

XT Sense

CPAP System

Instruction Manual



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IMPORTANT SAFEGUARDS – PLEASE MAKE THIS PRECAUTIONS LIST AVAILABLE TO ALL USERS and PROVIDERS

READ ALL INSTRUCTIONS BEFORE USING THIS SYSTEM

WARNING –

1. THIS DEVICE IS NOT INTENDED FOR LIFE SUPPORT. It may stop operating due to power interruption but no hazards to patient.
2. If oxygen is used with the device, the oxygen flow must be turned off when the device is not in use.
Explanation of the Warning: When the device is not in operation and the oxygen flow is left on, oxygen delivered into the tubing may accumulate within the device's enclosure and create a risk of fire.
3. Oxygen supports combustion. Oxygen should not be used while smoking or in the presence of an open flame.
4. Always ensured the device is generating airflow before the oxygen supply is turned. Always turn off the oxygen supply before stopping the airflow from the device.
5. This device **should not** be used in the vicinity of a flammable anesthetic mixture in combination with oxygen or air and nitrous oxide.
6. The airflow for breathing generated by this device may be as much as 7°C (12.6° F) higher than the room temperature. This device **should not** be used if the room temperature is warmer than 35°C (95°F) to prevent the airflow temperature from exceeding 40° C (104° F) and causing irritation to your airway.
7. If this device overheats, it will stop operating and show “**Error 002**” on the display. After cooling down to proper temperature, the device can restart again.
8. This machine should be used only with masks (and connectors) recommended by the manufacturer, or by your physician or respiratory therapist. A mask should not be used unless the CPAP machine is turned on and operating properly. The vent holes associated with the mask should never be blocked for proper exhaling purpose. If the vent hole is blocked, the CPAP machine will stop and show message “**Error 002**”, after cooling down, please re-connect the power cord to reset the machine.
9. At low CPAP pressure, some exhaled gas may remain in the mask and be re-breathed.

CAUTION –

1. Make sure the environment around the machine is dry and clean. Dust and foreign particles may affect the treatment. Keep the air inlet on the back of the machine clear to prevent overheating and damage of the device. Do not place the machine near a source of hot or cold air. Extreme cold or hot environment may damage user's respiratory airway.
2. If there is a possibility of electro-magnetic interference with mobile phones, please increase the distance between devices or turn off the mobile phone.
3. Do not connect the device to the personal computer for data downloading during the treatment. This may cause the CPAP system failure.
4. To prevent from potential allergic reactions to mask used in the device, This machine shall be used only with masks (and connectors) recommended by the manufacturer.
5. U.S. Federal law restricts this device to sale by or on the order of a licensed physician.

DANGER -To reduce the risk of electrocution:

1. Always unplug this product immediately after using.
2. Do not use while bathing.
3. Do not place or store product where it can fall or be pulled into a tub or sink.
4. Do not place in or drop into water or other liquid.
5. Do not reach for a product that has fallen into water. Unplug immediately.

WARNING -To reduce the risk of burns, electrocution, fire or injury to persons:

1. This product should never be left unattended when plugged in.
2. Close supervision is necessary when this product is used by, on, or near children or invalids. Choking resulting from a child swallowing a small part that has become detached from the device or its accessories.
3. Use this product only for its intended use as described in this manual, do not use attachments not recommended by the manufacturer.
4. Never operate this product if it has a damaged cord or plug, if it is not working properly, if it has been dropped or damaged, or dropped into water. Return the product to a service center for examination and repair.
5. Keep the cord away from heated surfaces.
6. Never block the air openings of this product or place it on a soft surface, such as a bed or couch, where their openings may be blocked. Keep the air opening free of lint, hair, and other similar particles.
7. Never drop or insert any object into any opening on the unit or into the tubing.
8. When disposing the unit because of damage or failure, you must comply with the regulations of your particular region or country for disposing such equipment
9. No modification of this equipment is allowed.
10. Do not modify this equipment without authorization of the manufacturer.
11. If this equipment is modified, appropriate inspection and testing must be conducted to ensure continued safe use of the equipment.
12. Do not leave long lengths of air tubing around the top of your bed. It could twist around your head or neck while you are sleeping and lead to strangulation.

1. Introduction

This manual is provided to explain initial set up and operating instructions. This manual should be stored in a safe location and be readily accessible for future reference.

1.1 General Information

Obstructive Sleep Apnea (OSA) is a condition that an intermitted and repetitive obstruction of the upper respiratory tract causes a complete (apnea) or partial (hypopnea) block of breathing airflow during sleep. The syndrome varies depending on the degree of relaxation of the tongue and soft palate muscle.

The most common treatment for OSA is Continuous Positive Airway Pressure (CPAP). CPAP devices can deliver a constant air pressure into your upper airway via a nasal mask. This constant air pressure can keep your airway open during sleep, therefore prevents the OSA.

This device is a micro-processor controlled continuous positive airway pressure device. It features the illuminated, menu-driven LCD display, universal power supply, and ramp time adjustment. The ramp time adjustment and ultra quiet operation ensure you to fall asleep comfortably while air pressure slowly build up to treatment level. The user compliance meter records the total system's operating time for physician's reference.

The system has been tested and successfully approved to the following standards:



- EN 60601-1
- EN 60601-1-2
- EN 61000-3-2 Class A
- EN 61000-3-3



For US and CANADA only

E348970

53DG



Medical Equipment- CPAP
**WITH RESPECT TO ELECTRICAL SHOCK,
FIRE AND MECHANICAL HAZARDS ONLY**
IN ACCORDANCE WITH ANSI/AAMI
ES60601-1 (2005,3rd ed.) and CAN / CSA C22.2
No. 60601-1 (2008)

Le produit à été testé avec des équipements médicaux et respecte les normes ANSI/AAMI ES60601-1 (2005,3rd ed.) and CAN / CSA C22.2 No. 60601-1 (2008). prévenant les choc électrique, le feu et les risques de blessures physiques.

EMC Warning Statement

This equipment has been tested and found to comply with the limits for medical devices to the EN 60601-1-2:2007. These limits are designed to provide reasonable protection against harmful interference in a typical medical installation. This equipment generates, uses and can radiate radio frequency energy and, if not installed and used in accordance with the instructions, may cause harmful interference to other devices in the vicinity. However, there is no guarantee that interference will not occur in a particular installation. If this equipment does cause harmful interference to other devices, which can be determined by turning the equipment off and on, the user is encouraged to try to correct the interference by one or more of the following measures:

- Reorient or relocate the receiving device.
- Increase the separation between the equipment.
- Connect the equipment into an outlet on a circuit different from that to which the other device(s) are connected.
- Consult the manufacturer or field service technician for help.

This system has been tested and compliance to the following volunteer standards: **FDA**.

1.2 Intended Use

This device is intended to provide continuous positive airway pressure (CPAP) for the treatment of adult Obstructive Sleep Apnea (OSA).



Cautions: Some patients might have pre-existing contraindications for CPAP therapy, or might experience some potential side effects of using CPAP device, please consult your physician if you have any questions concerning your therapy.

2. Product Description

Components including:

- (1) Main CPAP device
- (2) Detachable power cord
- (3) User manual
- (4) Flexible air tubing with 1.8 m length
- (5) Carrying bag
- (6) SD card

⚠ Note 1: ONLY for Physician or Technician to download data. Patient should not use this unction.

⚠ Note 2: Only applicable for devices with SD card slot.

⚠ Note 3: Please use SD card (smaller than 2GB) which comply with SDHC standard. Before using it, please format it to FAT16 to ensure correct data collection.



3. Installation

3.1 Unpacking

To secure its contents inside, the CPAP device and accessories are bundled in a paper packaged box. Unpack this box by removing the CPAP and its accessory and checking for any damage, which may have occurred during shipping. If there are damages, please contact your dealer immediately.

3.2 Setting Up



- 1) Connect the power cord to CPAP device and plug into main electrical outlet.

Once the power cord is plugged into the electrical outlet, the device is in ready to operate position (“**STANDBY**” sign appears in LCD display)

 **NOTE:** The plug is also served to disconnect the device.



- 2) Connect one end of the air tubing firmly onto the air outlet of the CPAP.



- 3) Connect the other end of the air tubing to the mask system. Putting on the mask and headgear according to the mask instruction manual.

4. Operation

 **NOTE:** Always read the operating instruction before use.

4.1 Control Panel Description

Buttons arrangement on control panel and main use of the buttons:



START/STANDBY

To start the treatment, simply press the **"START/STANDBY"** button. To stop the treatment, press the **"START/STANDBY"** button again. The display will switch between **[STANDBY]** and Therapy Pressure **[XX.X cmH₂O]** in cmH₂O unit.



MENU

Press the **"MENU"** button to enter the setting mode when device is in standby mode. The adjustment setting includes mode selection, ramp time selection, ramp starting pressure, therapy pressure adjustment, alarm ON/OFF, clock alarm and clock setting, compliance meter, and total operating meter. When each setting's value has been changed, press **"MENU"** for confirmation and press **"MENU"** again for next setting selection. Please refer to 4.2 Function Description section for detailed information.



UP

Press the **"UP"** button to increase the value.



DOWN

Press the **"DOWN"** button to decrease the value.

4.2 Function Description

Getting Started – First time use.

For first-time use, follow the steps below:

1. Make sure that your physician or care provider has initialized and configured the appropriate treatment values into the unit.
2. Make sure that your package contains a face or nasal mask with headgear. If not, your physician or care provider can recommend the type of mask and headgear you must obtain.
3. Attach tubing and mask (see Setup section).
4. Attach power cord to the unit and an electrical outlet (see Setup section). Unit will power-on automatically.
5. Use the **"MENU"** button to set the Current Time, Ramp Up Time, and Ramp Starting Pressure, (see **"MENU"** Button Section for description of these options).
Note that these values may have been already set by your physician or care provider.
6. Press the **"START/STANDBY"** button – Airflow to the mask will begin.
7. You can now place the Headgear and Mask over your head and face and begin treatment.

4.3 Menu Button - Option Descriptions

The following items describe the options presented to you after you press the **"MENU"** button on the control panel.

(1) Ramp Time

Ramp time function allows user to fall into sleep with a lower, comfortable pressure and helps users gradually get accustomed to increasing treatment pressure. The first selection of pressing **"MENU"** is **[Ramp XX MIN]**. When the **"MENU"** setting is in **[Ramp XX MIN]** mode, press **"UP"** or **"DOWN"** button to set the preferred ramp time and press **"MENU"** for confirmation. There are 10 adjustable levels, 0, 5, 10, 15, 20, 25, 30, 35, 40 and 45 minutes.

(2) Ramp Starting Pressure

Press **"MENU"** button to select **[Ramp P XX.X]** menu, press **"UP"** or **"DOWN"** button to set the preferred ramp starting pressure and press **"MENU"** for confirmation. The ramp starting pressure can be changed from 3 cmH₂O to "Therapy Pressure – 1" cmH₂O. For example, if your therapy pressure is 10 cmH₂O, the maximum ramp starting pressure you can select is 9 cmH₂O.

(3) Therapy Pressure

Press **"MENU"** button to select the **[P XX.XcmH₂O]** menu. You can view the current pressure setting displayed in cmH₂O unit. Therapy pressure is adjustable only by the provider, a respiratory therapist or doctor.

 **NOTE:** The therapy pressure is to only be prescribed by a physician.

(4) Total Meter

Press **"MENU"** button to select **[TM XXXX.X hr]** menu, the total compliance meter records the total number of hours that the device has been active. The meter should only be re-set by the provider, a respiratory therapist or by a physician.

(5) Compliance Meter

Press **"MENU"** button to select the **[CM XXXX.X hr]** menu. The compliance meter records the total therapy hours for the device. The compliance meter should be re-set only by the provider, a respiratory therapist or by a doctor.

(6) Alarm

Press **"MENU"** button to select the **[Alarm on/off]** menu, and press **"UP"** or **"DOWN"** button to set the alarm on or off. When the alarm is turned on, the audible alarm will activate with warning messages shown on the LCD display. Set alarm off to mute the audible alarm.

(7) Clock Alarm

Press **"MENU"** button to select the **[Clock Alarm on/off]** menu and press **"UP"** or **"DOWN"** button to set the clock alarm on or off. When the clock alarm is set on, the display will show the time on the left side. Press **"UP"** or **"DOWN"** button to set the time to wake you up. Once the clock alarm is activated, press the start/standby button to mute the audible alarm.

(8) Clock

Press **"MENU"** button to select the **[Clock XX:XX]** menu, press **"UP"** or **"DOWN"** button to set the current time.

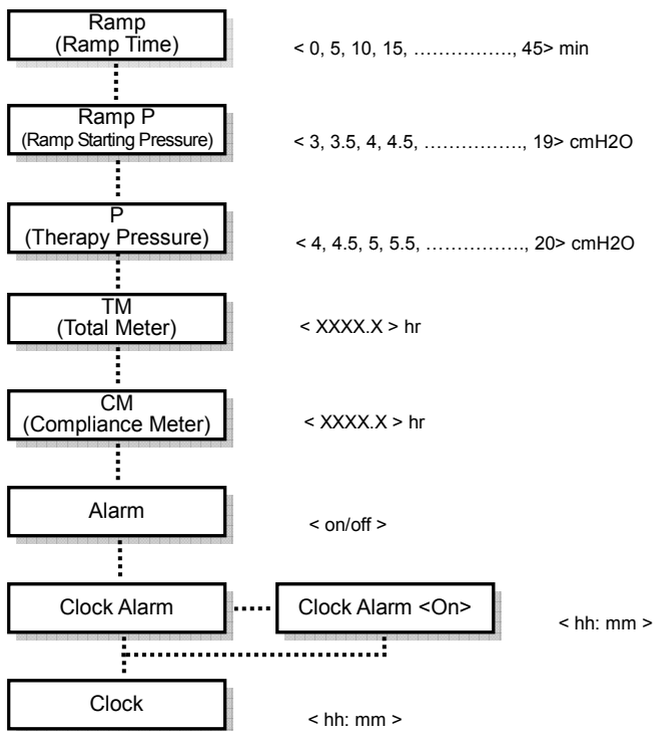
(9) Turning off the Device

Remove the power cord from the electrical outlet, and disconnect power cord from the power socket on the right side of the device.

 **NOTE:** Once the setting is confirmed, press the **"MENU"** button. Otherwise, the device will automatically go back to standby without saving the modification if no action is taken in 20 seconds.

4.4 Flowchart of Menu settings

Enter the user's menu mode by pressing the **"MENU"** button.



In each setting, when the preferred value has been selected, press **"MENU"** for confirmation and press **"MENU"** again to enter next selection.

NOTE: For physicians, please refer to a separated "Physician's Additional Instruction" page.

5. Adding a Humidifier



XT Sense CPAP device can be used with XT Heated Humidifier (**9S-006500**) which is available from the home care provider. The heated humidifier may reduce nasal dryness and irritation by providing adequate moisture and heat to the airflow. Please refer to the XT heated humidifier (**9S-006500**) instruction manual for complete setup information.



NOTE: When XT Sense CPAP device is used with the heated humidifier, its power supply is from the power socket outlet of the heated humidifier. Do not connect the power cord to CPAP device and plug into main electrical outlet.

6. Using the SD Card to Collect Data

If physicians need to review the usage data, they may ask you to use the SD card to copy data from the device, and to return the card to them. Data that is copied to the SD Card is still stored and available on the device.

1. Insert the SD Card when the device is in standby mode.
2. Data copying starts automatically when the SD Card is inserted into the slot.
3. The **[COPY] & [Card]** messages are displayed on the LCD while data is being copied.
4. The **[End] & [OK]** messages are displayed on the LCD when copying has finished.
5. Remove the SD Card and mail it to the clinician.
6. The SD card does not need to be uninstalled for the device to work properly.



CAUTION: Do not remove the SD card until **[End] & [OK]** messages are displayed on the LCD, or data copied to the card may be corrupted or missing.

7. Cleaning & Maintenance

7.1 Cleaning the Main Unit

The main unit should be cleaned regularly (at least every 30 days). To clean the unit, wipe it down with a damp cloth using a mild detergent. When selecting a cleaning agent, take precautions to choose one that is not chemically abrasive to the surface of the plastic case. All parts should be air-dried thoroughly before use. During storage, the unit should be kept in a dust-free location.

 **WARNING:** Don't try to open this device. Repairs and internal servicing should only be performed by an authorized service agent. Don't drop any subject into the air tube or air outlet.

7.2 Tubing and Mask

The tubing and mask should be checked and cleaned daily. Please refer to the cleaning instruction packaged with the accessories.

1. Disconnect the air tubing from the air outlet of the device.
2. Remove the air tubing and headgear straps from the nasal mask.
3. Wash the mask system according to the instructions supplied with it. Wash the air tubing in warm water using mild detergent.
4. Rinsed thoroughly, hang and allow to dry.
5. Before next use, assemble the mask and headgear according to the mask user instructions.
6. All items of the mask and air tubing are subject to normal wear and tear and may eventually be replaced. Replace the mask and the air tubing if they are damaged.

 **CAUTION:** Do not use bleach, chlorine-, alcohol-, or aromatic-based (including all scented oils), moisturizing or antibacterial soaps to clean the cushion, mask, air tubing. These solutions may cause hardening and reduce the life of the product.

 **CAUTION:** Do not wash or dry the mask or air tubing at a temperature above 70°C (160°F)

 **WARNING:** Do not use any cleaner containing fragrance or conditioners as they will leave a residue.

 **WARNING:** The mask must not be re-used by another person. This is to avoid the risk of cross-infection.

7.3 Air Filter Maintenance

For an optimum operation of the device, the air filter can be cleaned by the user at least once every two weeks or more often if this device is operated in a dusty environment. It is recommended that the filter be replaced with a new one after 6 months of use; however, depending on the air quality, the replacement time may vary.

CAUTION: Dirty air filter may cause high operating temperatures that affect device performance. Ensure the air filter is cleaned and fitted at all times.



1. Remove the dirty filter from the enclosure on the rear of the device.
2. Wash the filter in warm water with a mild detergent, and rinse with water. Allow the filter to air dry completely before reinstalling. Do not use a filter that is not completely dry. If the filter is torn, replace it.
3. Reinstall the filter.

NOTE: Please follow national requirements to dispose the unit properly.

8. Troubleshooting

The table below lists troubleshooting solutions for the problems that may happen. If the problem persists, contact your equipment provider service agent.

Problem	Possible Causes	Solutions
No display	<ol style="list-style-type: none"> 1. The power cord is not connected to the power socket. 2. LCD failure or controlled PCB failure. 	<ol style="list-style-type: none"> 1. Ensure the power cord is connected. 2. Contact your equipment provider for repair.
Display code incorrect	LCD failure or controlled PCB failure.	Contact your equipment provider for repair.
Illuminant under LCD is not on	LED failure	Contact your equipment provider for repair.
Buttons disable	Button failure	Contact your equipment provider for repair.
Air delivered is slow	<ol style="list-style-type: none"> 1. During ramp time. 2. Filter is too dirty. 3. Flow generator failure. 	<ol style="list-style-type: none"> 1. Check the ramp time setting 2. Change or clean the filter regularly. 3. Contact your equipment provider for repair.
Data can not be copied to the SD card	<ol style="list-style-type: none"> 1. SD card is full. 2. SD card is not inserted correctly. 3. Data on the SD card is corrupted. 	<ol style="list-style-type: none"> 1. Ensure the SD card has enough capacity. 2. Ensure the SD card is inserted into the slot. 3. Format the card.

Error / Warning Messages show in LCD.

Message type	Definition	Message in LCD
Error: Primary function can't execute.	Error for abnormal system settings	Error 001
	Error for flow generator failure	Error 002
	Error for abnormal timer setting or timer failure	Error 003
Warning:	Out of system memory	Warn 001
	System memory is nearly full	Warn 002
	SD card module communication failure	Warn 003
	Remove the SD card while data is being processed	Warn 004
	SD card is full	Warn 005
	Inconsistent ID no. of device with SD card	Warn 006
	Wrong firmware version of SD card module	Warn 007
	SD card is damaged	Warn 008

NOTE: When the warning message appears, contact your physician or equipment provider to download the memory data and reset the meter.

9. Technical Specifications

Item		Specifications
Power Supply		Universal power supply, AC100-240V, 50/60 Hz, 0.5-0.3A
Pressure Range		4-20 cmH ₂ O (adjustable in 0.5 cmH ₂ O increment)
Ramp Time		0-45 minutes (adjustable in 5-minute increment)
Ramp Starting Pressure		3-19 cmH ₂ O (adjustable in 0.5 cmH ₂ O increment)
Operating Altitude		up to 8,000 ft (2,438 m) when the pressure is set at 4-18 cmH ₂ O but limit to 5,000ft (1,524m) when the pressure is set at 18.5 - 20 cmH ₂ O
Dimensions (W x D x H)		14.5 x 13.0 x 10.0 cm or 5.7" x 5.1" x 3.9"
Weight		Approximately 800 g or 1.76 lb
Environment	Temperature	Operating: +5 °C to +35 °C (+41 °F to +95 °F) Storage: -15 °C to 50 °C (+5 °F to +122 °F) Shipping: -15 °C to 70 °C (+5 °F to +158 °F)
	Humidity	Operating: 15%RH to 95%RH non-condensing Storage: 10%RH to 90%RH non-condensing Shipping: 10%RH to 90%RH non-condensing
Classification:		Class II Type BF, Applied Parts: Mask Not suitable for use in the presence of a flammable anesthetic mixture IP21: Protected against solid foreign objects of 12.5 mm and greater; Protected against vertically falling water drops. Continuous operation.

Note: The manufacturer reserves the right to modify the specification without notice.

Symbols



BF symbol, which indicated this product, is according to the degree of protecting against electric shock for type BF equipment.



Attention, should read the instructions.



Refer to instruction manual



Attention, should read the instructions.



Class II

IP21 

Protected against solid foreign objects of 12.5 mm and greater; Protected against vertically falling water drops.



Disposal of Electrical & Electronic Equipment (WEEE):

This product should be handed over to an applicable collection point for the recycling of electrical and electronic equipment. For more detailed information about the recycling of this product, please contact your local city office, household waste disposal service or the retail store where you purchased this product.



Authorized representative in the European community



Manufacturer

10. Note, Caution, And Warning Statements

NOTE: Indicate information that you should pay special attention to.

CAUTION: Indicate correct operating or maintenance procedures in order to prevent damage to or destruction of the equipment or other property.

WARNING: Calls attention to a potential danger that requires correct procedures or practices in order to prevent personal injury.

11. Servicing

The XT series systems are intended to provide safe and reliable operation according to the instructions provided by Apex Medical. Apex Medical recommends that the XT system be inspected and serviced by authorized technicians if there is any sign of wear or concern with device function. Otherwise, service and inspection of the devices generally should not be required during the five year design life of the device.

12. Limited Warranty

Apex Medical Corp. (hereafter 'Apex') gives the customer a limited manufacturer warranty on new original Apex products and any replacement part fitted by Apex in accordance with the warranty conditions applicable to the product in question and in accordance with the warranty periods from date of purchase as listed below. This warranty does not cover damage caused by accident, misuse, abuse, alteration and other defects not related to material or workmanship. To exercise your rights under this warranty, please contact your local, authorized Apex dealer.

Product	Warranty period
CPAP devices	2 years
Water chamber	6 months
Mask systems	6 months
Disposable products	None

Appendix A: EMC Information

Guidance and Manufacturer's Declaration- Electromagnetic Emissions:

This device is intended for use in the electromagnetic environment specified below. The user of this device should make sure it is used in such an environment.

Emissions Test	Compliance	Electromagnetic Environment-Guidance
RF emissions CISPR 11	Group1	The device uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.
RF emissions CISPR 11	Class B	The device is suitable for use in all establishments, including domestic establishments and those directly connected to the public low-voltage power supply network.
Harmonic emissions IEC61000-3-2	Class A	
Voltage fluctuations / Flicker emissions IEC61000-3-3	Complies	

Guidance and Manufacturer's Declaration- Electromagnetic Immunity:

This device is intended for use in the electromagnetic environment specified below. The user of this device should make sure it is used in such an environment.

Immunity Test	IEC60601 test level	Compliance	Electromagnetic Environment-Guidance
Electrostatic Discharge (ESD) IEC61000-4-2	±6kV contact ±8kV air	±6kV contact ±8kV air	Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30 %.
Electrical fast transient/ burst IEC61000-4-4	±2kV for power supply line ±1kV for input/out line	±2kV for power supply line ±1kV for input/out line	Mains power quality should be that of atypical commercial or hospital environment
Surge IEC61000-4-5	±1kV for differential mode ±2kV for common mode	±1kV for differential mode ±2kV for common mode	Mains power quality should be that of atypical commercial or hospital environment.
Voltage dips, short interruptions and voltage variations on power supply input lines IEC61000-4-11	<5 % U_T (>95 % dip in U_T) for 0,5 cycle 40 % U_T (60 % dip in U_T) for 5 cycles 70 % U_T (30 % dip in U_T) for 25 cycles <5 % U_T (>95 % dip in U_T) for 5 sec	<5 % U_T (>95 % dip in U_T) for 0,5 cycle 40 % U_T (60 % dip in U_T) for 5 cycles 70 % U_T (30 % dip in U_T) for 25 cycles <5 % U_T (>95 % dip in U_T) for 5 sec	Mains power quality should be that of atypical commercial or hospital environment. If the user of this device requires continued operation during power mains interruptions, it is recommended that the device be powered from an uninterruptible power supply or a battery.
Power frequency (50/60Hz) magnetic field IEC61000-4-8	3 A/m	3 A/m	Power frequency magnetic fields should be at levels characteristic of atypical location in a typical commercial or hospital environment.

NOTE: U_T is the a.c. mains voltage prior to the application of the test level

Guidance and Manufacturer’s Declaration - Electromagnetic Immunity:

This device is intended for use in the electromagnetic environment specified below. The user of this device should make sure it is used in such an environment.

Immunity Test	IEC60601 test level	Compliance	Electromagnetic Environment-Guidance
<p>Conducted RF IEC 61000-4-6</p> <p>Radiated RF IEC 61000-4-3</p>	<p>3Vrms150 kHz to 80 MHz outside ISM bands^a</p> <p>3 V/m 80 MHz to 2.5 GHz</p>	<p>3 Vrms</p> <p>3 V/m</p>	<p>Portable and mobile RF communications equipment should be used no closer to any part of this device, including cables, than there commended separation distance calculated from the equation applicable to the frequency of the transmitter.</p> <p>Recommended separation distance $d = 1.2\sqrt{P}$ 150kHz to 80MHz</p> <p>$d = 1.2\sqrt{P}$ 150kHz to 80MHz $d = 2.3\sqrt{P}$ 80 MHz to 2.5G MHz</p> <p>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).^b</p> <p>Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey ^c, should be less than the compliance level in each frequency range^d.</p> <p>Interference may occur in the vicinity of equipment marked with the following symbol:</p> 
<p>NOTE 1: At 80 MHz and 800 MHz, the higher frequency range applies.</p> <p>NOTE 2: These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.</p>			

- a) The ISM (industrial, scientific and medical) bands between 150 kHz and 80 MHz are 6,765 MHz to 6,795 MHz; 13,553 MHz to 13,567 MHz; 26,957 MHz to 27,283 MHz; and 40,66 MHz to 40,70 MHz.
- b) The compliance levels in the ISM frequency bands between 150 kHz and 80 MHz and in the frequency range 80 MHz to 2.5 GHz are intended to decrease the likelihood that mobile/portable communications equipment could cause interference if it is inadvertently brought into patient areas. For this reason, an additional factor of 10/3 is used in calculating the recommended separation distance for transmitters in these frequency ranges.
- c) Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the device is used exceeds the applicable RF compliance level above, the device should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as reorienting or relocating the device.
- d) Over the frequency range 150 kHz to 80 MHz, field strengths should be less than 3 V/m.

Recommended separation distances between portable and mobile RF communications equipment and this device:

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

Rated maximum output power of transmitter W	Separation distance according to frequency of transmitter m		
	150 kHz to 80 MHz $d = 1.2\sqrt{P}$	80 MHz to 800 MHz $d = 1.2\sqrt{P}$	800 MHz to 2,5 GHz $d = 2.3\sqrt{P}$
0.01	0.12	0.12	0.23
0.1	0.38	0.38	0.73
1	1.2	1.2	2.3
10	3.8	3.8	7.3
100	12	12	23

For transmitters rated at a maximum output power not listed above, the recommended separation distance *d* in meters (m) can be estimated using the equation applicable to the frequency of the transmitter, where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer.

Note 1: At 80 MHz and 800 MHz, the separation distance for the higher frequency range applies.

Note 2: These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects, and people.

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