ND
200°	ND	ND	ND
225°	ND	ND	ND
245°	ND	ND	ND
270°	ND	ND	ND
290°	ND	ND	ND
315°	ND	ND	ND
335°	ND	ND	ND



Direction	Default type [mGy/h]	Rectangular 2x3 [mGy/h]	Rectangular 4x3 [mGy/h]
0°	ND	ND	ND
20°	ND	ND	ND
45°	ND	ND	ND
65°	ND	ND	ND
90°	ND	ND	ND
110°	ND	ND	ND
135°	ND	ND	ND
155°	ND	ND	ND
180°	ND	ND	ND
200°	ND	ND	ND
225°	ND	ND	ND
245°	ND	ND	ND
270°	ND	ND	ND
290°	ND	ND	ND
315°	ND	ND	ND
335°	ND	ND	ND

• ND: Not Detected. The detection limit is 0.00001 mGy per exposure.

10.4.3 Scattered Dose

<u>Scope</u>

IEC 60601-2-65 203.13

Requirements

ME EQUIPMENT shall be provided with means to optionally allow actuation of the EXPOSURE from a PROTECTED AREA after installation.

Relevant instructions shall be given in the ACCOMPANYING DOCUMENTS.

Results

The following exposure timetables were established with a unit equipped with a cone that corresponds to a focus-to-skin distance of 200 mm (8 inches), respectively.

	Test Equ	ipment	
Instrument	Manufacturer	Model	S/N
X / Gamma Survey Meter	Radcal Co.	9015/10X5-180	91-1470/19069

Method

PMMA Phantom aligned to 280 mm away from Focal Spot

Max. Exposure Condition

Measure point: 1000 mm from PMMA Phantom

Direction [9]	Result (Horizontal Plane) [µR]		
Direction [°]	Without Shield	With Shield	
0	63.4	68.14	
20	17.8	11.99	
45	18.5	13.01	
65	19.8	14.76	
90	27.1	20.65	
110	41.6	32.08	
135	63.4	46.18	
155	68.1	11.14	
180	3.29	2.893	
200	51.5	7.557	
225	47.5	48.63	
245	33.7	33.27	
270	23.4	33.25	
290	19.8	18.06	
315	17.2	15.23	
335	19.2	14.71	

Option 1: 60 kV / 2.5 mA / 1.0 s



Direction [8]	Result (Vertical Plane) [µR]		
Direction [°]	Without Shield	With Shield	
0	25.8	29.8	
20	27.3	26.8	
45	18.5	22.5	
65	15.9	18.5	
90	74.1	72.1	
110	15.9	18.5	
135	17.2	13.5	
155	19.9	17.9	
180	19.2	17.9	
200	27.8	31.4	
225	13.9	10.6	
245	38.4	8.6	
270	7.9	4.3	
290	48.4	11.9	
315	60.2	52.3	
335	41.7	45.7	



Direction [9]	Result (Horizontal Plane) [µR]		
Direction [°]	Without Shield	With Shield	
0	58.8	85.2	
20	22.5	30.4	
45	39.6	42.7	
65	47.6	40.3	
90	46.9	42.9	
110	47.6	66.7	
135	92.5	48.9	
155	87.9	34.3	
180	11.9	11.9	
200	94.5	28.4	
225	89.2	80.3	
245	86.2	49.5	
270	51.6	42.3	
290	47.6	43.6	
315	27.1	30.7	
335	40.4	23.1	

Option 2: 65 kV / 2.5 mA / 1.0 s



	Result (Vertical Plane) [µR]		
Direction [°]	Without Shield	With Shield	
0	64.7	55.4	
20	42.8	44.3	
45	26.3	34.8	
65	36.8	36.4	
90	57.9	82.4	
110	42.3	38.2	
135	20.7	25.9	
155	27.7	31.9	
180	36.2	52.6	
200	94.6	63.2	
225	64.2	49.8	
245	42.9	18.4	
270	11.2	11.3	
290	24.6	7.2	
315	52.8	26.5	
335	79.1	72.2	



10.5 Electromagnetic Compatibility (EMC) Information

Guidance and manufacturer's declaration - electromagnetic emissions

The VEX-P300 is intended for use in the electromagnetic environment specified below. The customer or the user of the VEX-P300 should assure that it is used in such an environment.

Emissions test	Compliance	Electromagnetic environment - guidance
RF emissions CISPR 11	Group 1 Class A	The EMISSIONS characteristics of the VEX-P300 make it suitable for use in industrial areas and hospitals (CISPR 11 class A). If used in a residential environment (for which CISPR 11 class B is normally required), this equipment might not offer adequate protection to radiofrequency communication services. The user might need to take mitigation measures, such as relocating or re-orienting the equipment.
Harmonic emissions IEC 61000-3-2	Applicable	The VEX- P300 is suitable for use in all establishments and may be used in domestic establishments and these directly connected to
Voltage fluctuations / flicker emissions IEC 61000-3-3	Applicable	and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.

Guidance and manufacturer's declaration - electromagnetic immunity

The VEX-P300 is intended for use in the electromagnetic environment specified below. The customer or the user of the VEX-P300 should assure that it is used in such an environment.

Test of Electronic Interference Resistance	IEC 60601-1-2 test level	Compliance level	Electromagnetic environment – guidance
Electrostatic discharge (ESD) IEC 61000- 4-2	Discharge by ±8 kV direct contact ±15 kV of Air-gap discharge	Discharge by ±8 kV direct contact ±15 kV of Air- gap discharge	Floors should be wood, concrete, or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30 %.
Radiated RF Electromagn etic Field IEC 61000- 4-3	3 V/m 80 MHz-2.7 GHz 80% AM at 1 kHz	3 V/m 80 MHz-2.7 GHz 80% AM at 1 kHz	The VEX-P300 is suitable to use in a professional healthcare environment.
Immunity to Proximity Fields from RF wireless Communicati ons Equipment according to Table 9 in IEC 60601- 1-2	28 V/m Max. 385-5785 MHz in according to table 9	28 V/m Max. 385-5785 MHz in according to table 9	RF communication equipment is used on closer than 30 cm to any part of the VEX-P300, including cables specified by VATECH
Electrical fast transient/bur st IEC 61000-4-4	 ± 2 kV for power supply lines ± 1 kV for input/ output lines 100 kHz repetition frequency 	± 2 kV for power supply lines ± 1 kV for input/ output lines 100 kHz repetition frequency	The quality of supplied power should be suitable for the general business site or hospital environment.
Surge IEC 61000- 4-5	± 1 kV line(s) to line(s) ± 2 kV line(s) to earth	± 1 kV line(s) to line(s) ± 2 kV line(s) to earth	The quality of supplied power should be suitable for the general business site or hospital environment.

Immunity to conducted disturbances induced IEC 61000-4-6	3 V 0.15-80 MHz 6 V in ISM bands between 0.15 and 80 MHz 80% AM at 1 kHz Power supply line & I/O lines	3 V 0.15-80 MHz 6 V in ISM bands between 0.15 and 80 MHz 80% AM at 1 kHz Power supply line & I/O lines	The strength of the RF field in the frequency range higher than 150 kHz ~ 80 MHz, the strength of the RF field is smaller than 3 V
The magnetic field of supply frequency (50/60 Hz) according to IEC 61000- 4-8	30 A/m	30 A/m	The magnetic field of power frequency should coincide with the genera level found in the business site or hospital environment
Voltage dips and short interruptions IEC 61000- 4-11	0 % <i>U</i> T: 0.5 cycle At 0°, 45°, 90°, 135°, 180°, 225°, 270° and 315° 0 % <i>U</i> T; 1 cycle and 70 % <i>U</i> T; 25/30 Cycles Single phase: at 0° 0 % <i>U</i> T; 250/300 cycle	0 % <i>U</i> T: 0.5 cycle At 0°, 45°, 90°, 135°, 180°, 225°, 270° and 315° 0 % <i>U</i> T; 1 cycle and 70 % <i>U</i> T; 25/30 Cycles Single phase: at 0° 0 % <i>U</i> T; 250/300 cycle	The quality of supplied power should be suitable for the general business site or hospital environment. If the user of the VEX- P300 image intensifier requires continued operation during power mains interruptions, it is recommended that the VEX- P300 image intensifier is powered from an uninterruptible power supply.

10.6 Abbreviations

Acronym	Name
AL	Aluminum
EMC	Electromagnetic Compatibility
ESD	Electrostatic Discharge
FOV	Field of View
IEC	International Electrotechnical Commission
ISO	International Standards Organization
LED	Light-Emitting Diode
ME	Medical Electrical
РММА	Poly Methyl Meth Acrylate
RF	Radio Frequency
SID	Source to Image receptor Distance
SIP	Signal Input Part
SOP	Signal Output Part
SSD	Source to Skin Distance

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CE 2460	The CE symbol grants this device compliance to the European Directive for Medical Devices 93/42/EEC as amended by 2007/47/EC as a class IIb device.
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