 

Model: UltraMint



***ULTRASONIC SCALER***

**USER MANUAL**

Version: 04

IFU-6635004

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Size：197mmX140mm

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# 1. Overview

## 1.1 Content



1. Main Unit
2. Power Adapter
3. Foot Pedal
4. Handpiece Support
5. Handpiece
6. Tip
7. External Water Supply Hose
8. Wrench

## 1.2 Packing list and coding

|  |  |  |
| --- | --- | --- |
| Main unit (1pcs)  6651026 | Power adapter (1pcs)  6651030 | Foot pedal (1pcs)  6651027 |
| Handpiece support (1pcs)  6604008 | External water supply hose (1pcs)  6641006 | Handpiece (1pcs)  Store in the Instrument Box  6651005 |
| Wrench (1pcs)  Store in the Instrument Box  6651010 | Tips (6pcs)  Store in the following box  G1(6626001), G2(6626002), G4(6626004), P1(6626006),  P3(6626007), E4(6626016) |  |

# 2. Symbol instruction

|  |  |
| --- | --- |
|  | If the instructions are not followed properly, operation may lead to hazards for the product or the user/patient. |
|  | Additional information, explanation of operation and performance. |
|  | Serial number |
|  | Class II equipment |
|  | Type B applied part |
|  | Alternating current |
|  | Dispose of in accordance with the WEEE directive |
|  | Keep dry |
|  | Can be autoclaved up to a maximum temperature of 134° Celsius |
|  | Temperature limitation |
|  | Humidity limitation |
|  | Atmospheric pressure limitation |
|  | Foot Pedal |
|  | Water inlet pressure: 0.01-0.5MPa |
| IPX0 | Ordinary equipment |
| IPX1 | Anti-drip equipment |
|  | Used indoor only |
|  | Catalogue number |
|  | Manufacturer |
|  | Date of manufacture |
|  | Lot of manufacture |
|  | Authorized Representative in the European Community |
|  | Manufacturer’s LOGO |
|  | Follow instructions for use |
|  | Washer-disinfector for thermal disinfection |
|  | Handpiece water control |

# 3. Foreword

## 3.1 Scope of application

UltraMint is an ultrasonic scaler intended for use during scaling, endodontic treatment and periodontal therapy to remove calculus deposits and stains from teeth by application of an ultrasonic vibrating scaler tip to the teeth.

This device must only be used in hospital environments, clinics or dental offices by trained and qualified dental personnel and not used in the oxygen-rich environment.

## 3.2 Contraindications

3.2.1 The patient who has hemophilia is not allowed to use in this equipment.

3.2.2 The patient or doctor who equips with cardiac implantable electronic devices is forbidden to use this equipment.

3.2.3 The heart disease patient, pregnant woman and children should be cautious to use the equipment.



Read the following warnings before use:

1. The device must not be placed in humid surroundings or anywhere where it can come into contact with any type of liquids.

2. Do not expose the device to direct or indirect heat sources. The device must be operated and stored in a safe environment.

3. The device requires special precautions with regard to electromagnetic compatibility (EMC) and must be installed and operated in strict compliance with the EMC information. In particular, do not use the device in the vicinity of fluorescent lamps, radio transmitters, remote controls and do not use this system near the active HF Surgical Equipment in the hospital. Portable RF communications equipment (including peripherals such as antenna cables and external antennas) should be used no closer than 30 cm (12 inches) to any part of the ultrasonic scaler, including cables specified by the manufacturer. Otherwise, degradation of the performance of this equipment could result. Do not charge, operate or store at high temperatures. Comply with the specified operating and storage conditions.

4. Gloves are compulsory during treatment.

5. If irregularities occur in the device during treatment, switch it off. Contact the agency.

6. Never open or repair the device yourself, otherwise, void the warranty.

# 4. Installation

## 4.1 Front view



## 4.2 Back view



## 4.3 Installation steps

|  |  |
| --- | --- |
| **4.3.1 Install the handpiece support**  Install the handpiece support to the main unit first.    Clamp the handpiece cord to the hanpiece support.    **4.3.2** **Install power adapter and foot pedal**  As shown in the figure, plug the power adapter and foot pedal into the corresponding sockets. Turn the power switch to the off state (press the side marked "O" on the switch), and insert the adapter power plug into the power supply to supply power to the main unit. | * The power adapter must be plugged into the main unit before being plugged into the power supply.   **4.3.3 Install external water supply hose**  Install the external water supply hose to the external water hose connector on the main unit as shown below.     1. Unscrew the lock nut on the external water hose connector counterclockwise. 2. Put the lock nut on the external water supply hose. 3. Insert the external water supply hose to the external water hose connector. 4. Tighten the lock nut clockwise to complete the installation of the external water supply hose. 5. Connect the other end of the external water supply hose to the clean water source. 6. See the next page for installation diagram. |
| The water source pressure should be in the range of 0.01-0.5MPa. |  |

## 4.4 Handpiece installation and removal

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| --- | --- |
| **4.4.1 Handpiece installation and removal**  Install and remove the handpiece as shown below.  Note that before installation, align the direction mark on the handpiece with the direction mark on the connector of handpiece cord, and then insert the handpiece into the connector of the handpiece cord.    **4.4.2 Lamp bead protective shield and light guide and lamp bead removal and installation**    Disassemble the light guide and lamp bead from the handpiece, before Cleaning, disinfection and sterilization handpiece. |  |

## 4.5 Tip installation and removal

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| --- | --- |
| **4.5.1 Tip installation**   1. Screw the tip to the handpiece. 2. Align the gap of the wrench with the tip. 3. Rotate the wrench clockwise until it turns 90 degrees. | **4.5.2 Tip removal**   1. Align the gap of the wrench with the tip. 2. Rotate the wrench counterclockwise until the tip becomes loose. 3. Unscrew the tip from the handpiece |

# 5. Product function and use

## 5.1 Operation panel instructions

|  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **5.1.1 Working mode**    This equipment can provide three working modes: Scaling mode, periodontal mode and endodontic mode.  Press the “” key to select the scaling mode, the indicator on the corresponding button is lit, select the tip marked with "G" at the tip when using it.  Press the “” key to select the periodontal mode, the indicator on the corresponding button is lit, select the tip marked with "P" at the tip when using it.  Press the “” key to select the endodontic mode, the indicator on the corresponding button is lit, select the tip marked with "E" at the tip when using it. | **5.1.2 Power adjustment**    The device can provide 10 (1-10) graduations of power configurations in each mode, and the user can adjust it according to the needs of use.  Press "-" key to decrease the power, press "+" key to increase the power.  "Power display window" and " power indicator" correspond to the power level.  Users can also adjust the power by tapping with their fingers or sliding “” left and right.   |  |  | | --- | --- | | Operating Power of the equipped tips | | | Tip model | Power | | G1 | 1-10(G) | | G2 | 1-10(G) | | G4 | 1-10(G) | | P1 | 1-10(P) | | P3 | 1-6(P) | | E4 | 1-6(E) | |

## 5.2 Function mode and use

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| --- | --- |
| **Water supply function and use**  Connect the other end of the external water supply hose to the connector that can provide water for the device.  **Water flow adjustment**  Adjust the water flow through the water control knob on the handpiece cord. |  |

## 5.3 Operating the device

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| --- | --- |
| 5.3.1 Install the device correctly according to the installation steps. The operator is facing the device, and the water control knob on the handpiece cord is adjusted to the maximum.  5.3.2 Press the power switch on the main unit. At this time, the scaling mode  indicator is lit, the power display window shows 05, the power indicator lights half.  5.3.3 Select the appropriate working tip as required, and tighten it to the handpiece by using the provided wrench.  5.3.4 Hold the handpiece according to the using habits, generally adopt the pen holding position.  5.3.5 When the device is working normally, the frequency is extremely fast. When scaling, ensure that the tip vibrates normally and the water is atomized normally, then gently touch the tooth surface and reciprocate at a certain speed to remove the dental calculus. Keep the water flow smoothly to avoid tip overheat. Never use excessive force or stay for too long when cleaning the teeth.  5.3.6 Power adjustment: Adjust the power depending on the application. Generally, the medium power is enough, and the output power can be adjusted at any time in clinical practice according to the sensitivity of the patient's teeth and the hardness of calculus. | 5.3.7 Water flow adjustment: Step on the foot pedal to activate the tip vibration. Turn the water control knob on the handpiece cord to form water spray to cool the tip and the cleaned tooth surface.  5.3.8 During clinical use, the cusp of the tip should not be in vertical contact with the tooth surface, and heavy pressure should not be applied in order to avoid damage to the tooth and the tip.  5.3.9 After the operation is completed, adjust the water flow of the handpiece to the maximum, let the device work for about 30 seconds to rinse the handpiece and the tip, and then remove the handpiece and the tip for cleaning, disinfection and sterilization. |

|  |  |
| --- | --- |
| 1. Keep the device clean before and after use.  2. Before each use, always check that the device is placed on a secure and flat surface. And adjust the water flow to the maximum, and then start the device to work for about 10 seconds in order to eliminate the residual liquid in the liquid circuit.  3. The operator should take sufficient protection (such as wearing goggles, face mask, etc.) to prevent cross infection. Use of an antiseptic mouth rinse prior to procedure and use of a high volume evacuator during the scaling procedure are recommended.  4. The use of the product must meet the requirements of the relevant operating specifications and relevant regulations of the treatment department, and is limited to the use of trained professionals (such as dentists) in the hospital or clinic.  5. Do not screw the tip or pull out the handpiece while the machine is running.  6. The tip must be tightened and there must be fine spray coming out from the tip when operating. Refer to the user manual of tips for detailed operating instructions.  7. Change a new one when the tip or handpiece is damaged or there are visible signs of wear.  8. Do not twist or rub the tip.  9. Do not use impure water sources, and never use saline instead of pure water. | 10. If use the water source without hydraulic pressure, the water surface should be 1 meter higher than the head of the patient.  11. Do not pull the handpiece cord with force to avoid damage to the tail wire.  12. Do not knock or rub the handpiece.  13. After operating, turn off the power switch and pull out the power plug.  14. Always use original parts. Using non original instruments may damage the device, and operator or patient may be injured.  15. No modification shall be made on this product.  16. The ultrasound power must be adjusted in accordance with the tip used and the required treatment.  17. Always check that the cords or cables will not rub against the front face during the operation since this could eventually modify the selected settings. |

## 5.4 Advanced settings

|  |  |
| --- | --- |
| **5.4.1"Cleaning" mode**  It is recommended to flush the liquid circuit of the device after scaling at least once a day. The "Cleaning" mode allows for cleaning the liquid circuit in order to reduce the accumulation of crystals and bacteria in the liquid circuit.  **Steps:**   1. Install the water hose to the water inlet correctly. 2. Press and hold “” and “” at the same time for about 2 seconds. At this time, the buzzer beeps once and enters the “Cleaning” mode. At this time, the LED digital tube on the panel alternately displays “CL” and cleaning time. The default cleaning time is 30 Seconds, press "+" or "-" key to adjust. The adjustment range is 10 ~ 60 seconds. 3. Step on the foot pedal to start cleaning the liquid circuit. At this time, the foot pedal can be released. 4. After the cleaning countdown, the device will automatically stop and exit the "Cleaning" mode. During the cleaning process, step on the foot pedal again or press the "" key to stop cleaning and exit the "Cleaning" mode. | **5.4.2 Handpiece LED light delay adjustment**  The handpiece LED light will be lit during the operation. The device will stop running after the foot pedal is released. The handpiece LED light will be delayed for a certain time before extinguishing. The default delay time is 10 seconds, and the delay time can be adjusted as needed.  **Steps:**  Press and hold the “” key and the “” key at the same time for about 2 seconds, at this time the, the buzzer beeps once and enter the “Cleaning” mode, and then press the “” key to enter the handpiece LED light adjustment state, at this time, the LED digital tube display the delay time. The default is 10 seconds, you can press the "+" or "-" key to proceed adjustment. The adjustment range is 10 ~ 20 seconds. Press the "G" button or no operation for about 5 seconds to exit the setting of the handpiece LED light delay. |

# 6. Cleaning, Disinfection and Sterilization

## 6.1 Foreword

The parts for clinical application contamination are the outer surfaces of the handpiece, tip and wrench. For hygiene and sanitary safety purpose, these components must be cleaned, disinfected and sterilized before each usage to prevent any contamination. This concerns the first use as well as the subsequent uses. Comply with your national guidelines, standards and requirements for cleaning, disinfection and sterilization.

Reprocessing procedures have only limited implications to these dental instruments. The limitation of the numbers of reprocessing procedures is therefore determined by the function / wear of the device. From the processing side there is no maximum number of allowable reprocessing. The device should no longer be reused in case of signs of material degradation.

In case of damage, the device should be reprocessed before sending back to the manufacturer for repair.

## 6.2 General recommendations

* The user is responsible for the sterility of the product for the first cycle and each further usage as well as for the usage of damaged or dirty instruments, where applicable after sterility.
* For your own safety, please wear personal protective equipment (gloves, safety glasses, etc.).
* Use only a disinfecting solution which is approved for its efficacy (VAH/DGHM-listing, CE marking, and FDA approval) and in accordance with the DFU of the disinfecting solution manufacturer.
* The water quality has to be convenient to the local regulations especially for the last rinsing step or with a washer-disinfector.
* Thoroughly clean and wash the components before autoclaving.
* Do not use bleach or chloride disinfectant materials.

## 6.3 Autoclavable components

|  |  |  |  |
| --- | --- | --- | --- |
| **Autoclavable Components** | | | |
| Handpiece | | Tip | Wrench |
| * Only the components above can be autoclaved. * Before first use and after each use, sterilize the above components. | | | |
| **Reprocessing Instructions** | | | |
| **Preparation at the Point of Use:** | Before cleaning, disconnect the handpiece and tips from the main unit. Disassemble the light guide and lamp beads from the handpiece. Refer to Chapter 4.4 and 4.5 of this manual for disassembly instructions. Remove gross contaminations from the components with code water (<40°C) immediately after use. Don’t use a fixating detergent or hot water (>40°C) as this can cause the fixation of residuals which may influence the result of the reprocessing process.  Store the instruments in a humid surrounding.    Do not submerge the components or wipe them with any of the following functional water (acidic electrolyzed water, strong alkaline solution, or ozone water), medical agents (glutaral, etc.), or any other special types of water or commercial cleaning liquids. Such liquids may result in metal corrosion and adhesion of the residual medical agents to the components. | | |
| **Transportation:** | Safe storage and transportation to the reprocessing area to avoid any damage and contamination to the environment. | | |
| **Preparation for Decontamination:** | The devices must be reprocessed in a disassembled state.    Observe suitable personal protective measures. | | |
| **Pre-Cleaning:** | Do a manual pre-cleaning, until the components are visually clean. Submerge the components in a cleaning solution and flush the lumens with a water jet pistol with cold tap water for at least 10 seconds. Clean the surfaces with a soft bristle brush. | | |
| **Cleaning:** | Regarding cleaning/disinfection, rinsing and drying, it is to distinguish between manual and automated reprocessing methods. Preference is to be given to automated reprocessing methods, especially due to the better standardizing potential and industrial safety.  Automated Cleaning:  Carefully put the components into the washer-disinfector on a tray and set the parameters as follows, then start the program:   * 4 mins pre-washing with cold water (<40°C); * emptying * 5 mins washing with a mild alkaline cleaner at 55°C; * emptying * 3 mins neutralizing with warm water (40°C); * emptying * 5 mins intermediate rinsing with warm water (40°C); * emptying   *The automated cleaning processes have been validated by using 0.5% neodisher MediClean forte (Dr. Weigert).*  Note Acc. to EN ISO 17664 no manual reprocessing methods are required for these devices. If a manual reprocessing method has to be used, please validate it prior to use.     * Use only approved washer-disinfectors according to EN ISO 15883, maintain and calibrate it regularly. * Follow instructions and observe concentrations given by the manufacturer (see general recommendations). | | |
| **Disinfection:** | Automated Thermal Disinfection in washer/disinfector under consideration of national requirements in regards to A0 value (see EN ISO 15883).  A disinfection cycle of 5 mins disinfection at 93°C has been validated for the device to achieve an A0 value of 3000.  After cleaning, the instruments should be automated disinfected immediately. A manual disinfection is not recommended. Please use fully demineralized water. | | |
| **Drying:** | Automated Drying:  Drying the instruments according to drying cycle of washer/disinfector. If needed, additional manual drying can be performed through lint free towel. Insufflate cavities of instruments by using sterile compressed air. | | |
| **Functional Testing, Maintenance:** | Visual inspection for cleanliness of the instruments and reassembling. Functional testing according to instructions of use. If necessary, perform reprocessing process again until the instrument is visibly clean.  Before packaging and autoclaving, make sure that the components have been maintained according to the manufacturer’s instruction. | | |
| **Packaging:** | Pack the instruments in an appropriate packaging material for sterilization.     * Check the validity period of pouch given by the manufacturer to determine the shelf life. * Use pouches which resist to a temperature up to 141℃ and in accordance with EN ISO 11607. | | |
| **Sterilization:** | Sterilization of instruments by applying a fractionated pre-vacuum steam sterilization process (according to EN 285/EN 13060/EN ISO 17665) under consideration of the respective country requirements.  Minimum requirements: 5 mins at 134 °C.  Maximum sterilization temperature: 137°C.  Drying time: at least 8mins.  Flash sterilization is not allowed on lumen instruments!     * Use only approved autoclave devices according to EN 13060 or EN 285. * Respect the maintenance procedure of the autoclave device given by the manufacturer. * Use only this recommended sterilization procedure. * Control the efficiency (packaging integrity, no humidity, color change of sterilization indicators, physicochemical integrators, digital records of cycles parameters). * The sterilization procedure must comply with EN ISO 17665. * Waiting for cooling before touching. | | |
| **Storage:** | Storage of sterilized instruments in a dry, clean and dust free environment at modest temperatures, refer to label and instructions for use.     * Sterility cannot be guaranteed if packaging is open, damaged or wet. * Check the packaging before using it (packaging integrity, no humidity and validity period). | | |
| **Reprocessing validation study information:** | The above-mentioned reprocessing process (cleaning, disinfection, sterilization) has been successfully validated. Refer to cleaning/disinfection validation test reports No. RDS2020D0076 001 and RDS2020D0073 001; sterilization validation test reports No. RDS2020S0084 001 and RDS2020S0081 001. | | |
| * Before sterilization, please remove the tip. * Make sure that the handpiece is intact and not damaged before sterilization or use, and do not apply any protective oil on the handpiece. * The two O-rings on the handpiece cord (and the handpiece insertion point) will be subject to force and wear during insertion and removal. Users can apply dental lubricant to the O-ring in daily use. If the O-ring is damaged or severely worn, causing water leakage or loose connection, please replace the O-ring. * The instructions provided above have been validated by the manufacturer of the medical device as being capable of preparing a medical device for use. It remains the responsibility of the processor to ensure that the processing, as actually performed using equipment, materials and personnel in the processing facility, achieves the desired result. This requires verification and/or validation and routine monitoring of the process. Likewise, any deviation by the processor from the instructions provided should be properly evaluated for effectiveness and potential adverse consequences. | | | |

## 6.4 Disinfection components

|  |  |  |
| --- | --- | --- |
| **Disinfection components** | | |
| Main unit | Power adapter | Foot pedal |
| Handpiece support | External water supply hose |  |
| Wipe all the surfaces with a cloth lightly moistened with Ethanol for Disinfection (Ethanol 70 to 80 vol%) at least 2mins, repeat for 5 times. | | |
| * Do not use anything except Ethanol for Disinfection (Ethanol 70 to 80 vol%). * Do not use too much ethanol as it’s going into machine and damage the components inside. * Do not spray any liquid directly on the machine. Do not allow any moisture to get into the machine. | | |

# 7. Maintenance

When the device is not in use, the power switch should be turned off and the power plug should be unplugged. When not in use for a long time, it should be powered and watered once a month for about 5 minutes each time.

# 8. Troubleshooting

When trouble is found, check the following points before contacting your distributor. If none of these are applicable or the trouble is not remedied even after action has been taken, the product may have failed. Contact your distributor.

|  |  |  |
| --- | --- | --- |
| Malfunction | Causes | Methods |
| The LED digital tube and the corresponding button indicator on the rear panel are not lit after power on. | Fuse is broken | Replace T0.5AL 250V fuse in power adapter.  Replace T1.6AL 250V fuse in main unit. |
| Poor contact of power plug | Plug in the power plug properly |
| After the power is turned on and the foot pedal is pressed, the tip does not vibrate and there is no water spray | Poor foot pedal contact | Plug in the foot pedal properly |
| After the power is turned on and the foot pedal is pressed, the tip does not vibrate but there is water spray | The tip is loose | Tighten the tip properly |
| Tail wire failure | Contact your local distributor or our company |
| Handpiece failure | Contact your local distributor or our company |
| The connecting plug of the tail wire and the circuit board is loose | Contact your local distributor or our company |
| After the power is turned on and the foot pedal is pressed, the tip vibrates but there is no water spray | The water control knob on the handpiece is not on | Turn on the water control knob on the handpiece |
| There are impurities in the solenoid valve | Contact your local distributor or our company |
| Blocked waterway | Drain the waterway with dental gun |
| There is still water spray after releasing the foot pedal | There are impurities in the solenoid valve | Contact your local distributor or our company |
| Handpiece heat | Water control knob is switched too small | Turn up the water control knob |
| The water spray is too small | Water control knob is switched too small | Turn up the water control knob |
| Water pressure is not enough | Increase water pressure |
| Blocked waterway | Drain the waterway with dental gun |
| The tip vibration is weakened | Working tip is loose | Tighten the tip |
| Working tip is broken | Replace the tip |
| Control panel malfunction | Control panel circuit board damaged | Contact your local distributor or our company |
| Water seepage at the connection between the handpiece and the handpiece cord | Damaged waterproof O-ring | Replace waterproof O-ring |
| The root canal file does not vibrate or the file holder makes noise | Nut is not tightened | Tighten the nut |
| Root canal adapter damaged | Replace the file holder |
| LED light is not on | LED light is damaged | Replace the LED light |
| Poor contact | Check the circuit |

# 9. Technical Data

|  |  |
| --- | --- |
| Manufacturer | Changzhou Sifary Medical Technology Co.Ltd. |
| Model | UltraMint |
| Box dimensions | 395mm×230mm×90mm |
| Total weight | 2.8kg |
| Power supply | ~220-240V 50/60Hz |
| Main unit input | ~25V 50/60Hz 1.3A |
| Output primary tip vibration excursion | 1μm-200μm |
| Output tip vibration frequency | 25kHz~42kHz |
| Output half-excursion force | 0.1N~2N |
| Output power | 3W~20W |
| Power adapter fuse | T0.5AL250V |
| Main unit fuse | T1.6AL250V |
| Water pressure at inlet | 0.1bar~5bar (0.01MPa~0.5MPa) |
| Electrical safety class | ClassⅡ |
| Applied part | B |
| Ingress protecting rating | Ordinary equipment (IPX0),  Foot pedal (IPX1) |
| AP / APG type equipment | None |
| Anti-defibrillation application part | None |
| Operating mode | Continuous operation |
| Operating conditions | Use: in enclosed spaces  Ambient temperature: 5°C ~ 40 °C  Relative humidity: <80%  Altitude max. 3000m |
| Transport and storage conditions | Ambient temperature: -20 °C ~ +55 °C  Relative humidity: 20% ~ 80 %  Atmospheric pressure: 70kPa ~ 106kPa |

# 10. EMC Tables

|  |  |  |
| --- | --- | --- |
| **Guidance and manufacturer’s declaration – electromagnetic emissions** | | |
| The **UltraMint** is intended for use in the electromagnetic environment specified below. The customer or the user of the **UltraMint** should assure that it is used in such an environment. | | |
| **Emissions test** | **Compliance** | **Electromagnetic environment - guidance** |
| RF emissions CISPR 11 | Group 1 | The **UltraMint** 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 **UltraMint** is suitable for use in all establishments, including domestic establishments and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes. |
| Harmonic emissions IEC61000-3-2 | Class A |
| Voltage fluctuations/flicker emissions IEC 61000-3-3 | Complies |

|  |  |  |  |
| --- | --- | --- | --- |
| **Guidance and manufacturer’s declaration – electromagnetic immunity** | | | |
| The **UltraMint** is intended for use in the electromagnetic environment specified below. The customer or the user of the **UltraMint** should assure that it is used in such an environment. | | | |
| **Immunity test** | **IEC 60601 test level** | **Compliance level** | **Electromagnetic environment - guidance** |
| Electrostatic discharge (ESD) IEC 61000-4-2 | +/- 8 kV contact  +/- 2 kV, +/- 4 kV, +/- 8 kV, +/- 15 kV air | +/- 8 kV contact  +/- 2 kV, +/- 4 kV, +/- 8 kV, +/- 15 kV 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  transients/bursts  IEC 61000-4-4 | ±2kV  100kHz repetition frequency | ±2kV  100kHz repetition frequency | Mains power quality should be that of a typical commercial or hospital environment. |
| Surge  IEC 61000-4-5 | Line to line: ±0.5kV, ±1kV  Line to earth: ±0.5kV, ±1kV, ±2kV | Line to line: ±0.5kV, ±1kV  Line to earth: ±0.5kV, ±1kV, ±2kV | Mains power quality should be that of a typical commercial or hospital environment. |
| Voltage dips  IEC 61000-4-11  Voltage interruptions  IEC 61000-4-11 | 0% UT; 0.5 cycle  at 0°, 45°, 90°, 135°, 180°, 225°, 270°, and 315°  0% UT; 1 cycle and 70% UT; 25/30 cycles  sine phase at 0°  0% UT; 250/300 cycle | 0% UT; 0.5 cycle  at 0°, 45°, 90°, 135°, 180°, 225°, 270°, and 315°  0% UT; 1 cycle and 70% UT; 25/30 cycles  sine phase at 0°  0% UT; 250/300 cycle | Mains power quality should be that of a typical commercial or hospital environment. If the user of devices require continued operation during power mains interruptions, it is recommended that devices be powered form an uninterruptible power supply or a battery |
| Rated Power frequency magnetic field IEC 61000-4-8 | 30 A/m  50Hz or 60Hz | 30 A/m  50Hz or 60Hz | Power frequency magnetic field should be at levels characteristic of a typical location in a typical commercial or hospital environment. |
| Note: UT: rated voltage(s); E.g. 25/30 cycles means 25 cycles at 50Hz or 30 cycles at 60Hz | | | |

|  |  |  |  |
| --- | --- | --- | --- |
| **Guidance and manufacturer’s declaration – electromagnetic immunity** | | | |
| The **UltraMint** is intended for use in the electromagnetic environment specified below. The customer or the user of the **UltraMint** should assure that it is used in such an environment. | | | |
| **Immunity test** | **IEC 60601 test level** | **Compliance level** | **Electromagnetic environment - guidance** |
| Conducted dis-turbances induced by RF fields  IEC 61000-4-6 | 3 V  0.15 MHz – 80 MHz, 6 V in ISM bands be-tween 0.15 MHz and 80 MHz, 80 % AM at 1 kHz  3 V/m, 80 MHz – 2,7 GHz, 80 % AM at 1 kHz  See the RF wireless communication equipment table in "Recommended minimum separation distances" | 3 V | Portable and mobile RF communications equipment should be usedno closer to any part of the **UltraMint**, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter.  **Recommended minimum separation distances**  See the RF wireless communication equipment table in "Recommended minimum separation distances" |
|  |  |
| Radiated RF EM fields  IEC 61000-4-3 | 3V/m |
| Proximity fields from RF wireless communication equipment | Complies |
| IEC 61000-4-3 |

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
| **Recommended minimum separation distances** | | | | | | |
| Nowadays, many RF wireless equipments have being used in various healthcare locations where medical equipment and/or systems are used. When they are used in close proximity to medical equipment and/or systems, the medical equipment and/or systems’ basic safety and essential performance may be affected. The **UltraMint** has been tested with the immunity test level in the below table and meet the related requirements of IEC 60601-1-2:2014. The customer and/or user should help keep a minimum distance between RF wireless communications equipments and the **UltraMint** as recommended below. | | | | | | |
| **Test frequency**  **(MHz)** | **Band**  **(MHz)** | **Service** | **Modulation** | **Maximum power**  **(W)** | **Distance**  **(m)** | **Immunity test level**  **(V/m)** |
| **385** | **380-390** | **TETRA 400** | **Pulse modulation**  **18Hz** | **1.8** | **0.3** | **27** |
| **450** | **430-470** | **GMRS 460**  **FRS 460** | **FM**  **± 5 kHz deviation**  **1 kHz sine** | **2** | **0.3** | **28** |
| **710** | **704-787** | **LTE Band 13, 17** | **Pulse modulation**  **217Hz** | **0.2** | **0.3** | **9** |
| **745** |
| **780** |
| **810** | **800-960** | **GSM 800/900,**  **TETRA 800,**  **iDEN 820,**  **CDMA 850,**  **LTE Band 5** | **Pulse modulation**  **18Hz** | **2** | **0.3** | **28** |
| **870** |
| **930** |
| **1720** | **1700-1990** | **GSM 1800;**  **CDMA 1900;**  **GSM 1900;**  **DECT;**  **LTE Band 1, 3,**  **4, 25; UMTS** | **Pulse modulation**  **217Hz** | **2** | **0.3** | **28** |
| **1845** |
| **1970** |
| **2450** | **2400-2570** | **Bluetooth,**  **WLAN,**  **802.11 b/g/n,**  **RFID 2450,**  **LTE Band 7** | **Pulse modulation**  **217Hz** | **2** | **0.3** | **28** |
| **5240** | **5100-5800** | **WLAN 802.11**  **a/n** | **Pulse modulation**  **217Hz** | **0.2** | **0.3** | **9** |
| **5500** |
| **5785** |



1. Use of accessories and cables other than those specified or provided by the manufacturer of **UltraMint** could result in increased electromagnetic emissions or decreased electromagnetic immunity of **UltraMint** and result in improper operation.

**Cable information:**

|  |  |  |  |
| --- | --- | --- | --- |
| Cable Name | Cable Length (m) | Shielded or not | Remark |
| Power adapter input cable | 1.4 | NO | / |
| Power adapter output cable | 1.4 | NO | / |
| Pedal cable | 2.5 | NO | / |
| Handpiece cord | 2.0 | NO | / |

1. Use of **UltraMint** adjacent to or stacked with other equipment should be avoided because it could result in improper operation. If such use is necessary, **UltraMint** and the other equipment should be observed to verify that they are operating normally.

# 11. Statement

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| **Service Life**  The service life of **UltraMint** series products is 5 years. |
| **Maintenance**  MANUFACTURE will provide circuit diagrams, component part lists, descriptions, calibration instructions to assist to SERVICE PERSONNEL in parts repair. |
| **Disposal**  The package should be recycled. Metal parts of the device are disposed as scrap metal. Synthetic materials, electrical components, and printed circuit boards are disposed as electrical scrap. Please deal with them according to the local environmental protection laws and regulation. |
| **Rights**  All rights of modifying the product are reserved to the manufacturer without further notice. The pictures are only for reference. The final interpretation rights belong to CHANGZHOU SIFARY MEDICAL TECHNOLOGY CO., LTD. The industrial design, inner structure, etc, have claimed for several patents by SIFARY, any copy or fake product must take legal responsibilities. |



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