

ARFLOW[®] PROPHYLAXIS MASTER

INSTRUCTIONS FOR USE

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1. BEFORE USE

CONGRATULATIONS!

You are now the owner of this new EMS device!

Please read the instructions carefully before use \rightarrow

▲ TO AVOID the risk of electric shock, this equipment must only be connected to a mains supply with protective earth/grounding. This device uses a Class-I insulating system that requires protective earth.

FOR USA AND CANADA: GROUNDING RELIABILITY CAN ONLY BE ACHIEVED WHEN EQUIPMENT IS CONNECTED TO AN EQUIVALENT RECEPTACLE MARKED "HOSPITAL ONLY" OR "HOSPITAL GRADE".

- ▲ DO NOT modify this equipment and/or any of its accessories. No modification of any part of this medical device is allowed.
- ▲ DO NOT open the device. There are no serviceable parts inside.
- If any serious incident occurs that is directly or indirectly related to the device, report it immediately to the manufacturer and to the competent authority of your country and of where the patient is established (if different).
- Disconnect the mains plug from electrical outlet for the purposes of maintenance, in the case of malfunction or when the device is left unattended.
- Turn off the water inlet when not in use. The device is not equipped with Aquastop and the EG-110 water hose may disconnect or leak: risk of flooding.

The Instructions for Use of the device, as well as the Treatment Recommendations (FB-648) and Piezon Treatment Recommendations (FB-652), are provided in electronic format and are part of the product documentation. However, if you want these in hard copy, you can request one set free of charge on our website, by telephone or in writing, and receive it within 7 days.

- The Instructions for Use of the device (FB-618), as well as the Treatment Recommendations (FB-648) and Piezon Treatment Recommendations (FB-652), are available for download in PDF format at www.ems-instruction.com using the Product/Key Code FT-229. A PDF reader is required and, in case of need, it can be downloaded from the same web site.
- It is essential to first read and understand all the Instructions for Use of the device before operating it and using the related accessories. The Treatment Recommendations are an integral part of the device's Instruction for Use and each one document is complementary to the other. Always keep this documentation close at hand.
- We recommend that you visit our website regularly to consult and/or download the latest version of the documentation for your device at www.ems-instruction.com
- Please contact EMS technical support or your local EMS representative for further information and support.



1.1. Intended Use

The device is a fixed table top unit combining:

- PIEZON[®]: ultrasound scaler
- AIRFLOW[®]: air polishing technology

Intended for use in

PREVENTION, MAINTENANCE AND TREATMENT

during dental prophylaxis to remove biofilm and calculus from natural teeth, restorations and implants

1.2. Application Fields

Application on natural teeth including all smooth surfaces, pits, fissures and interproximal areas, dental restorations and dental implants.

AIRFLOW[®] applications include:

- Plaque removal for the placement of sealants
- Surface preparation prior to the bonding/cementation of inlays, onlays, crowns and veneers
- Surface preparation prior to placing composite restorations
- Effective plaque and stain removal for orthodontic patients
- Cleaning prior to the bonding of orthodontic brackets
- Cleaning the implant fixture prior to loading
- Stain removal for shade determination
- Plaque removal prior to a fluoride treatment
- Plaque and stain removal prior to a whitening procedure

PERIOFLOW[®] applications include:

- Maintenance of periodontal deep pockets up to 9 mm following initial treatment
- Removal of periodontal biofilm
- Cleaning of implants

PIEZON[®] applications include:

- Removal of supragingival calculus
- Removal of subgingival calculus
- Periodontal treatment
- Preparation of approximal cavities
- Luting tooth-shaded inlays and onlays with highly thixotropic, dual-curing cements
- Endodontics: preparation, cleaning and irrigation of root canals
- Endodontics: retrograde preparation of root canals
- Endodontics: condensing gutta-percha
- Endodontics: removal of crowns and bridges
- Restorative: Preparation of cavities
- Restorative: Cementation of restorations
- Restorative: Condensing of amalgams
- Restorative: Removal of crowns, bridges, inlays and posts



1.3. Intended Users

Only dental professionals must use this device by fully complying with their respective country's regulations, accident prevention measures, and strictly follow these instructions for use.



maintained only by persons who have been instructed in infection control, personal protection and patient safety.

The device must be prepared and Improper use (e.g. due to lack of hygiene or routine maintenance), non-compliance with our instructions, or the use of accessories and spare parts that are not approved by EMS invalidates all claims under warranty and any other claims.

No specific training other than initial professional training is required to use this medical device. The practitioner is responsible for performing the clinical treatments and for any dangers that may arise due to a lack of skill and/or training.

For optimal patient comfort, safety and efficiency, we suggest that you regularly follow our:

SWISS DENTAL ACADEMY Training Program



Professional product installation and product introduction by EMS certified person is highly recommended for optimal setup and reliability.

1.4. Patient Population

PIEZON[®] devices are intended for use on patients requiring dental treatment, including scaling (e.g. subgingival and supragingival calculus, stains), endo (e.g. root canal treatment), restorative (e.g. cavities, amalgams), periodontics and dental prophylaxis, regardless of age or gender.

AIRFLOW[®] devices are intended for use on patients requiring dental treatment, including cleaning and polishing of teeth (natural or implant) by the projection of water, air and dental powders onto the tooth surface, regardless of age or gender.

A This medical device is not intended for use on newborn (neonate) and infant (< 2 years old) patient populations.



1.5. Contraindications

▲ Treatments contraindications:

Suggestion for alternatives:

| AIRFLOW® and PERIOFLOW® | are contraindicated with | Patients with severe or unstable upper respiratory tract infections, chronic bronchitis/asthma ¹ . | PIEZON® |
|-------------------------|--------------------------|---|---|
| PERIOFLOW® | is contraindicated with | Pregnant and breastfeeding patients | AIRFLOW [®] and PIEZON [®] PS |
| PERICIPLOW® | is contraindicated with | Patients with severe inflammation and/or osteonecrosis. | AIRFLOW [®] PLUS and PIEZON [®] PS |
| PIEZON® | is contraindicated with | Patients with a cardiac pacemaker, defibrillators and any implantable electronic device. | AIRFLOW [®] PLUS |

The decision to use AIRFLOW[®] and/or PERIOFLOW[®] on contagious patients or on patients with risk of infection, has to be taken by the dentist/medical doctor on an individual basis following practitioner protection level, patient risk assessement and specific country regulations.

On patients under Bisphosphonate therapy, the decision to use AIRFLOW[®] and/or PERIOFLOW[®] has to be taken by the dentist/medical doctor depending on the oral health of the patient.

| ▲ AIRFLOW [®] powders cor | ntraindications: | | Suggestion for alternatives: |
|------------------------------------|-------------------------|--|--|
| CLASSIC powder | is contraindicated with | Low-salt diet patients. | AIRFLOW [®] PLUS |
| Flavored CLASSIC powder | is contraindicated with | Patients allergic to flavor aroma. | AIRFLOW [®] PLUS/PERIO and CLASSIC NEUTRAL |
| PLUS powder | is contraindicated with | Patients allergic to Chlorhexidine. | AIRFLOW® PERIO |
| PERIO & SOFT powder | is contraindicated with | Patients allergic to Glycine (Glycocoll). | AIRFLOW® PLUS |

1.6. Compatibility

This device is compatible with the following accessories:

| AIRFLOW [®] Powders | PLUS powders: DV-082, DV-086 series CLASSIC powders: DV-048 series PERIO and SOFT powders: DV-070, DV-071 series |
|--|--|
| AIRFLOW [®] Handpiece | EL-308 |
| PERIOFLOW [®] Handpiece | EL-354 |
| PIEZON [®] Handpieces | EN-060, EN-061 |
| PIEZON® Scaling and Periodontal instruments | PS, A, B, C, P, PSR, PSL, PL1, PL2, PL3, PL4, PL5, HPL3, DPL3, PI |
| PIEZON® Endodontics instruments | RT1, RT2, D, H, ESI, Files ISO 15, 20, 25, 30, 35, Endochuck 180°, 120° and 90°, RT3, RE2, BERUTTI |
| PIEZON® Conservative measures instruments | E, F, G, SP |
| PIEZON® Cavity preparation instruments | SM, PF, SD, VE, SB, SBD, SBM |

¹ Related to possible powder inhalation during AIRFLOW[®] treatment.



Applied Parts

The following items are Medical Device Applied Parts:

- AIRFLOW[®] (EL-308) Handpiece
- PERIOFLOW[®] (EL-354) Handpiece
- PIEZON[®] (EN-060 and EN-061) Handpieces

Applied Parts, under certain operating conditions, may exceed 41°C of temperature and reach a maximum temperature of 51°C.

1.7. General Precautions







USE EMS ACCESSORIES ONLY!

The use of any other accessories could lead to patient injury, malfunction or damage to the device

DO NOT use this device in the presence of flammable anesthetics or oxidizing gases (such as nitrous oxide (N2O) and oxygen) or in close proximity to volatile solvents (such as ether or alcohol), as explosion may occur.

DO NOT store the powder near acids or heat sources.

TAKE the following precautions to prevent any adverse events to the patient and/or to the user in case of electromagnetic disturbances:

- Always refer to the information listed in the chapter " Electromagnetic Compatibility".
- In case of a wireless pedal malfunction, presumably caused by electromagnetic disturbances, use the wired pedal instead.
- In case of a device malfunction, presumably caused by electromagnetic disturbances, first verify the cabling, and then move any portable RF communications equipment and mobile devices placed nearby as far away as possible to rule out interference.
- Stop using the device if the electromagnetic disturbances persist and contact EMS technical support for assistance.



2.INSTALLATION

2.1. Equipment included in the box

Check contents for any damage that may have occurred during transportation.



BIOFILM DISCLOSER

BIOFILM DISCLOSER³ DV-158

² Not for end point sterilisation

³ If available in your country.





- 1 EL-308: AIRFLOW[®] Handpiece
- 2 AB-470A/A: Easy Clean
- 3 Ultra FS ClasenUNO Cannula
- 4 EL-651: Cord gaskets
- 5 EI-600: Water filter
- 6 EL-599: Air filter



PIEZON 🖸

- 1 EN-060 : PIEZON® Handpiece
- 2 3x DS-016A : Instrument PS
- **3** 4x AB-340: Light guide



- 1 EL-354: PERIOFLOW[®] Handpiece AB-358/B Nozzle extractor (under)
- 2 10x AB-327A/A: PERIOFLOW[®] nozzle
- 3 6x DT-064: Instrument PI
- 4 DT-018: Flat wrench (on top)
- 5 DS-010: Endochuck 120°

⁴ AIRFLOW[®] application box FS-442 integrates ClasenUNO Cannula in the European Union. In the rest of the world, the reference is FS-447.



2.2. Step-by-step installation

Find an appropriate area to place the device.

Place the medical device (control unit) within the dental cabinet in a suitable position for your activity and leave enough free space to allow easy handling and proper aeration.

A Keep a minimum of 10 cm clearance around the unit. Do not stack it with other equipment.

The medical device must be placed on a secure and flat surface (with a maximum slope of 5 degrees).

Check for proper water and air supply lines.

Verify that your dental cabinet has a filtered tap water source and a compressed air source using air and water hoses EG-110 and EH-142, respectively.

U In case your cabinet water and air lines are not provided with the required hoses EG-110 and EH-142, a proper installation by qualified personnel is required. Call EMS Service for support.

▲ In order to prevent retro contamination, connect the cable to EN-1717 or DVGW⁵ compliant fluid sources.

Check for a proper and safe power grid.

- <u>This device uses a Class-I insulating system that requires protective earth.</u>
- Plug the unit only into an FI protected mains supply (FI = Residual current protection). For USA and Canada: connect only to a hospital-grade outlet.
- ▲ Check that the rated voltage of the device is suited for the local line voltage to prevent damaging the unit, risk of fire and electric shock.
- A The mains plug of the unit must be accessible at all times.

O DO NOT INSTALL the device in case your dental cabinet does NOT have protective earth. If you have any concerns about this, call EMS Service for on-site support by qualified personnel.

Be aware

▲ The use of cables and accessories other than those supplied by EMS may negatively affect EMC performance. Use only parts supplied by EMS.

A The device uses a low power radio, 8 dBm EIRP max, Bluetooth[®] 2.4 GHz, to communicate with the wireless pedal. Interference may occur in the vicinity of this equipment.

The Bluetooth® radio is automatically disabled (powered off) when a wired pedal is connected.

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 device, including cables. Otherwise, degradation of the performance of this equipment could result.

⁵ German Technical and Scientific Association for Gas and Water



Connect air and water hoses

Turn the device over and place it upside down.

Connect the air hose EH-142 to the cabinet/dental unit. Push the hose connector into the air jack firmly (it may be hard).

Connect the water hose EG-110 to the cabinet/dental unit.

To prevent retro contamination, connect the cable to an EN-1717 or DVGW compliant fluids source.

O NOT install the PIEZON[®] or NIGHT CLEANER bottle before connecting the air and water lines.

Pressure: 4.5 to 7 bar Dry air. Max. humidity: 1.032 g/m³ Filtration: max. 1 µm

Drinking water Pressure: 2 to 5 bar Salinity: max. 0.2% Temperature: 10°C to 30°C

Install accessories

Continue to keep the device upside down and disconnected from the power grid!





Check the cord connections



The handpiece cord is not fully connected.



The system is well connected & locked.

To disconnect the handpiece cord system, unlock the connection and pull at the same time.

Fix the device

You will find a "Master Screw" provided on the bottom center of the device.

Unscrew the Master Screw first and use it to secure the device firmly to a table or onto the AL-125 device support in your cabinet (the AL-125 part is available through our after-sales support and dealers).



Fix your device with the provided "Master Screw" in order to ensure that the unit cannot be removed without the use of a tool.

Check the position of the medical device so that it corresponds to your line of sight and the characteristics of your personal workstation (the lighting and the distance between the user and the device). The device must remain quickly and easily accessible at all times.

U Check that the water and air lines and the power cord do not hinder physical movement.



Power your device

You can now connect the power cord to the mains grid. <u>Protective earth is required!</u> Be sure your power grid has an efficient protective earth.

Voltage: 100-240 Vac Frequency: 50 to 60 Hz. Operating current: 4 A max.

Installation of the wireless pedal



Insert two (2) AA 1.5V lithium batteries into the wireless pedal. Close the cover and operate your device.

Risk of fire: use only batteries that have current limiter/short-circuit and over-temperature protection (compliant to IEC 60086-4:2014 Safety of lithium batteries).

The optional wireless pedal supplied with your device is already paired and ready to use (Note: A pedal can only command one single device at a time. Pairing is maintained even if the batteries are removed).

When you receive your new machine, all you need to do is insert the two (2) AA lithium batteries into the wireless pedal and your device is ready to work.

In case you replace your pedal, you will need to pair it with your device. For instructions, please read the specific Maintenance & Troubleshooting chapter.

The Bluetooth® radio is automatically disabled (powered off) when a wired pedal connected.

The wireless pedal uses a low power, 8 dBm EIRP max, Bluetooth[®] 2.4 GHz radio, to communicate with the control unit. Interference may occur in the vicinity of this equipment.

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 device, including cables. Otherwise, degradation of the performance of this equipment could result.



2.3. Powder Chambers

Clinical risk: Only use PLUS or PERIO Powder with the PLUS Powder chamber.

Clinical risk: Only use PLUS Powder chamber (red) for subgingival treatments.



The PLUS Powder chamber is designed for the PLUS Powder. It can be used for supra and subgingival treatments. Pressure is automatically reduced for compatibility with subgingival treatments,

including Perioflow treatments (Supra applications also possible). Compatible EMS Powders: PLUS and PERIO (refer to paragraph "Compatibility" for details).



The CLASSIC Powder chamber is designed for the CLASSIC Powder and can only be used for supragingival treatments.

Sodium Bicarbonate: Use only this powder and chamber for supragingival applications.

Compatible EMS Powders: CLASSIC and SOFT (refer to paragraph "Compatibility" for details).

Check bottle and powder chamber integrity: There should be no crack on the body.

 Δ The powder chamber is pressurized during use. Replace faulty parts immediately.

U Make sure that the powder chambers are dry.

Use only PLUS or PERIO Powders for restorations, crowns, bridgework, implants and orthodontics.

O not sterilize the powder chambers and their caps/parts by steaming or dry thermal reprocessing. Use only ambient temperature active disinfectant and cleaning agents.





By hand only: remove the powder chamber cap to refill powder up to the indicated MAX level, then insert the cap back fully onto the bottle.

Pour the powder in freely. <u>The central tube can be fully</u> <u>filled without problem</u>.

O not fill the chamber higher than the indicated MAX level. The powder level will go down slightly a few minutes after the filling (powder compaction).

Before pressurizing, position the powder chamber into the device. Magnetic attraction will position it correctly.

🛇 Do not insert upside-down.

2.4. Water supply and PIEZON[®] bottle

Without Bottle:

PIEZON[®] & AIRFLOW[®] use external water supply.





The CLIP+CLEAN shall be previously cleaned and disinfected before use. Non-disinfected CLIP+CLEAN may contaminate the device. Place the CLIP+CLEAN into the device's bottle receptacle for dust protection.

With Bottle connected:

AIRFLOW[®] uses external water supply. PIEZON[®] uses bottle liquid supply.



▲ Only use the PIEZON[®] bottle EG-111 (transparent) for disinfectant solutions.

The PIEZON® bottle is compatible with the following solutions:

Avoid unwanted chemical reactions or ingestion of residues of solution. Always rinse the liquid circuit with drinking water before using a different solution or undergo a regular treatment requiring water only.

• For treatment application use the percentage of solution as per discretion and per local country regulations and recommendations.

| Solution | Concentration | |
|------------------------------|---------------|------------|
| Sodium Hypochlorite | Up to 5% | EMS |
| Chlorhexidine | Up to 1% | |
| Carbanilides | Up to 0.05% | |
| Povidone iodine | Up to 12% | Piche |
| Quaternary ammonium compound | Up to 0.7% | SHULL MALL |
| Cetylpyridin chloride | Up to 1% | |
| Citric acid | Up to 5% | |
| Physiological water (NaCl %) | Up to 0.9% | |
| Ethanol | Up to 5% | 125-5 |
| Isopropanol | Up to 5% | a - m |

O not sterilize the PIEZON[®] bottle and its nozzle cap by steaming or dry thermal reprocessing. Use only ambient temperature active disinfectant and cleaning agents.



2.5. AIRFLOW[®] and PERIOFLOW[®] Handpieces

AIRFLOW[®] and PERIOFLOW[®] Handpieces are reusable, but they shall have been previously reprocessed: cleaned, disinfected and sterilized. Not reprocessed Handpieces and accessories may cause bacterial or viral infections.



Connect the AIRFLOW[®] or PERIOFLOW[®] Handpiece.

A Follow the "Reprocessing of EMS parts" instructions and the present-day regulations on reprocessing enforced in your country.

In case the AIRFLOW[®] Handpiece gets clogged, refer to the "Maintenance & Troubleshooting" section for instruction.

2.6. PIEZON[®] Handpiece and Instruments

PIEZON[®] Instruments and Handpieces are reusable, but they shall be reprocessed before use: cleaned, disinfected and sterilized. Not reprocessed handpieces and accessories may cause bacterial or viral infections.

Follow the "Reprocessing of EMS parts" instructions and the present-day regulations on reprocessing enforced in your country.

Check tip length and tip thread through the cover right folder of your Quick Guide.

If tip extremity is in the red area, it can have excessive and uncontrolled vibration. Replace the tip.



Accessories are available from EMS and authorized dealers.



Mount the tip / insert using the EMS CombiTorque tool

Once the instrument is screwed all the way in, give an extra quarter of a turn to obtain the required torque and remove the CombiTorque.

Use only the CombiTorque to tighten the EMS Instrument on the handpiece to the correct torque to avoid tip unscrewing or breaking.

Connect the PIEZON® Handpiece

Blow-dry the connections to remove any presence of liquid and to ensure a proper electrical contact.



Nose cap and light guide

Always use the handpiece with the light guide and the nose cap installed and correctly tightened.

For replacing the light guide, refer to the "PIEZON[®] light guide check & replace" section.



3.DEVICE USE

3.1. Interfaces



PIEZON® power setting



The unit is equipped with NO PAIN technology which provides an adaptive response in function according to the load applied to the instrument.

The following table shows the maximum output power as per user power setting:



A Risk of tip breakage: with ENDO files, do not exceed 2.5W (power setting "3" max.)

AIRFLOW[®] pressure setting



Both the PLUS and CLASSIC powder chambers have an integrated dynamic pressure regulator that automatically set the optimal pressure range for the selected powder chamber and related powder type as detailed in chapter "Powder Chambers".

The following table shows the static and approximate dynamic pressures⁶ as per selected powder chamber and user power setting:

| AIRFLOW® | Pressure Setting | 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
|-----------------|-----------------------|---|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|
| Pressure | Static [Bar] | | 2.5 | 2.7 | 3.0 | 3.2 | 3.5 | 3.7 | 4.0 | 4.2 | 4.5 | 4.7 |
| | CLASSIC dynamic [Bar] | | 1.9 | 2.1 | 2.3 | 2.6 | 2.8 | 3.0 | 3.2 | 3.5 | 3.7 | 3.9 |
| | PLUS dynamic [Bar] | | 1.5 | 1.7 | 1.9 | 2.0 | 2.2 | 2.4 | 2.6 | 2.7 | 2.9 | 3.1 |

⁶ Dynamic pressures depend on handpiece and powder type too. The listed pressures are for information purpose and referring to the commonly used EL-308 AIRFLOW[®] Handpiece with DV-082 and DV-048 powders.

PIEZON[®] and AIRFLOW[®] BOOST



Pressing hard on the center of the wireless pedal activates the BOOST mode and results in an increase of power, as the following table shows:

| AIRFLOW ® | Power Setting | 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
|------------------|---------------------------|---|---|---|---|---|----|----|----|----|----|----|
| Boost | Corresponding Boost Level | 0 | 6 | 7 | 8 | 8 | 8 | 9 | 10 | 10 | 10 | 10 |
| | | | | | | | | | | | | |
| PIEZON ® | Power Setting | 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| Boost | Corresponding Boost Level | 0 | 6 | 7 | 8 | 9 | 10 | 10 | 10 | 10 | 10 | 10 |

▲ Risk of tip breakage: use BOOST only with a tip suited for high power usage.
 ◇ DO NOT use BOOST with ENDO files.

Wireless pedal battery saving

Each time the wireless pedal is released, it enters into a low power mode. Even if unused for long, it is not required to remove the batteries.

To avoid an involuntary depletion of the wireless pedal batteries, in case the pedal remains pressed without interruption for 10 minutes, it will automatically enter into switch-off mode.

To resume from the switch-off mode, it is required to first release the wireless pedal and then power cycle the device (switch off for 30s and then power on again).

Water temperature and acoustic feedback settings

AIRFLOW® and PIEZON® liquids' temperature is 40°C by default.

To adjust the water temperature or the acoustic feedback, follow the procedure below:

- 1. Turn the device ON.
- 2. Securely place both handpieces (AIRFLOW® and PIEZON®) back into their holders.
- 3. Press (1) + (1) simultaneously to access the menu. (See image below place fingers in the groove below the numbers)



- 4. Colors will appear on the numbers:
 - 0 to 4 for setting water temperature (5 is not used)
 - 6 to 10 for setting acoustic feedback (5 is not used)

| Water temperature ⁷ | | | | | Acoustic feedback | | | | | |
|--------------------------------|-----|-----|-----|-----|-------------------|---------------|------------------|----------------|-------------------|--|
| 0 | 1 | 2 | 3 | 4 | 6 | 7 | | | 10 | |
| No Heating | 25° | 30° | 35° | 40° | No sound | Low volume | Medium volume | High volume | Maximum volume | |

- 5. Change the settings according to your wish.
- 6. Press the ON/OFF button to save the setting and exit.

Note:

- Changes are applied to both AIRFLOW[®] and PIEZON[®] liquid temperatures.
- After a few seconds of keyboard inactivity, the device automatically exits the mode.

⁷ The target temperature is determined into the device's body.

On AIRFLOW[®] side, water temperature decreases along the cord. Air spray also decreases the temperature. Final temperature of AIRFLOW[®] spray is lukewarm, lower than 40°C.

On PIEZON[®] side, PIEZON[®] Handpiece warms up the waterline which compensates natural cooling along the cord. Please adjust the temperature setting for maximizing patient comfort.



3.2. Treatment sequence

Ocnsult the Treatment Recommendations (FB-648) before starting any treatment to the patient.

AIRFLOW®

- 1 Position the powder chamber.
- 2 Pressurize the chamber.
- 3 Set the AIRFLOW[®] power.
- 4 Set the water flow.
- 5 Take the AIRFLOW[®] Handpiece.
- 6 Press the pedal to start treatment.
- 7 [Step hard on the center of the BT pedal for BOOST.]
- 8 Release the pedal to stop treatment.
- 9 Put the handpiece back into its holder.



PIEZON®

- 1 Set the PIEZON[®] power.
- 2 Set the water flow.
- 3 Connect the PIEZON[®] bottle (if required).
- 4 Take the PIEZON[®] Handpiece.
- 5 Press the pedal to start treatment.
- 6 [Step hard on the center of the BT pedal for BOOST.]
- 7 Release the pedal to stop treatment.
- 8 Put the handpiece back into its holder.



Treatment does not stop immediately. Beware there is a small delay between the release of the pedal and the effective stop of the treatment (approximately 0.2 second).

Risk of patient injury. If you are not trained on a specific treatment, do not execute it. Always get trained before executing new treatments.



4.OPTIONAL EQUIPMENT

4.1. PERIOFLOW® Nozzles



Single-use nozzle.



🗥 Cannot be reprocessed. DO NOT use the nozzle if the package is damaged or open.



Fully connect the nozzle by pushing on a hard surface.

Make sure the nozzle is correctly attached = fully inserted.



Remove the nozzle by using the nozzle extractor.

A Risk of injury: Always USE the nozzle extractor AB-358/A. DO NOT remove by hands.

4.2. Endochuck & PI Instrument

The Endochuck file holders are available in 90°, 120° or 180° orientations to adapt to particular configurations.

 \odot Do not tighten the chuck nut when no file or instrument is installed as this may damage it.



A Verify that the plastic coating is not worn or damaged before use.



4.3. Mirror Suction Cannula

Ultra FS ClasenUNO Mirror Suction Cannula is only available in the European Union.



Ultra FS ClasenUNO Mirror Suction Cannula is a combination of a dental mouth mirror and a medical suction cannula. The device is designed to improve the view of the area under treatment and/or for the suction of fluids and particles from the patient's mouth cavity.

The ClasenUNO Cannula must be reprocessed before use: cleaned, disinfected and sterilized.

Follow the ClasenUNO Reprocessing instructions and the present-day regulations on reprocessing enforced in your country.

Connect it to the high-speed suction hose of your dental unit and check for compatibility before use (It may not be compatible with your dental unit suction hoses).

U The mirror surface shall be dried thoroughly. Chalky coating on the mirror may be difficult, if not impossible, to remove.

The Ultra FS ClasenUNO Mirror Suction Cannula has been designed for a large number of sterilization cycles. Its service live is predominantly determined by wear and tear through use. Always replace the medical device as it presents any sign of worn-out or damage.



5. CLEANING & REPROCESSING

5.1. Water Line Cleaning & Disinfection

Keeping the device's water lines clean and disinfected is mandatory to prevent patient infection.

NIGHT CLEANER⁸ ensures the decontamination and prevents biofilm formation in water lines of AIRFLOW Prophylaxis Master.

NIGHT CLEANER⁹ removes and protects from algae and limescale, after longer idle times or heating of the process.

The water supply hose and related device connection will not be cleaned by this procedure.



Each morning before the first patient: Rinsing



Place a fully filled water bottle onto the device

To reduce the risk of ingestion of the cleaning agent by the patient, always use a fully filled 800ml water bottle.



Set water to 10 Turn the device ON

Set both water regulators to 10 to ensure optimal rinsing.



Hold both cords over a sink with CLIP+CLEAN

Contamination prevention: Do not make any contact between the sink and the cords.



Press the pedal once, release, and then wait 1 minute

The white and blue countdown indicates remaining time.

Cleaning can be paused and reset by pressing the pedal again.

Risk of ingestion of the cleaning agent. Check that no more blue residue of NIGHT CLEANER⁹ is flushing out of the cord. Otherwise, repeat the rinsing procedure.

Always empty out and wash the water bottle used for rinsing before any new use. EMS recommends a weekly use of a bottle cleaning agent (e.g. BC-San 100 from Alpro Medical GMBH).

Risk of ingestion of residue of cleaning agent. During rinsing, a small quantity of cleaning agent flows back into the water bottle.

⁸ If available in your country. Not for end point sterilization.



Between each patient

Overall cleaning and disinfection



Clean the external surface of the device with a cloth and alcohol Clean the unit only with an alcohol-based (ethanol, isopropanol), colorless disinfectant.

Never use scouring powder or an abrasive sponge. It will damage its surface.



A Reprocess handpieces and instruments See the specific following chapters.

Risk of contamination. Always disinfect the bottom and top areas of device air connections.

End of day: Overnight cleaning

Use only EMS NIGHT CLEANER⁹ as a cleaning agent.
Other products may damage or not clean the unit, and cause patient intoxication.



Place the NIGHT CLEANER bottle onto the device



Each cleaning consumes 30ml of NIGHT CLEANER¹⁰.

Before cleaning, check that the liquid level is above the black flange of the bottle's neck.



Set water to 10 Turn the device ON

Set both water regulators to 10 to ensure the flow of the cleaning agent.



Hold both cords over a sink with CLIP+CLEAN

Contamination prevention: Do not make any contact between the sink and the

cords. CLIP+CLEAN shall be reprocessed after each use.

Press the pedal once, release, and then wait 1 minute

The white and blue countdown indicates remaining time.

Cleaning can be paused and resumed by pressing the pedal again.

Once completed, leave the NIGHT CLEANER bottle on the device overnight at least 12h minimum.

⁹ If available in your country. Not for end point sterilization.

The NIGHT CLEANER¹⁰ agent can remain active in the device's water lines (weekend, holidays or over night), and requires at least 12 hours (3 months maximum) of contact time for an optimal efficacy.



Refill the blue NIGHT CLEANER bottle with NIGHT CLEANER¹¹ agent only NIGHT CLEANER¹¹ has the following properties:

- Bactericidal/ fungicidal
- Removes and prevents lime and algae formation
- Remains stable in the NIGHT CLEANER bottle
- Blue color increases user awareness of the cleaning procedure

Do not sterilize the NIGHT CLEANER bottle and its nozzle cap by steaming or dry thermal reprocessing. Use only ambient temperature active disinfectant and cleaning agents.
 Do not use Hydrogen Peroxide like EMS Ultra Clean. It deactivates after some time in the device's bottle.

5.2. Safety Information on NIGHT CLEANER¹¹

 \bigcirc DO NOT mix NIGHT CLEANER¹¹ with other cleaning solutions.

NIGHT CLEANER¹¹ should not be swallowed. Keep this product away from children. In case of ingestion, rinse the mouth with water. Do not induce vomiting. In case of discomfort, consult a medical doctor.

▲ NIGHT CLEANER¹¹ should not be inhaled.

In case of inhalation, supply fresh air and if necessary consult a physician.

Avoid contact with eyes. In case of contact, flush eyes with flowing water for several minutes holding eyelids apart. Remove contact lenses , if present and easy to do.

I Manipulate the product with gloves. In case of skin contact, wash it with water and soap.

In case of soiled clothes, take off these immediately. If you have any contamination concern, promptly consult a medical doctor.

• For further information refer to the specific NIGHT CLEANER¹¹ instructions for use provided with the product.

Manufacturer information and point of contact

For any information and/or complaints, you can also contact the legal manufacturer: **ALPRO MEDICAL GMBH** Mooswiesenstrasse 9 78112 St. Georgen, GERMANY Phone: +49 7725 9392-0 www.alpro-medical.com

¹⁰ If available in your country. Not for end point sterilization.

5.3. Reprocessing of EMS parts

EMS recommends the use of cleaning, disinfection, packaging for sterilization and sterilization procedures accordingly with ISO 17664.

I Always report adverse events related to device reprocessing directly to EMS.

Reusable products must be cleaned, disinfected and, if applicable, sterilized prior to first use. Do not reprocess the products over the allowed number of sterilization cycles, but replace: refer to the "Service life" section of the "Technical Description" chapter.

Concentrations and contact times specified by the manufacturer of the cleaning and disinfection agent must be followed.

Remember that sterilization cannot be achieved unless the elements of the assembly are cleaned and disinfected first.



If there is anything in the following instructions that is not clear or seem to be inadequate, do not hesitate to contact/inform EMS.

The following instructions have been validated as being capable of preparing for re-use the EMS medical devices and parts listed in the "Intended Use and Compatibility" chapter. It remains the responsibility of the user to ensure that the processing as actually performed using equipment, materials and personnel in the processing facility achieve the desired result. This requires 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.

The user shall also observe any applicable legal requirements in their country as well as the hygiene regulations of the hospital or clinic. This applies especially with regard to the additional requirements for the inactivation of prions.

Preparation

Manual pre-cleaning is required:

Immediately after use, rinse the lumen(s) line of the handpiece/instrument with water for 20 seconds. Coarse soiling must be removed immediately after application.

• For AIRFLOW[®] and PERIOFLOW[®]: always carry out handpiece powder unclogging and check for both lumens (water and powder) clearage before proceeding.

• For PIEZON®:

- Remove any installed instrument.
- Remove the nose cap and separate aside the light guide and the O-ring.



• For Endochuck file holder:

- Remove any installed file.
- Remove the screw and separate aside the small O-ring.



U For any Instrument mounted on the CombiTorque:

• Separate aside the Instrument and the CombiTorque.



Safely transport to the reprocessing area to avoid any damage to the parts and contamination to the environment and to the people involved in the reprocessing process.

Cleaning shall need to be performed within 1 hour from the use.

Cleaning

Any part can be cleaned manually or automatically by washer or disinfector. EMS recommends the use of an ISO 15883 compliant automatic washer-disinfector (WD) for an optimal effectiveness and part service life.

O DO NOT use any Ultrasound Bath cleaning procedure with the PIEZON[®], AIRFLOW[®] and PERIOFLOW[®] handpieces: it may destroy the products.

Manual Cleaning (without ultrasound bath)

The following validated process can be used with any EMS part:

- Remove away any surface soiling (gross contamination) on the product with a wetted cloth and tap water.
- Brush the devices in a cleaning solution of 0.5% neodisher MediClean (Dr. Weigert Hamburg) in deionized water at 40°C with a suitable soft bristled brush (Medisafe MED100.33) until all visible residues have been removed.
- For parts having lumens, flush all lumens with a spray gun (water jet gun, with a static water pressure of 2 bar) with cold tap water for 15 seconds.
- Immerse the part in 0.5% solution of neodisher MediClean with deionized water for 15 minutes at 40°C. Taken care that all lumens are filled with cleaning solution (use a syringe if needed). All surfaces must be moistened.
- Rinse all lumens flushing with a spray gun (water jet gun, with a static water pressure of 2 bar) with cold deionized water for 15 seconds, and further rinse the whole part under cold running deionized water for 10 seconds.
- Using an air pistol (compressed air) fully dry the lumen and the whole part until no more residues of water are present (visible or detectable).

Automated Cleaning

The washer-disinfector shall comply with ISO 15883 standard, must have suitable baskets to hold small fragile products and must have rinsing connections with diameter of approximately 16mm for the attachment to product lumen.

Correctly place the instrument into a suitable rack, connect all lumens to the rinsing connectors and start the automated cleaning. The following validated (For example on Miele Professional G 7836 CD having Miele Rack E429) automated cleaning process can be used:

- 2 minutes pre-washing with cold water.
- Drain.
- 5 minutes cleaning with tap water and 0.5% detergent of neodisher MediClean Dental (Dr. Weigert, Hamburg) at 55°C.
- Drain.
- 3 minutes rinsing and neutralization cold with deionized water.
- Drain.
- 2 min final rinse with cold deionized water.
- Drain.

Also instructions of the manufacturer of the washer disinfector shall be followed.



Manual Cleaning with Ultrasound Bath O NOT USE WITH HANDPIECE BODY

The following validated process is intended ONLY for the EMS PIEZON Instruments and file holders:

- Remove away any surface soiling (gross contamination) on the product with a wetted cloth and tap water.
- For part having lumens, flush all lumens with a spray gun (water jet gun, with a static water pressure of 2 bar) with cold tap water for 15 seconds.
- Using an appropriate sieve for sonication, immerse the part in 0.5% solution of neodisher MediClean with deionized water and sonicate for 10 minutes at 40°C (Bandelin, Sonorex 1028K, 35kHz). Taken care that all lumens are filled with cleaning solution (use a syringe if needed). All surfaces must be moistened.
- Rinse all lumens flushing with a spray gun (water jet gun, with a static water pressure of 2 bar) with cold deionized water for 15 seconds, and further rinse the whole part under cold running deionized water for 10 seconds.
- Using an air pistol (compressed air) fully dry the lumen and the whole part until no more residues of water are present (visible or detectable).



Example of correct placement of the parts in the WD Miele Professional G 7836 CD using the Mobile Injector Unit (Rack) Miele E429 Mettre nouvelle photo et remplacer le texte par Exemple de positionnement des appareils dans un thermodesinfecteur compatible ISO ---



Disinfection

Any part can be disinfected manually or automatically by a disinfector.

HIGHLY RECOMMENDED! Automated Disinfection

EMS highly recommends the use of an ISO 15883 compliant automatic washer-disinfector (WD) for an optimal effectiveness and part service life

All EMS parts can undergo an Automated Thermal Disinfection in a washer-disinfector compliant with the ISO 15883 standard.

The following validated (For example on Miele Professional G 7836 CD using the Rack Miele E429) process can be used to achieve an A₀ Level of 3000:

- Execute a 3 minutes thermal disinfection (final rinse) with deionized water at 93°C minimum.
- Drain.
- Drying for at least 20 minutes at 100°C.

Also special instructions and warnings of the manufacturer of the washer disinfector have to be followed.

In case of use of a chemical disinfectant, follow carefully the instructions provided by the disinfection solution manufacturer.



Example of correct placement of the parts in the WD Miele Professional G 7836 CD using the Mobile Injector Unit (Rack) Miele E429

ALTERNATIVE Manual Disinfection

DO NOT use any Ultra Sound Bath disinfecting procedure with the PIEZON®, AIRFLOW® and PERIOFLOW® Handpieces: it may destroy the products.

- ASP CIDEX OPA[®] 0.55% disinfectant solution shall be used not diluted and within its Use-Life and Shelf-Life, respecting the manufacturer's Warnings and precisely following its Instruction for Use.
- Disinfectant shall be poured into a proper tray having secure lid. Tray size shall have to allow the complete submersion of the device and to facilitate the evacuation of air from the inside.
- Warm up the disinfectant to at least 20°C.
- Immerse the product completely, and if applicable, fill all lumens and eliminate air pockets by means of a syringe fulfilled of disinfectant (to guaranty full internal lumen contact with the disinfectant).
- Close the tray with the secure lid.
- Wait for at least 5 minutes and be sure that the temperature of the disinfectant solution doesn't drop below 20°C.
- The ASP CIDEX OPA disinfectant requires a total of three (3) rinses, with large volumes of fresh water to properly remove its residues. Sterile water shall be poured into proper rinsing trays. Tray size shall have to allow the complete submersion of the device and to facilitate the evacuation of air from the inside.
 - For each of the 3 rinses do:
 - Flush all lumens with large volumes (not less than 100 ml) of rinse water. Use a syringe or a water jet pistol to correctly accomplish this task.
 - Keep the device totally immersed for a minimum of 1 minute in duration.
 - Remove the device from the rinsing tray and discard the rinse water.
- Finally, fully dry it internally and externally.

Always use fresh volumes of water for each rinse. Do not reuse the water for rinsing or any other purpose. Residues of disinfectant may cause serious side effects.

A Final rinse water requirements:

Free from optional pathogenic micro-organisms. Deionized water is recommended to avoid deposits or crystallization on the medical device. Potential bacterial contamination must be taken into account depending on the water treatment process used.

A Sterilization shall be performed immediately after cleaning-disinfection.

Inspection and final dry before sterilization

If stains are still visible on the part after cleaning/disinfection, the entire cleaning/ disinfection procedure must be repeated. Parts with visible damage, chip/flake loss, corrosion or are bent out of shape must be disposed of (no further use is permissible). Check also the integrity of O-rings and gaskets and replace if damaged or out of shape.

• Verify the part to be fully dry. In case of detection of residues of water, remove these using an air pistol (clean compressed air). Fully dry the lumen and the whole part until no more residues of water are present (visible or detectable).

Reassembly and packaging for sterilization

▲ Only previously cleaned and disinfected parts can be sterilized.

Effective sterilization can take place only on fully dry parts. Ensure each part (internal lumens and any surface) to be perfectly dry before reassembling and packing.

Prior to sterilization, the parts need to be reassembled to their readiness of use and placed in a suitable sterilization packaging.

• For PIEZON®:

• Reinstall the O-ring first, then put the light guide into the nose cap and screw it on the handpiece.





• For Endochuck file holder:

• Reinstall the small O-ring (gasket) first, and then gently screw the retention bolt without tightening it.



For any Instrument having its CombiTorque:

• Reinstall the Instrument on the CombiTorque.



Any EMS part can be correctly packaged using the following validated procedure:

• Single or Double Pouches suitable for Pre-Vacuum Moist Heat Sterilization compliant with ISO 11607-1 or EN 868, resistant to 138°C and having adequate steam permeability (e.g. Wipak STERIKING flat rolls Type R43 and R44).

Sterilization

A Sterilization must be performed immediately after cleaning-disinfection.

 \odot DO NOT exceed the maximum number of sterilization cycles allowed.

 \bigcirc DO NOT exceed a sterilization temperature of 138°C and a holding time of 20 minutes.

 \odot DO NOT use hot-air sterilization and radio-sterilization procedures: they destroy the products.

Moist heat sterilization of parts shall be performed according to ISO 17665 and under consideration of the respective country requirements.

The following validated Pre-vacuum Moist Heat (steam) process can be used with any EMS part packaged in appropriate single or double pouches:

Parameters for the Pre-vacuum Moist Heat cycle:

- 3 pre-vacuum phases
- Sterilization temperature of 132°C
- Pressure of 3 bar (absolute pressure)
- Humidity of 100%
- Holding time of 3 minutes minimum (full cycle)
- Drying time of 20 minutes minimum

▲ It is the duty of the user to ensure that the reprocessing processes, including resources, materials and personnel, are capable to reach the required results and maintained over the time: keeping actual the validation of the reprocessing processes is under the responsibility of the user.

Storage

Store the sterilized instruments in a dry, clean and dust free environment at a temperature of 5°C to 40°C.

Service life

If the number of permissible re-sterilization cycles is restricted, this will be stated in the product's specific Instructions for Use (if any) and/or in the "Service life" section of the "Technical Description" chapter.

The products have been designed for a large number of sterilization cycles. The materials used in their manufacture were selected accordingly. However, with every renewed preparation for use, thermal and chemical stresses will result in the ageing of the products.

Always replace products that present sign of worn-out or of early degradation, regardless of the number of sterilization cycles left unused.

 \odot DO NOT expose the products to temperature exceeding the 138°C.

5.4. Reprocessing of ClasenUNO Cannula

The following instructions are from the Cleverdent documentation "ClasenUNO Instructions" edition 03/2016 and are current at the date of issue. We recommend that you regularly consult Cleverdent website or contact them for the latest version of their instructions for use and reprocessing.

The Ultra FS ClasenUNO Mirror Suction Cannula requires an EN ISO 17664 compliant reprocessing. Check the cannula regularly before use and replace it as signs of wear are detected.

Cleaning and disinfection

Only disinfectants that are suitable for polypropylene (PP) and used according to the guidelines may be employed for cleaning and disinfection. To avoid the risk of scratching the mirror and cannula, do not use hard brushes (wire brushes) for cleaning. The requirements stated in EN ISO 17664 must be observed. Firstly, remove the coarse dirt, and then rinse the ClasenUNO under running water. For the ClasenUNO with Ultra mirror, use distilled water for rinsing. Place the contaminated ClasenUNO in a suitable disinfectant solution. Follow the recommendations of the solution manufacturer regarding the concentration levels of the disinfectant and the duration of disinfection. Rinse it well with water after disinfection and dry carefully. Next, disinfect/sterilize using one of the methods below.

Ultrasonic disinfection

Make sure that the surface of the mirror is completely dry, particularly with the Ultra version, as lime residue could be burnt in otherwise. Place the ClasenUNO in an ultrasonic bath (e.g. Bandelin Sonorex Super RK 514). Add a cleaning and disinfection agent suitable for polypropylene (PP) (e.g. 0.55% Cidex OPA) and set the washing cycle as per the manufacturer's instructions. Keep it in the ultrasonic bath for 12 minutes and ensure that the temperature does not drop below 18°C. Then, rinse with sterile water until all cleaning agent residue has been removed. Check to ensure that the ClasenUNO is thoroughly clean and, if necessary, repeat the cleaning cycle. Finally, dry the ClasenUNO carefully.

Disinfection with a thermal disinfector

To clean and disinfect with a thermal disinfector, use a device that corresponds to EN ISO 15883 (e.g. Belimed WD 100) and observe the manufacturer's instructions when choosing the cleaning and disinfection cycle and cleaning agent. Position the ClasenUNO in the thermal disinfector in such a way that the inside surfaces are rinsed and the water can flow off. 0.5% (V/V) deconex 24 LIQ has proven suitable as the cleaning solution and 0.2% (V/V) deconex 26 Plus as the neutralizing solution. Disinfection is performed at 90°C for a hold time of 5 minutes. At the end of the cleaning and disinfection cycle, ensure that the ClasenUNO is thoroughly clean and, if necessary, repeat the cleaning cycle.

Sterilization

Steam sterilization must be performed using a device that complies with EN 13060 or EN 285 (e.g. autoclave with fractionated pre-vacuum, W&H, type LISA 517), observing the sterilization procedure as per EN ISO 17665-1. The cycle must be conducted at a sterilization temperature of 134°C with a hold time of 5 minutes, or at a sterilization temperature of 121°C with a hold time of 12 minutes.

Manufacturer information and point of contact

For any information and/or complaints, you can also contact the legal manufacturer: **Cleverdent Ltd.**, Theresiengrund 31, DE - 48149 Münster, Germany. Tel: +49 (0) 251 98292828 Website: www.clasen.uno - Email: info@clasen.uno



6. MAINTENANCE & TROUBLESHOOTING

6.1. AIRFLOW[®] Handpiece powder unclogging

In case of a clogged handpiece and before the reprocessing of AIRFLOW[®] and PERIOFLOW[®] Handpieces.



Provided in your **AIRFLOW**® Application box



The Easy Clean tool can be thermally disinfected and also sterilized at up to 135°C in the autoclave.

6.2. AIRFLOW[®] handpiece leakage

disposable syringe filled with more than 2 ml of drinking water

In case of leakage at the AIRFLOW[®] handpiece connection with the AIRFLOW[®] cord, replace the o-rings of the cord with the spare provided in the EL-651 Kit located in the **AIRFLOW**[®] application box.



6.3. PIEZON[®] light guide check & replace

The light guide loses its transparence after undergoing repeated reprocessing cycles. Check the transparency of the light guide every month and do the following:



- 1. Remove the tip and unscrew the handpiece nose cap by hand.
- 2. Take off the light guide and inspect it.
- 3. Place in a new light guide AB-340 (provided in the PIEZON® application).
- 4. Screw the nose cap on again, by hand only.



6.4. Handpiece cord replacement

Bisconnect the mains plug for purposes of maintenance and in case of malfunction.

A Depressurize the powder chamber before disconnecting the AIRFLOW[®] cord.

In case of persistent malfunction or damage to the PIEZON[®] or AIRFLOW[®] Handpiece cord system, the part can be easily replaced by the user. Follow the directions for replacement provided with the spare part supply.



Handpiece cord disconnecting procedure:

- 1. Unlock the cord system by pushing the lock switch to the front (Switch located under the device).
- 2. The cord system is now unlocked and can be removed by pulling it.

6.5. Monthly check

Each month check both air and water filters for cleanliness.

 Disconnect the mains plug for purposes of maintenance and in case of malfunction. No maintenance is allowed while in use with a patient.



Check water and air filter cleanliness.

Good

Worn-out

Filter color has to be white without significant visible impurities. If not, replace the filter.

If the water filter needs to be changed more than 3 times a year, please check the quality of your water line.

Air filters usually remain cleaner for longer periods of time. Replace once a year. (The yearly maintenance service includes the replacement of both filters.)

- 1. E Disconnect the power cord from the grid first.
- 2. Disconnect the water hose by pulling it off the connector.
- 3. Pull the filter off by hand or by using a small flat screwdriver.
- 4. Replace with a new filter and reconnect the hose.

6.6. Yearly maintenance & repair



This device must only be maintained and/or repaired by EMS and by authorized EMS repair centers.



A yearly preventive maintenance or 2000 hours usage maintenance (LED ① solid orange), whichever comes first, is required as means of safety and performance guarantee for both the patient and the user. Qualified service repair may also be required anytime persistent malfunctioning is detected by the user and/or reported by the device diagnostic.



When returning the device for service, it is recommended that you ship the device with its pedal, powder chamber, bottle and cords in its original packaging for optimal protection against damage during transportation. Provide the contact details of your EMS dealer for a quicker service process (see § 6.9).

6.7. Pairing a new pedal



- 1. Remove one battery from the pedal (no need to remove both).
- 2. Place the two handpieces in their holders.
- 3. Turn the machine OFF, wait 10 seconds, then turn it ON again.
- Press (1) + (5) first, then also press (10) simultaneously. A sonar sound will start playing (if not, repeat step 4). Respect the order and the three-finger sequence (see figure below – place fingers in the groove below the numbers).
- 5. While the sonar sound plays, replace the lithium batteries into the wireless pedal.
- 6. Within a short time (less than 15 seconds), the pairing will be complete, the white LEDs will blink for a while and the device is then ready for use.



If the process takes longer than 1 minute, it means the pairing has failed and the device will automatically exit the mode. (No more sonar sound and no blinking at exit).

In case of this process failure, redo it from the beginning.



6.8. Troubleshooting



The device is whistling or making strange noises



First disconnect the mains plug.

This symptom is generally caused by a problem to the pressure regulator (fault or low temperature) or by a crack in the water bottle.

1° Stop using your device immediately and disconnect it from the grid.

2° Check the bottle in use for crack or any damage and, if the case, replace it with a new one.

3° Check the supplied air pressure: it shall be 4.5 bar minimum.

 4° lf the device temperature is below 10°C (device too cold), wait for it to warm-up at ambient

temperature and then reconnect to the power grid and switch it on again.

5° If the device temperature is over 10°C, or the problem recurs, definitively stop using it and contact EMS aftersales service.

The device is making smoke (and fire)







First disconnect the mains plug.

Stop using your device immediately, disconnect it and contact EMS aftersales service.

Cord or device leakage

A Risk of fire and electric shock.

First disconnect the mains plug.

1° If the leak is from the AIRFLOW® handpiece, replace the o-rings.

2° If the leak is from the device (handpiece support and water regulator), replace the complete handpiece cord.

3° If still not solved, contact EMS aftersales service.

LED 1 is SOLID orange

Automatic maintenance reminder. It is time to send your device to yearly maintenance service. Promptly contact EMS aftersales service.

LED 1 BLINKING orange

A Permanent or transitory hardware fault condition detected.

1° Unplug the device power cord, wait for 30 seconds, then plug it back again and restart the device (to check for effective permanent fault condition).

The wireless pedal's 2x AA lithium batteries are depleted. Replace both with new AA high-quality lithium

2° If the error is still present, contact EMS aftersales service for repair.



LED 3 SOLID orange

batteries having current limiter protection.

LED 2 SOLID orange

The problem may have multiple causes. A step-by-step multiple checks are required.

1° No pedal detected (at least one pedal must be connected to operate the device):

- Wired pedal may be disconnected. Check if the jack is fully inserted. Restart the device.
- Wireless pedal is not paired. Execute the procedure "Pairing of new pedal"

2° If the error is still present, contact EMS aftersales service for repair.

LED 3 BLINKING orange

Both the AIRFLOW[®] and PIEZON[®] cord systems are not detected or missing. At least one cord system is required to operate the device.

1° First, switch OFF the device, then disconnect both AIRFLOW® and PIEZON® handpiece cords and clean the electric contacts (jacks) present on the cord system connections. Also blow air to clean the device connection receptacles.

2° Reinstall both handpiece cords and start the device again.

3° If error is still present, contact EMS aftersales service.





LED 4 BLINKING orange





1° Your device is too hot. Unplug it, wait for 1 hour and start the device again.

2° If error is still present, contact EMS aftersales service.

Note: This error also shows up when the device is operating below the minimum temperature. In this case, just wait for the device to warm up to ambient temperature.



Water filter leakage

First disconnect the mains plug.

1° Replace the water filter (blue cartridge).

2° If still not solved, contact EMS aftersales service.

Bottle or bottle connection leakage

- 1° Ensure the bottle cap has been correctly closed.
- 2° Clean the connection: cap and device sides.
- 3° Replace the bottle.
- 4° If still not solved, contact EMS aftersales service.

AIRFLOW® connection leakage

- 1° Ensure the handpiece has been correctly connected to the cord.
- 2° Clean the interior of the handpiece and the cord terminating end.
- 3° Replace the AIRFLOW® cord gasket as described in paragraph "AIRFLOW® handpiece leakage". 4° If still not solved, contact EMS aftersales service.

Insufficient or no water from handpiece

1° Make sure you have set your water regulators to 10 (maximum flow on the cord) and verify that the handpiece is not clogged by removing it and checking the water flow without handpiece. 2° Check your water filter cleanliness and replace it if necessary.

Bisconnect the mains plug before servicing any filter.

2° Make sure you have well-connected and sufficient pressure from your water supply. 3° If still not solved, contact EMS aftersales service.

Still some blue liquid remaining after rinsing

- 1° Make sure you have set your water regulators to 10: maximum flow on the cords.
- 2° Make sure you have well-connected and sufficient pressure from your water supply.
- 3° Perform a second rinsing phase before treatment.
- 4° If still not solved, contact EMS aftersales service.

The unit does not start

1° Check the electrical connection and power socket. 2° Check the fuses at the back of the unit:

First disconnect the mains plug.

- Fuses are housed within the power cord socket.
- 1° Remove the power cord from the device.
- 2° With the help of a small flat screwdriver, open the fuse-holder cover.
- 3° Replace fuses only with the exact type required (refer to the "Technical Description" section).
- 4° If still not solved, contact EMS aftersales service.



Wireless pedal does not work

In the case is evident that the pedal remained pressed for longer than 10min, simply release the pedal and power cycle the device. If not this case, the problem may have multiple causes. A step-by-step multiple checks are required:

1° Switch-off the device and disconnect and reconnect both the PIEZON® and AIRFLOW® cord systems. Try again.

2° Perform a new pairing. This procedure is explained in the paragraph "Pairing a new pedal". Try again. 3° Change the 2x AA lithium batteries and try again.

4° If still not solved, contact EMS aftersales service.





Night Cleaner









Wired pedal does not work

1° Disconnect and reconnect the pedal. Check the cable for damage. Restart the device. 2° If still not solved, contact EMS aftersales service.

No pressurization of the powder chamber

- 1° Check that your device is ON: at least 1 LED light should be ON.
- 2° Check that the AIRFLOW[®] cord system is well connected (full green mark on the lock actuator).
- 3° lf still not solved, contact EMS aftersales service.

Powder chamber white light is BLINKING at pressurization attempt

- Either the air line is not connected or there is not enough air pressure.
- 1° Check the air line for no kinking and check the air compressor unit.
- 2° Check air filter for cleanliness and replace if dirty.
- 3° If still not solved, contact EMS aftersales service.

Powder chamber white light is BLINKING at depressurization

- 1° The handpiece could be clogged. Unclog with Easy Clean (see paragraph below).
- 2° AIRFLOW® cord could be clogged. Dismount and clean the airflow cord extremity.
- 3° If still not solved, contact EMS aftersales service.

Powder sprays out of chamber at depressurization

- 1° Powder chamber is filled beyond the maximum level marked.
- 2° Remove the powder exceeding the MAX sign on the bottle.



Powder leaks under the AIRFLOW® Handpiece cord system

The AIRFLOW® pinching element might be worn out or the air interface is dirty and is leaking powder. 1° Disconnect the cord, clean the air jack and connect again. If problem persists, go to Step 2. 2° Replace your AIRFLOW® Handpiece cord with a new one.

3° If still not solved, contact EMS aftersales service.

Powder chamber is leaking

 1° Clean the chamber with a wet cloth, in particular the cap and the bottom o-rings. Also clean the connecting elements on the device.

2° If still not solved, replace the powder chamber with a new one.

White LED PIEZON® is not working

- 1° Clean and dry the handpiece connection and try again.
- 2° Your PIEZON® Handpiece LED might have been switched off by activity time-outs:
- after 10 minutes of continuous operation,
- or after 20 seconds of inactivity off the holder.
- In both cases, put back the handpiece into the holder, wait 1 minute and try again.
- 3° If still not solved, contact EMS aftersales service.

Insufficient lighting

1° Check the light guide and replace if necessary.

2° If the light is still weak, replace the handpiece.

Damaged light guide

Replace the light guide.

Low or no mechanical power delivered by PIEZON® or vibration perceived

- 1° Make sure that the PIEZON® Instrument (tip) is correctly screwed on (use CombiTorque tool).
- 2° Check the wear of the instrument (tip) and replace it if necessary.
- 3° Clean and dry the handpiece and the cord system electric connections.
- 4° Replace the PIEZON[®] Handpiece first.
- 5° Replace the PIEZON® Handpiece cord.
- 6° If still not solved, contact EMS aftersales service.



6.9. To contact EMS Service support

E.M.S. Electro Medical Systems S.A. Ch. de la Vuarpillière 31 1260 Nyon – Switzerland Phone: +41 (0) 22 99 44 700

Fax: +41 (0) 22 99 44 700 Fax: +41 (0) 22 99 44 701 Email: <u>TSAV@ems-ch.com</u>

6.10. To report an Adverse Event

If any serious incident occurs that is directly or indirectly related to the treatment, report it immediately to EMS and to the competent authority of your country and of where the patient is established (if different).

Adverse Event notification to EMS

By email: <u>vigilancemailbox@ems-ch.com</u> By fax: +41 (0) 22 99 44 701 By post: E.M.S. Electro Medical Systems S.A., Ch. de la Vuarpillière 31, 1260 Nyon – Switzerland



7.SUSTAINABILITY

7.1. Disposal of waste parts



The device must not be discarded in domestic household waste. Should you wish to definitively dispose of the device, please comply with the regulations that apply in your country.

Other parts of this device, including tips/inserts, and chemicals must be disposed of according to your country's regulations.

Waste Electrical and Electronic Equipment belonging to customers located in the European Union may be shipped to EMS for recycling in accordance to the WEEE regulations. The costs of recycling, exclusive of shipping fees, are covered by EMS.



Keep the original packaging until the device is to be disposed of permanently. It can be used for shipping or storing.

7.2. Sustainable design



The device, on a voluntary basis, respects the latest Eco design low energy standby and off mode consumption regulation¹¹. Packaging cardboards are recycled and recyclable.



Concentration
 Concent

Printed instructions are aligned with a sustainable development policy and are certified « Myclimate neutral imprimerie » and « FSC ».

8.WARRANTY

Warranty is void if the device has been used with non-original EMS powder, instruments and handpieces. Warranty is void if the device has been opened.

EMS and the distributor of this device accept no liability for direct or consequential injury or damage resulting from improper use, arising in particular through non-observance of the instructions for use, or improper preparation and maintenance.

EMS declines the responsibility for the safety of the device and declares the warranty null and void if service or repair is carried out by an unauthorized third party or if non-genuine spare parts are used.

¹¹ European Commission Regulation N°1275/2008 of 17 December 2008 regarding the Eco design requirements for standby and off mode electric power consumption of electronic household and office equipment.



9. TECHNICAL DESCRIPTION

| Manufacturer | EMS ELECTRO MEDICAL SYSTEMS SA, CH-1260 Nyon, Switzerland |
|---|---|
| Models | AIRFLOW Prophylaxis Master, product code FT-229 |
| Classification IEC 60601-1 | Electrical Insulation Class-I Applied part Type B IP20 Control unit IP21 Foot pedal |
| Classification EU MDD 93/42/EEC | Medical Device Class Ila |
| Essential Performance | This medical device, in the meaning of the EU MDD 93/42 has no Essential Performance |
| Operating mode | Continuous operation |
| Power supply | 100-240Vac, 50-60Hz, 4A max. |
| Power consumption | OFF-mode / Stand-by: 0.5W max. Max: 700VA |
| Ultrasonic module | Max Output Power: 8W under fully-loaded mechanical condition. Frequency: 24-32kHz. Primary tip vibration excursion: 200um max. |
| Fuse | 5A, T (slow), 250Vac, H type (=T5H250V) |
| Wireless communication module | Max 8 dBm EIRP, 2.4GHz band, Bluetooth [®] radio module |
| Weight | Control Unit 5kg max. (full operating condition) Foot pedal: 0.35kg max. (wireless pedal) |
| Dimensions | Control Unit: Height: 245 mm, Width: 260 mm, Length: 290 mm Wireless pedal: Diameter 135 mm, Height 35 mm |
| Operating conditions | Temperature: 10°C to 35°C Humidity: 30% to 75% Altitude: Max 2000m |
| Storage conditions (device) | Temperature: -10°C to 30°C, no water inside Humidity: 10% to 95% not condensed Pressure: 500hPa to 1060hPa |
| Storage conditions (application box) | Temperature: up to 40°C |
| Transport conditions | Temperature: -29°C to 38°C, no water inside Humidity: 10% to 95% not condensed Pressure: 500hPa to 1060hPa |
| Input fluids | Water: pressure 2-5bar, temperature 10-30°C, salinity 0.2% max., hardness from 8 to 12°dH, minimum flow-rate 100ml/min, RECTUS 20KA connector type. EN-1717 compliant water network/inlet is required. Air: pressure 4.5-7bar, dry-only (humidity 1.032g/m3 max.), oil filtered 0.1mg/m3 max., minimum flow-rate 20 NI/min at 4.5bar, RECTUS 21KA connector type |
| Output fluids | Water: min. 40ml/min. for AIRFLOW [®] , min. 30ml/min. for PIEZON [®] Air: max pressure 5bar for Airflow |
| Shelf life / lifetime | PIEZON [®] and NIGHT CLEANER [®] bottles: 5 years Handpieces (main bodies): 1000 sterilization cycles CombiTorque: 1000 sterilization cycles |
| Expected service life | Device: 7 years, having regular yearly preventive maintenance |



9.1. Symbols

| $\mathbf{\Lambda}$ | General Warning |
|---|--|
| 4 | Warning Electricity |
| | Non-ionizing radiation (radio communication) |
| | Read the operation instructions |
| Ă | Device requiring protective earth |
| Ê | Disconnect the mains plug for purposes of maintenance and in case of malfunction |
| | Electronic instructions for use |
| | Mandatory action |
| | Expiration date |
| | Single use. Do not re-use. |
| Ŏ | Do not do. |
| | Disposal of old electronic equipment (European Union & other countries with separate collection systems) |
| 135°C | Sterilizable at up to 135°C in the autoclave |
| 本 | Thermal disinfection |
| \rightarrow | Input |
| \bigcirc | Output |
| | Fuse |
| 2 | Wired foot pedal connection |
| IP | Protection against water permeability |
| | Applied part, type B |
| | Manufacturer |
| ₩ | Manufacturing date |
| SN | Serial number |
| REF | Catalog number / Product reference |
| 0124 | Nedical Device compliant with EU Directive 93/42/EEC Number of the Notified Body |
| PG | GOST R for products in conformance with Russian standards |
| ULTER23 | Ukrainian Technical Regulation compliance marking for wireless equipment UA – Symbol of Ukraine; TR – Provisional symbol of the Conformity Assessment Body that is assigned to perform conformity assessment to the requirements of technical regulations; 028 – Identification number of the designated Conformity Assessment Body. |
| AGREE PAR L'ANRT MAROC Numéro d'agrément: MR 17713 ANRT 2018 / MR 14883 ANRT 2017 | Moroccan ANRT compliance marking for wireless equipment MR 17713 ANRT 2018: Wireless pedal approval number MR 14883 ANRT 2017: Device approval number |
| Date d'agrément: 16-10- 2018 / 09-10-2017 | |
| TRA REGISTERED NO: ER64514/18 ER67538/18 DEALER NO: DA76058/18 | United Arab Emirates TRA compliance marking for wireless equipment ER64514/18: BLE113 Bluetooth module approval number ER67538/18: BLE121LR Bluetooth module approval number |
| \bigotimes | Australian RCM compliance marking for wireless equipment |



R-NZ Complies with IMDA Standards (DB106919) CMIIT ID: 2018DJ3393

R-RMH-E23-FT-229 KCC-CRM-BGF-BLE13 CCCCRM-BGF-BLE13 TA-2017/2826 TA-2018/3027 APPROVED New Zealand R-NZ compliance marking for wireless equipment Singaporean IMDA compliance marking for wireless equipment DB106919: Dealer's Licence No. Chinese SRRC compliance marking for wireless equipment

2018DJ3393: System approval number Korean KC compliance marking for wireless equipment

R-RMM-E23-FT-229: System approval number KCC-CRM-BGT-BLE113: Bluetooth module approval number

South African ICASA compliance marking for wireless equipment TA-2017/2826: BLE113 Bluetooth module approval number TA-2018/3027: BLE121LR Bluetooth module approval number



9.2. Electromagnetic Compatibility

The use of parts other than those supplied or listed as accessory may negatively affect EMC performance. The device has embedded a low power, 8 dBm EIRP max, Bluetooth 2.4 GHz module, for communication with the

wireless pedal. This radio module is disabled when a wired pedal is connected (device reboot required). The Bluetooth module complies with all the restrictions foreseen by the ERC recommendations 70-03 for the CEPT

countries concerning the Annex 3 (Wideband Data Transmission System band A 2400-2483.5 MHz) without requiring any modifications of the products by the user.

The product is intended for use and Basic Safety is maintained in the electromagnetic environment specified below. The customer or the user of this product should assure that it is used in such an environment.

Electromagnetic immunity compliance

| Immunity test | IEC 60601 test level | Compliance level | Electromagnetic environment - guidance |
|---|--|---|---|
| Electrostatic discharge (ESD) IEC 61000-4-2 | ± 8 kV c ± 15 k | ontact V air | Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, relative humidity should be > 30%. |
| Electrical fast transient/burst IEC 61000-4-4 | ± 2 kV for power supply I freque ± 1 kV for input/output li freque | ines 100 kHz repetition ency nes 100 kHz repetition ency | Mains power quality should be that of a typical commercial or hospital environment. |
| Surge IEC 61000-4-5 | ± 1 kV line(s ± 2 kV line(s | s) to line(s) s) to earth | Mains power quality should be that of a typical commercial or hospital environment. |
| Voltage dips IEC 61000-4-11 | <5 % UT (>95 % dip 40 % UT (60 % dip 70 % UT (30 % dip in 0 % UT for 0,5 cycle at 0 225°, 270° 0 % UT for 1 cyc | in UT) for 0,5 cycle in UT) for 5 cycles n UT) for 25 cycles °, 45°, 90°, 135°, 180°, and 315° le single phase | Mains power quality should be that of a typical commercial or hospital environment. If the user of the product requires continued operation during power mains interruptions, it is recommended that the product be powered from an uninterruptible power supply or a battery. |
| Voltage interruptions IEC 61000-4-11 | <5 % UT (>95 % o 0% UT for 2 | dip in UT) for 5 s 250 cycles | |
| Power frequency (50/60 Hz) magnetic field IEC 61000-4-8 | 30 A/m | 30 A/m (50 Hz or 60 Hz) | Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment. |
| Conducted RF IEC 61000-4-6 | 3 V 150 kHz to 80 MHz 6V in ISM bands 150kHz and 80 MHz 80 % AM at 1 kHz | 3 V | Portable and mobile RF communications equipment should be used no closer to any part of the product, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter. Recommended separation distance $d = 1.17 \sqrt{P}$, $d = 1.17 \sqrt{P}$ 80 MHz to 800 MHz, $d = 2.4 \sqrt{P}$ 800 MHz to 2.7 GHz |
| Radiated RF IEC 61000-4-3 | 3 V/m 80 MHz to 2.7 GHz 80% AM at 1 kHz | 3 V/m | where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and d is the recommended separation distance in meters (m). Field strengths from fixed RF transmitters, as determined by an electromagnetic site survev ¹² , should be less than the compliance |
| Proximity fields from RF wireless communications equipment IEC 61000-4-3 | See Table below | | level in each frequency range ¹³ . Interference may occur in the vicinity of equipment marked with the following symbol: ((())) or () |

Notes:

UT is the a. c. mains voltage prior to application of the test level.

At 80 MHz and 800 MHz, the higher frequency range applies.

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

¹² 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 product is used exceeds the applicable RF compliance level above, the product should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as re-orienting or relocating the product.

¹³ Over the frequency range 150 kHz to 80 MHz, field strengths should be less than 3 V/m.



Proximity fields from RF wireless communications equipment

| Test Frequency | Band | Service | Modulation | Maximum power | Distance | Immunity test level |
|----------------------|-------------|--|--------------------------------------|---------------|----------|------------------------|
| (MHz) | (MHz) | | | (VV) | (m) | (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 |
| 710 745 780 | 704-787 | LTE Band 13, 17 | Pulse Modulation 217 Hz | 0,2 | 0,3 | 9 |
| 810 | | GSM 800/900, | Pulse | | | |
| 870 | 800-960 | iDEN 820, CDMA | Modulation | 2 | 0,3 | 28 |
| 930 | | 850, LTE Band 5 | TOTIZ | | | |
| 1845 | 1700-1990 | GSM 1800, CDMA 1900, GSM 1900, DECT, | Pulse Modulation | 2 | 0,3 | 28 |
| 1970 | | 25, UMTS | 217 HZ | | | |
| 2450 | 2400 -2570 | Bluetooth, WLAN, 802.11b/g/n, RFID 2450, LTE Band 7 | Pulse Modulation 217 Hz | 2 | 0,3 | 28 |
| 5240 5500 5785 | 5100 – 5800 | WLAN 802.11a/n | Pulse Modulation 217 Hz | 0,2 | 0,3 | 9 |

Recommended separation distances

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

| Rated maximum | Separation distance according to frequency of transmitter (m) | | | | | |
|-------------------------------|--|---|--|--|--|--|
| output power of transmitter W | 150 kHz to 80 MHz d = $\frac{3.5}{v} \sqrt{P}$ with V=3V | 80 MHz to 800 MHz d = $\frac{3.5}{E} \sqrt{P}$ with E=3V/m | 800 MHz to 2,5 GHz d = $\frac{7}{E} \sqrt{P}$ with E=3V/m | | | |
| 0.01 | 0.12 | 0.12 | 0.24 | | | |
| 0.1 | 0.37 | 0.37 | 0.74 | | | |
| 1 | 1.17 | 1.17 | 2.34 | | | |
| 10 | 3.69 | 3.69 | 7.4 | | | |
| 100 | 11.67 | 11.67 | 23.4 | | | |

For transmitters rated at a maximum output power not listed above, the recommended separation distance d in metres (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: This equipment has been tested and found to comply with the limits for a Class B digital device, pursuant to part 15 of the FCC Rules. These limits are designed to provide reasonable protection against harmful interference in a residential installation. This equipment generates, uses and can radiate radio frequency energy and, if not installed and used in accordance with the instruction, may cause harmful interference to radio communications. However, there is no guarantee that interference will not occur in a particular installation. If this equipment does cause harmful interference to radio or television reception which can be determined by turning the equipment off and on, the user is encouraged to try to correct interference by one or more of the following measures:

- Reorient or relocate the receiving antenna.

- Increase the separation between the equipment and receiver.

- Connect the equipment into an outlet on a circuit different from that to which the receiver is connected.

- Consult the dealer or an experienced radio/TV technician for help.

Electromagnetic emissions compliance

| Emissions test | Compliance | Electromagnetic environment - guidance | | |
|---|------------|--|--|--|
| RF emissions CISPR 11 | Group 1 | The product 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 product is suitable for use in all establishments, including domestic establishments and those directly connected to the public low-voltage power supply estuard that supplies buildings used for domestic purposes. | | |
| Harmonic emissions IEC 61000-3-2 | Class A | | | |
| Voltage fluctuations / flicker emissions IEC 61000-3-3 | Complies | network that supplies buildings used for domestic purposes. | | |



9.3. Radio Equipment Compliancy

This Medical Device and all of its accessories having radio equipment are compliant with the European Directive 2014/53/EU (RED – Radio Equipment Directive).



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