

INSTRUCTIONS FOR USE



This IFU is only valid for the United States.

Rx ONLY

EMS 



SUMMARY

BEFORE USE	5
INTENDED USE	5
INTENDED USERS	5
PATIENT POPULATION	6
INTENDED ENVIRONMENT	6
CONTRAINDICATIONS.....	6
RECOMMENDATIONS AND SAFETY PRECAUTIONS	6
COMPATIBILITY	7
CYBERSECURITY	7
1- GENERAL INSTRUCTIONS AND RECOMMENDATIONS	7
2- NETWORK INTERFACES AND CONNECTIVITY	7
3- SOFTWARE MANAGEMENT AND UPDATES.....	8
4- RESPONSE TO CYBERSECURITY INCIDENTS	8
5- SUPPORT AND LIFECYCLE	8
6- DECOMMISSIONING.....	8
INSTALLATION	9
1- EQUIPMENT INCLUDED IN THE BOX.....	9
2- STEP-BY-STEP INSTALLATION	10
2.1- Find an appropriate area to place the device	10
2.2- Check for proper water and air supply lines.....	10
2.3- Check for a proper and safe power grid	10
2.4- Be aware	10
2.5- Device connectivity	10
2.5.1- Connecting / Disconnecting from the Wi-Fi.....	11
2.6- Connect air and water hoses.....	11
2.7- Install accessories.....	12
2.8- Check the handpiece cord system connections.....	12
2.9- Fix the device	12
2.10- Power your device	13
2.11- Installation of the wireless pedal	13
3- POWDER CHAMBERS	13
4- WATER SUPPLY AND WATER BOTTLE	14
5- AIRFLOW® MAX AND PERIOFLOW® MAX HANDPIECES	14
5.1- Before use	14
5.2- Attaching and removing AIRFLOW® MAX or PERIOFLOW® MAX handpiece	15
5.3- Attaching and removing PERIOFLOW® nozzles.....	15
6- PIEZON® HANDPIECE AND INSTRUMENTS	15
6.1- Before use	15
6.2- Attaching and removing the PIEZON® handpiece.....	15
6.3- Attaching and removing the PIEZON® PI MAX instrument tip.....	16
6.4- Attaching and removing PIEZON® instruments	17
DEVICE USE	17
1- GBT SETTINGS	17
2- INTERFACES	17
2.1- PIEZON® power setting	18
2.2- AIRFLOW®/PERIOFLOW® pressure setting.....	19
2.3- Wireless pedal battery saving	19
3- TREATMENT SEQUENCE.....	19
3.1- Patient and dental professional precautions	19
3.1.1- Patient preparation	19

3.1.2- Dental professional preparation	19
3.2- Disclose	19
3.3- AIRFLOW® and PERIOFLOW®	20
3.3.1- Risk of emphysema	20
3.3.2- AIRFLOW® treatment.....	20
3.3.2.1- Recommendations	20
3.3.2.2- Recommended position and movement.....	20
3.3.2.3- Settings	21
3.3.3- PERIOFLOW® treatment	21
3.3.3.1- Absolute restrictions.....	21
3.3.3.2- Recommended use.....	21
3.3.3.3- Settings	22
3.3.4- How to start AIRFLOW®/PERIOFLOW® treatment	22
3.4- PIEZON® treatment	22
3.4.1- Recommended use	22
3.4.2- Use and settings.....	23
3.4.3- How to start PIEZON® treatment.....	24
3.4.4- End of treatment.....	24
3.4.4.1- Fluoride protection.....	24
3.4.4.2- Post-treatment recommendations.....	24
4- CLEANING AND REPROCESSING	24
4.1- Water lines cleaning	24
4.2- Device cleaning and parts reprocessing	26
5- MAINTENANCE	27
5.1- AIRFLOW® MAX/PERIOFLOW® MAX handpiece daily maintenance	27
5.2- AIRFLOW® MAX/PERIOFLOW® MAX handpiece leakage	27
5.3- PIEZON® handpiece leakage	27
5.4- Light guide check & replace	27
5.5- Wear.....	28
5.6- Handpiece cord system replacement	28
5.7- Monthly check.....	28
5.8- Preventive maintenance and repair	29
5.9- Pairing a (new) pedal	29
6- TROUBLESHOOTING	30
6.1- For AIRFLOW®/PERIOFLOW® products	30
6.2- For PIEZON® products.....	30
6.3- For the device.....	31
6.3.1- Device troubleshooting	31
6.3.2- Symbols troubleshooting.....	32
6.4- Contact EMS aftersales service	33
6.5- Report an adverse event	33
SUSTAINABILITY	34
1- DISPOSAL OF WASTE.....	34
2- SUSTAINABLE DESIGN	34
WARRANTY	34
TECHNICAL DATA COLLECTION AND PRIVACY POLICY	34
TECHNICAL DESCRIPTION	35
SYMBOLS.....	36
ELECTROMAGNETIC COMPATIBILITY	37
1- INTENDED ELECTROMAGNETIC ENVIRONMENT	37
2- DEVICE PERFORMANCE AND EMC DISTURBANCE EFFECTS	37
3- CLINICAL IMPACTS	37
4- OPERATOR ACTIONS.....	37
5- COMPLIANCE LEVELS	37
6- PROXIMITY FIELDS FROM RF WIRELESS COMMUNICATIONS EQUIPMENT.....	38
7- ELECTROMAGNETIC EMISSIONS COMPLIANCE.....	38
RADIO EQUIPMENT COMPLIANCY	39
1- FCC STATEMENTS.....	39




1.1- RF Exposure mobile Device	39
1.2- RF Exposure portable device only for RFID	39
2- ISED STATEMENTS.....	39
2.1- RF Exposure mobile Device	39
2.2- RF Exposure portable device only for RFID	40
3 - WIRELESS COMMUNICATION MODULE, ONLY FOR COSTA RICA	40


BEFORE USE

Before using this device, please carefully read and follow the recommendations of this instruction manual, the reprocessing manual and the powder instruction manual. Please pay special attention to the safety precautions.



Any serious incident that has occurred in relation to the device should be reported to the manufacturer and the competent authority of the Member State in which the user and/or patient is established.

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 device or any of its accessories or component.

 Always keep these Instructions close at hand. These Instructions are only applicable to the equipment they were delivered with.

To prevent injury to persons or damage to property, please observe the corresponding directives and symbols.

  The Instructions for Use of the device is part of the product documentation and is provided in electronic format. However if you want it 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 electronic Instructions for Use (eIFU) of the device is available for download in PDF format at www.ems-instruction.com using the Product Name: GBT Machine AIRFLOW® Prophylaxis Master or Reference: FT-300. A PDF reader is required and, in case of need, it can be downloaded from the same web site.

► 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.

INTENDED USE

The GBT Machine AIRFLOW Prophylaxis Master combines the functions of an ultrasonic scaler and air-polishing unit within a single chassis. The GBT Machine AIRFLOW Prophylaxis Master is intended for use in the following dental and periodontal applications:

- Removing supra and subgingival calculus deposits and stains from teeth
- Periodontal pocket lavage with simultaneous ultrasonic tip movement
- Scaling and root planing

The GBT Machine AIRFLOW Prophylaxis Master is intended for use in the cleaning and polishing of teeth by the projection of water, air, and dental powders onto the tooth surface. The device removes dental plaque, soft deposits, and surface stains from pits, grooves, interproximal spaces, or smooth surfaces of teeth.


The GBT Machine AIRFLOW Prophylaxis Master can be used for the following cleaning procedures:

- plaque removal for placement of sealants
- surface preparation prior to 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 bonding ortho brackets
- cleaning implant fixture prior to loading
- stain removal for shade determination
- plaque removal prior to fluoride treatment
- plaque and stain removal prior to whitening procedure

The GBT Machine AIRFLOW Prophylaxis Master is also intended for use as an air-polisher in patients suffering from periodontal disease. The GBT Machine AIRFLOW Prophylaxis Master is indicated for the non-surgical removal of subgingival plaque in pockets up to 5 mm after initial periodontal treatment.

INTENDED USERS

Only qualified 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.

 The device must be prepared and maintained only by persons who have been instructed in infection control, personal protection and patient safety.

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. Please contact your local EMS representative for further information.

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

PATIENT POPULATION

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.

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


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


INTENDED ENVIRONMENT

The device is intended to be used in a dental cabinet/hospital complying with each country's regulations.

CONTRAINDICATIONS

AIRFLOW®


 Patients suffering from chronic bronchitis or asthma must not, under any circumstances, be treated with an air polishing device. The jet of air and powder could cause respiratory difficulties.


 Patients on a low salt diet must not be treated with the powder containing sodium bicarbonate. For patients on a low salt diet use the PLUS powder without sodium bicarbonate provided by EMS.


PERIOFLOW®


The treatment of deep periodontal pockets can cause bacteraemia. Please apply appropriate restrictions for the treatment of risk patients:

- ▶ Endocarditis
- ▶ Pregnancy, breast feeding
- ▶ Contagious disease
- ▶ Immune deficiency (neutropenia, angranulocytosis, diabetes, hemophilia)
- ▶ Patients under treatment (radiotherapy, chemotherapy, antibiotics)

 The air jet and powder may cause breathing difficulties. Please apply appropriate restrictions for the treatment of risk patients: Patients suffering from chronic bronchitis or asthma must not be treated under any circumstances with this product.

 Predisposed persons may be sensitive to the powder. If allergic reactions are observed, stop using the product and completely remove it.

 The single use nozzle must be used for one single patient only. Never reuse a nozzle because treatment will be ineffective and the risk of emphysema would increase.

 The use of any other powder than the EMS powders for subgingival application would reduce the nozzle's service life. As a result the treatment would become ineffective and would increase the risk of emphysema.

PIEZON®

EMS recommends not to treat patients with a cardiac pacemaker or a defibrillator with this product. The functionality of these devices may be affected by the high frequencies of the ultrasonic oscillations.


Powders


Refer to the instructions for use of the specific powder.

RECOMMENDATIONS AND SAFETY PRECAUTIONS


Only use EMS products together.


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

 Only use this product for the intended indications. Please refer to treatment section. Carefully read these operating instructions before using the product. This also applies to any product used with this system. Failure to observe the operating instructions may result in the patient or user suffering serious injury or the product being damaged.


 Follow the recommendations of the "Reprocessing Instructions" manual (FB-358/NA) regarding procedure. Always examine your AIRFLOW®, PERIOFLOW® and/or PIEZON® products for damage before starting treatment. Damaged product must not be used and must be replaced. Only use original EMS spare parts and accessories.

 DO NOT modify this equipment and/or any of its accessories. No modification of any part of this medical device is allowed.

 DO NOT direct the AIRFLOW® output towards openings of the salivary ducts as this may cause temporary pain and redness.

 Most dental procedures involve contaminated aerosols which could represent a risk factor. Follow the recommendations of treatment and patient and dental professional precautions, in order to minimize the risk.

Specific recommendations for the device:

 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.

 Keep a minimum distance of 25 cm from any source 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 an explosion may occur.

⚠ 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 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.

⊘ DO NOT open the device. There are no serviceable parts inside.

⊘ DO NOT store the powder near acids or heat sources.

⊘ DO NOT direct the jet of powder toward fillings, crowns or bridgework as this could damage these restorations.

! 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.

! Do not leave the device unattended when the water supply is under pressure: risk of flooding. The water supply must be turned off when the device is not in use.

! The installation and connection of the unit must be carried out by a qualified technician.

! EMS cannot be held liable for damage caused by noncompliance with these Warning and Safety instructions.

! Use secure networks to prevent any unauthorized access to your data, which are encrypted for security purposes.

! If a vulnerability is detected, contact technical support via emsrepairs@ems-na.com and disconnect the device from the network if necessary.

COMPATIBILITY

This device is compatible with the following accessories:

AIRFLOW® Powders	series of DV-164, DV-165, DV-175
AIRFLOW® MAX Handpiece	EL-308
PERIOFLOW® MAX Handpiece	EL-354
PIEZON® Handpieces	EN-060, EN-061
PIEZON® Periodontal instruments	DS-001A, DS-011A, DS-016A, DS-083A, DS-084A
PIEZON® PI MAX instruments	DS-010A, DT-065A
GBT Station	DW-100

Applied Parts:

The following items are Medical Device Applied Parts:

- ▶ AIRFLOW® MAX Handpiece (EL-308)
- ▶ PERIOFLOW® MAX Handpiece (EL-354)
- ▶ PIEZON® Handpieces (EN-060, EN-061)

⚠ Applied Parts, under certain operating conditions, may exceed 41°C of temperature and reach a maximum temperature of 48°C.

CYBERSECURITY

1- GENERAL INSTRUCTIONS AND RECOMMENDATIONS

- ▶ Use secure networks to prevent any unauthorized access to your data, which are encrypted for security purposes.
- ▶ The device requires a 2.4 GHz Wi-Fi network.
- ▶ If a vulnerability is detected, contact technical support via emsrepairs@ems-na.com and disconnect the device from the network if necessary.

2- NETWORK INTERFACES AND CONNECTIVITY

- ▶ Two jack connectors: input and output, used for connection to satellites and for maintenance by EMS technicians only.
- ▶ Wi-Fi (Radio Interface No. 1): input and output, used for cloud connection (data transfer and updates).
- ▶ LTE (Radio Interface No. 2): input and output, used for cloud connection (data transfer and updates).
- ▶ RFID (Radio Interface No. 3): input, for handpiece recognition.
- ▶ Bluetooth Low Energy (Radio Interface No. 4): input, for wireless pedal connection to the device.

Detailed technical specifications of each interface are provided in the table "Wireless Communication Module." in the section Technical Description.

3- SOFTWARE MANAGEMENT AND UPDATES

- ▶ The device automatically receives software updates via the cloud.
- ▶ The user is notified by an icon displayed on the screen.
- ▶ Only software versions digitally signed by EMS can be installed.
- ▶ Users also receive proactive notifications in case of potential malfunction or maintenance needs.

4- RESPONSE TO CYBERSECURITY INCIDENTS

- ▶ In case of a security event, EMS may interrupt the cloud connection.
- ▶ EMS customer service will contact the user if necessary.
- ▶ The device continues to operate without cloud connection.

5- SUPPORT AND LIFECYCLE

Software support is provided as long as the device is commercially available.

6- DECOMMISSIONING



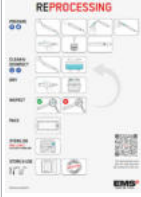


















- ▶ The device does not transmit or store any sensitive data.
- ▶ Therefore, no specific data deletion procedure is required upon decommissioning.

INSTALLATION

1- EQUIPMENT INCLUDED IN THE BOX

! Check contents for any damage that may have occurred during transportation (device, consumable or accessory).

GBT Machine AIRFLOW® Prophylaxis Master CLASSIC CONFIGURATION/ GBT READY CONFIGURATION

	GBT Machine AIRFLOW® Prophylaxis Master Unit with Master Screw, water & air filters installed 1x FT-300/*¹		Quick Guide 1x FB-1023/US		Reprocessing posters 1xFA-844 1xFA-887/EN
	Customer Flyer 1xFA-888/EN		AIRFLOW® COMFORT 1x DV-164/MIN		AIRFLOW® PLUS 1x DV-165/Z (3x DV-165/Z)
	CLEANER bottle 1x EG-1000		Water bottle 2x EG-121		GBT Machine Wireless Pedal 1x EK-1055 with 2x AA 1.5V type lithium batteries
	US Power cord 1 x CD-137		Air hose 1x EH-142 Water hose 1x EG-110		Powder chambers 1x EL-607 (Plus) 1x EL-606 (Classic)
	AIRFLOW® MAX Handpiece cord (1,80m) 1x EM-1007		PIEZON® Handpiece cord (1,80m) 1x EM-1009		PSL/PSR Instruments 1x FS-461
	AIRFLOW® MAX application 1x EL-308: AIRFLOW® MAX Handpiece 1xAB-470A/A (FV-083 ²): Easy Clean 1x FS-465 (3x FS-465)		PERIOFLOW® MAX application 1xEL-354: PERIOFLOW® MAX Handpiece 20x AB-1010 : PERIOFLOW® Nozzle 1xFS-474		GBT Machine AIRFLOW® Prophylaxis Master Maintenance KIT 1xEL-605 Perio cap 1xEL-655 Set CLIP+CLEAN 2xEL-599 Air cartridge assembly 2xEL-1207 Filters enclosure assembly 2xAB-348/B Flat seal ø13x1.1x2.05 3xBC-1039 O'ring Ø1.50x1.00 VMQ Precision 1xEL-1211 AIRFLOW cord seals set for AFPM 2.0 1xFS-600
	PI MAX Introduction Kit 1x FV-117 PI MAX Tool 1x DS-010A/A PI MAX Instrument 1x DT-065A/A 4x PI MAX Instrument tips 1xFS-295#C		PIEZON® 1PS application 1xEN-060 : PIEZON® LED Handpiece 1x DS-016A : Instrument PS 4x AB-340 (FV-065): Light guide 1x FS-455 (3x FS-455)		GBT training tool kit 1x FV-122

¹Two versions of the device exist to meet local requirements: with LTE (FT-300/A) and without LTE (FT-300/B). This difference is not visible on the device.
²Reference to be used for individual accessory order

2- STEP-BY-STEP INSTALLATION

Scan for support on installing your device on the chair or station



2.1- Find an appropriate area to place the device

! Place the device with the recommended GBT Station within the dental cabinet in a suitable position for your activity and leave enough free space to allow easy handling and proper aeration.

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

! Avoid using this device near other equipment, because it could result in improper function. Keep a minimum of 10 cm clearance around the unit.

If not possible, verify that the device and other equipments are functioning normally.

The device must be placed on a secure and flat surface (slope < or = 5°).

2.2- Check for proper water and air supply lines

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

! 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 hoses (cable) to EN-1717 or DVGW³ compliant fluid sources.

2.3- Check for a proper and safe power grid

! This device uses a Class-I insulating system that requires protective earth.

! 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.

! The mains plug of the unit must be accessible at all times.

⊘ 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.

2.4- 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.

! The device uses a low power radio, 6 dBm EIRP max, Bluetooth[®] 2.4 GHz, to communicate with the wireless pedal. Interference may occur in the vicinity of this equipment. The device uses Wi-Fi 2.4 GHz and Mobile network are used for the connectivity services.

The device uses also RFID/NFC for AIRFLOW[®] MAX and PIEZON[®] handpieces recognition.

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.

2.5- Device connectivity

Your GBT Machine AIRFLOW[®] Prophylaxis Master can connect to the internet via Wi-Fi. To connect through Wi-Fi, please refer to the instructions in the section 2.5.1- Connecting / Disconnecting from the Wi-Fi.

The GBT Machine AIRFLOW[®] Prophylaxis Master's connectivity enables continuous treatment data collection and transmission to Electro Medical Systems. The device collects data such as PIEZON activity, AIRFLOW activity, type of handpieces used, hourly usage, etc., to monitor the state of your device and provide access to this information through your dashboard on my.ems-dental.com. This allows for regular software updates and proactive notifications in case of potential failures or maintenance needs. Our goal is to help our customers and technical team improve treatment plans and device usage.

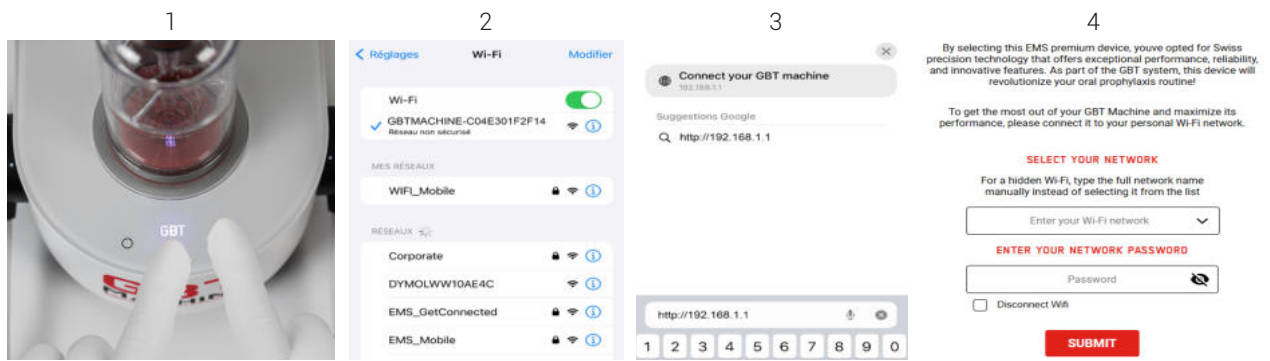
Ensure you are in an area with strong signal reception and avoid sources of electromagnetic interference. Use secure networks to prevent unauthorized access to your data, which is encrypted for security. The device adheres to relevant data protection regulations, such as GDPR/HIPAA. Contact customer support if you encounter any connectivity issues. If connectivity is lost, the Wi-Fi logo on your GBT Machine AIRFLOW[®] Prophylaxis Master will turn orange. Check if the connectivity works.

2.5.1- Connecting / Disconnecting from the Wi-Fi

Setting up the Wi-Fi connection/ disconnection is straightforward—just follow the steps below during installation:

Put your device (phone, computer, etc.) in airplane mode.

⚠️ Ensure that only one user is connected to the device's WiFi at a time.



Connect

Press simultaneously the GBT level and high level buttons and hold them for 2 seconds.



A sonar sound will start playing (if not, repeat the step above) and the Wi-Fi symbol will light up and fade several times, connection search is in progress.

Connect to "GBTMACHINE-XXXX" Wi-Fi network with your phone.

Go on internet browser. Tap <http://192.168.1.1> in the search bar or scan the QR code.



If the web page does not open, try another browser.

Choose your Wi-Fi network you want to connect your device to with the drop-down list or enter the Wi-Fi network name manually (if hidden) and enter your password.

If successful, the Wi-Fi logo & the LEDs on the machine will blink twice.

Your GBT Machine AIRFLOW® Prophylaxis Master is connected to the Wi-Fi !

Disconnect

Same that above

Same that above

Same that above

Check the Disconnect Wi-Fi box, then click on the Submit button.

If you enter your login credentials incorrectly during the 4th step of your device's connection process, the sonar sound will persist. Please turn your device off and on again, then restart the connection process from the first step.

2.6- Connect air and water hoses

Turn the device over and place it upside down.

- | | | |
|---|---|---|
| 1 | <p>In the absence of the GBT Station, connect the air hose EH-142 to the cabinet/dental unit.
Push the hose connector into the air jack firmly (it may be hard).</p> | <p>Pressure: 4.5 to 7 bar. Ideally 6 bar
Dry air. Max. humidity: 1.032 g/m3
Filtration: max. 1 µm</p> |
| 2 | <p>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.
🚫 DO NOT install the PIEZON® or CLEANER bottles before connecting the air and water lines.</p> | <p>Drinking water
Pressure: 2 to 5 bar
Salinity: max. 0.2%
Temperature: 10°C to 30°C</p> |

2.7- Install accessories

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

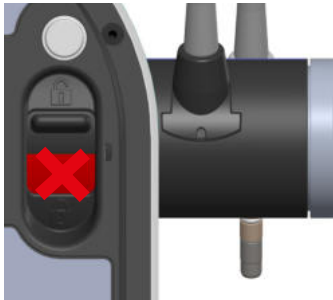


- 1 **EH-142**
Air hose – filter pre-installed
USE HARD FORCE
- 2 **EG-110**
Water hose – filter pre-installed
- 3 **EG-121**
Water bottle - filter pre-installed
For the use of bottled water, the water goes through the dedicated filter.
- 4 Power cord into socket
(Fuse holder in the socket)
- 5 **EM-1007**
AIRFLOW® MAX handpiece cord + lock actuator
USE FORCE
- 6 **EM-1009**
PIEZON® handpiece cord + lock actuator
USE FORCE

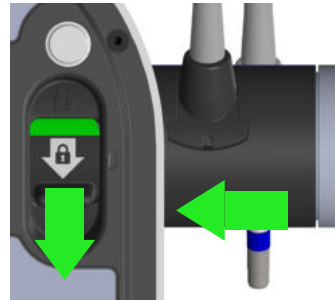
2.8- Check the handpiece cord system connections

 First disconnect the mains plug before connecting/disconnecting any handpiece cord system.

Scan for support



The handpiece cord system is not fully connected.



USE FORCE to lock in.

The system is well connected & locked.

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

2.9- Fix the device

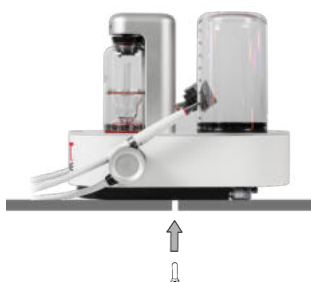
Fix the device to the recommended GBT Station

Follow the quick guide delivered with your GBT Station

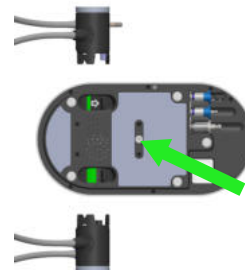
Fix the device to the working area other than the GBT Station

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).



Master Screw usage



Master Screw placement

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

2.10- Power your device

You can now connect the power cord to the mains grid.

- ⚠ Protective earth is required! Be sure your power grid has an efficient protective earth.
Settings: Voltage: 100-240 Vac - Frequency: 50 to 60 Hz - Operating current: 4 A max.

2.11- Installation of the wireless pedal



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

The 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).

Scan for support





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

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

- ⚠ The wireless pedal uses a low power, 6 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.

3- POWDER CHAMBERS

<p>PLUS</p> 	<p>⚠ The PLUS Powder chamber (with the red base) is designed for the PLUS Powder. It can be used for supragingival and subgingival treatments.</p> <p>Pressure is automatically reduced for compatibility with subgingival treatments, including PERIOFLOW® treatments.</p> <p>Compatible EMS Powders: series of DV-070, DV-082 (PLUS and PERIO)</p>
<p>CLASSIC/COMFORT</p> 	<p>⚠ The CLASSIC/COMFORT Powder chamber (with the white base) is designed for the CLASSIC/COMFORT Powder and should be used only for supragingival treatments.</p> <p>Compatible EMS Powders: series of DV-164 (CLASSIC/COMFORT)</p>

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

- ❗ Check powder chamber for any cracks: There should be no crack on the body.

- ❗ Make sure that the powder chambers are dry.

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

- ❗ Only use powders for their own intended use. Please refer to the powder specific instructions for use.



Filling the powder chambers :



- ❗ 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.
- ⚠️ DO NOT fill the powder chamber beyond the MAX level. Pour the powder in freely. The central tube can be fully filled without problem.
- ❗ Shake the powder chamber after each filling and at least once a day.



- Before pressurizing, position the powder chamber into the device. Magnetic attraction will position it correctly.
- ❗ DO NOT insert upside-down.

4- WATER SUPPLY AND WATER BOTTLE

What water should I include in my water bottle?

EMS recommends the use of:

- ▶ Filtered drinking water
- ▶ Drinkable tap water (≤ 500 CFU/mL bacteria)
- ▶ Distilled water

The temperature must be between 10°C and 30°C. All other liquids can damage your device and void your warranty.

How to fill your water bottles ?

- ▶ Fill both bottles each morning with filtered water or drinkable tap water (keep one as a back up to ensure uninterrupted treatment flow).
- ▶ To remove and place the bottle, use a straight up and down movement (do not shake the bottle to avoid damages).
- ▶ At the end of every day: empty your bottles and allow to dry.
- ▶ Once a week: wash your water bottles with warm soapy water, rinse thoroughly and allow to dry.
- ▶ It is also recommended to use a bottle cleaning agent weekly.

Without Bottle:

PIEZON® & AIRFLOW® use external water supply.



With Bottle connected:

PIEZON® & AIRFLOW® use bottle liquid supply.



⚠️ The CLIP+CLEAN shall be previously cleaned and sterilized before use.

Non-sterilized CLIP+CLEAN may contaminate the device.

❗ Place the CLIP+CLEAN into the device's bottle receptacle for dust protection.

Connect the water bottle

❗ DO NOT use disinfectant solutions (e.g., chlorhexidine) inside the water bottle during treatment.

❗ DO NOT sterilize the water bottle and its nozzle cap by steaming or dry thermal reprocessing. Use only ambient temperature active disinfectant and cleaning agents.

5- AIRFLOW® MAX AND PERIOFLOW® MAX HANDPIECES

5.1- Before use

⚠️ EMS AIRFLOW® MAX and/or PERIOFLOW® MAX handpieces are supplied non-sterile and must be cleaned and sterilized before first use and between patient uses. Non reprocessed products may cause bacterial or viral infections.

⚠️ Follow the recommendations of the "Reprocessing Instructions" manual (FB-358/NA) regarding procedure for cleaning and sterilizing the components, and the present-day regulations on reprocessing in effect in your country.

5.2- Attaching and removing AIRFLOW® MAX or PERIOFLOW® MAX handpiece

In order to ensure perfect electronic connection, the individual components must be dry.

Attaching the AIRFLOW® MAX or PERIOFLOW® MAX handpiece



Connect the handpiece to the AIRFLOW® MAX handpiece cord.

Removing the AIRFLOW® MAX or PERIOFLOW® MAX handpiece



Disconnect the handpiece from the AIRFLOW® MAX handpiece cord by turning and pulling at the same time.

5.3- Attaching and removing PERIOFLOW® nozzles



Single-use PERIOFLOW® nozzle.



Cannot be reprocessed.



DO NOT use the PERIOFLOW® nozzle if the package is damaged or opened.



Fully connect the PERIOFLOW® nozzle by pushing on a hard surface.

Make sure the PERIOFLOW® nozzle is correctly attached = fully inserted and right way



Remove the nozzle by using the PIEZON® instrument check tool after treatment.



Risk of injury: Always USE the PIEZON® instrument check tool.



DO NOT remove by hand.

Scan for support



6- PIEZON® HANDPIECE AND INSTRUMENTS

6.1- Before use

⚠ EMS PIEZON® products (handpieces, instruments and tools) are supplied non-sterile and must be cleaned and sterilized before first use and between patient uses. Non reprocessed products may cause bacterial or viral infections.

⚠ Follow the recommendations of the "Reprocessing Instructions" manual (FB-358/NA) regarding procedure for cleaning and sterilizing the components, and the present-day regulations on reprocessing in effect in your country.

⚠ Check regularly instrument length with the PIEZON® instrument check tool.

If PIEZON® instrument extremity reaches or is shorter than the limit indicated by the GBT Logo, it can have excessive and uncontrolled vibrations. Replace the tip.

6.2- Attaching and removing the PIEZON® handpiece

In order to ensure perfect electronic connection, the individual components must be dry.

Attaching the PIEZON® handpiece



Connect the PIEZON® handpiece to the PIEZON® handpiece cord.

Removing the PIEZON® handpiece



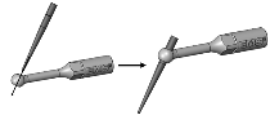
Disconnect the PIEZON® handpiece from the PIEZON® handpiece cord.

6.3- Attaching and removing the PIEZON® PI MAX instrument tip

In order to ensure perfect electronic connection, the individual components must be dry.
The PIEZON® PI MAX instrument is composed of:



Attaching the
PIEZON® PI MAX
instrument tip



Pre-Insert the TIP in the HOLDER.



Take the PI TOOL. Remove the SCREW from the PI TOOL base.
Gently insert the TIP in the PI TOOL feature marked CONNECT.



Gently tilt the PI in the PI TOOL base. Slide it down to STOP.



Insert the SCREW and screw until the head of the screw is in contact with the body of the tool.



Unscrew the SCREW.

Gently remove the PIEZON® PI MAX with new TIP inserted.



If needed, remove the small plastic chip with your finger.

Removing the
PIEZON® PI MAX
instrument tip



Take the PI TOOL. Remove the SCREW from the PI TOOL base.
Gently insert the TIP in the PI TOOL feature marked DISCONNECT.



Gently tilt the PI in the PI TOOL base. Slide it down to STOP.



Insert the SCREW and screw until the head of the screw is in contact with the body of the tool.

The TIP is disconnected from the HOLDER. Take it with fingers.



Unscrew the SCREW and get back the HOLDER.

6.4- Attaching and removing PIEZON® instruments

In order to ensure perfect electronic connection, the individual components must be dry.

Attaching the
PIEZON® instrument



Mount the PIEZON® instrument using the CombiTorque®.

⚠ Only use the CombiTorque® to tighten the PIEZON® instrument on the PIEZON® handpiece to the correct torque to avoid instrument unscrewing or breaking

! 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®. Gently remove the CombiTorque® following the shape of the PIEZON® instrument.

Removing the
PIEZON® instrument



Gently place the CombiTorque® following the shape of the PIEZON® instrument. Unscrew it counterclockwise.

DEVICE USE

1- GBT SETTINGS

Guided Biofilm Therapy (GBT) is a standardized protocol to remove biofilm, stains and calculus from natural teeth, restorations and implants. The default GBT settings set at medium power and high water levels for both PIEZON and AIRFLOW technologies - are one of the innovations of the GBT Machine AIRFLOW® Prophylaxis Master, allowing it to:



Ensure Consistency: Easy implementation.






Improve Ergonomics: For better clinician experience

The GBT settings are designed to benefit both patients and clinicians. For more understanding, please refer to the Swiss Dental Academy (SDA).

2- INTERFACES



1	ON/ OFF- Standby mode	ON: the device goes into operating mode. OFF: the device reverts back to standby. After 1 hour of inactivity, the device switches automatically to standby mode, and the powder chamber depressurizes automatically.	
2	Powder chamber	To pressurize or depressurize the powder chamber, use the button on the caliper. A white light will turn on to indicate the powder chamber is pressurized. When depressurizing, the AIRFLOW® MAX handpiece cord will automatically purge, and the white light will turn off when done. ! Depressurizing the powder chamber can take up to 10 seconds. When not in use, keep the AIRFLOW® MAX handpiece in its holder with the nozzle facing down to prevent upward spraying of air and powder. If the powder chamber is not pressurized, the device will operate in water-only mode.	

3	Power setting	<p>First lift the handpiece from his holder. Press the touch buttons to adjust AIRFLOW® and PIEZON® power. White LED lights indicate your selection:</p> <ul style="list-style-type: none"> • Low level: Left LED light • GBT power setting: Left and central LED lights • High level: Left, central and right LED lights <p>By default, each time you put AIRFLOW® MAX or PIEZON® handpiece back on the device, it goes back to GBT power setting.</p>	<p>Scan for support</p> 
4	Water setting	<p>First lift the handpiece from his holder. Adjust PIEZON® and AIRFLOW® water flow rate by turning the control knob on the handpiece cord system (touch buttons does not work for water setting):</p> <ul style="list-style-type: none"> • AIRFLOW®: Right control knob • PIEZON®: Left control knob <p>Blue LED lights indicate your selection:</p> <ul style="list-style-type: none"> • Low level: Left LED light • Medium level: Left and central LED lights • GBT water setting: Left, central and right LED lights <p>By default, each time you put AIRFLOW® MAX or PIEZON® handpiece back on the device, it goes back to GBT water setting.</p>	 <p>AIRFLOW</p>  <p>PIEZON</p>
5	Pedal (normal)	<p>Press the edge of the foot pedal for normal operation. At least one handpiece cord system is required to operate the device. The foot pedal cannot be activated when both handpieces are placed in their holders, and when the device is not in use.</p>	
6	Pedal BOOST	<p>Using BOOST provides a convenient way of temporarily increasing the power/pressure within the current setting (Low, GBT, High). BOOST is activated by pressing hard on the center of the wireless pedal with the heel up and deactivated by returning the foot to normal pedal activation (heel down). See Sections 2.1 and 2.2 below for the BOOST power and pressure levels.</p>	 <p>NORMAL Press gently the border</p> <p>BOOST Press hard the center</p>

2.1- PIEZON® power setting

The unit is equipped with a technology which provides a dynamic power regulation in function according to the load applied to the instrument.



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

Power settings	Low	GBT	High
Max output power (W)	1.5	3.4	4.4
BOOST (W)	2.4 - 4.8	3.4 - 6.8	4 - 8

2.2- AIRFLOW®/PERIOFLOW® pressure setting

Both the PLUS and CLASSIC/COMFORT 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 pressures⁴ as per selected powder chamber and user power setting:

Pressure settings	Low	GBT	High
Static (Bar)	2.3	2.65	3.15
AIRFLOW® PLUS dynamic (Bar)	1.6	1.95	2.45
BOOST AIRFLOW® PLUS (Bar)	2.45	2.8	3.0
Pressure settings	Low	GBT	High
PERIOFLOW® dynamic (Bar)	2.0	2.3	2.7

2.3- 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).

3- TREATMENT SEQUENCE

3.1- Patient and dental professional precautions

- ⚠ For your protection and the protection of your patient, start by gargling patient mouth with antimicrobial mouth rinse.
- ⚠ Always refer to treatment recommendations manual (FA-892/US) before using the device.

3.1.1- Patient preparation

⚠ Eye protection is mandatory.

It is also recommended to remove the patient's glasses and optical lenses.



⚠ GBT VISIGATE for lip and cheek retractor protection is recommended for clinician convenient.

Not available in the device configuration.

GBT VISIGATE® is an auxiliary aid for clinical use which allows the working field to be accessed with ease. It provides retraction of lips and cheeks during dental treatment, offers the clinician increased visibility and accessibility and facilitates saliva and moisture control in the oral cavity.

The device is placed in the mouth by the clinician before treatment to retract the lips and cheeks, then removed after the procedure.

For detailed instructions on use and contraindications, please refer to the GBT VISIGATE® Instructions for Use

⚠ If PERIOFLOW® treatment has to be performed, radiographs is mandatory to correlate clinical probing depth

3.1.2- Dental professional preparation

Protect yourself with the following measures:



Wear protective mask



Wear protective glasses



Wash your hands



Wear protective gloves

Additional personal protective equipment can be used.

⚠ GBT FLOWCONTROL® and saliva ejector are mandatory. They evacuate the air/powder mixture deviated by the treated tooth. Study⁵ on aerosol management shows that if AIRFLOW® is used as recommended, the risk is negligible for the clinician.

3.2- Disclose

Use the Biofilm Discloser to reveal biofilm on all hard tissues. Remove excess disclosing using AIRFLOW® MAX handpiece without pressurizing the powder chamber (water-only mode) and with High level water, with the GBT Flowcontrol. Please refer to the instruction provided with the product.

⁴ 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® MAX handpiece with DV-082 powder.

⁵ Aerosols in Dentistry: The Bacterial Contamination of the Room Air During an AIRFLOW® Treatment. Marcel Donnet, Klaus-Dieter Bastendorf, Magda Mensi, Adrian Lussi. www.ZM.ONLINE.de 12/2020

3.3- AIRFLOW® and PERIOFLOW®

⚠ Please refer to the section "CONTRAINDICATIONS" for information regarding AIRFLOW® and PERIOFLOW®.

3.3.1- Risk of emphysema

⚠ To limit the risk, always follow contraindications, recommendations and detailed instructions.

Subcutaneous emphysema occurs as a result of an abnormal introduction or presence of air or gas into tissue or tissue spaces. It has been recognized and documented as a complicating factor of any dental procedure where pressurized air has been used. Expedient diagnosis and management of subcutaneous emphysema are important to facilitate recovery.

Clinical signs of emphysema:

- ▶ Crackling of mucosa upon pressure (subcutaneous crepitation), pain, swelling, tenderness and discomfort.
- ▶ Often accompanied by facial or neck swelling.

In case of emphysema :

- ▶ Immediately stop the procedure
- ▶ Determine the extension and location
- ▶ Observe the patient for 30min and if the emphysema is not extended outside the oral cavity, compress to reduce it (from the apex).
- ▶ Patient may need antibiotics and anti-inflammatory (according to patient health), in case the dental professional cannot prescribe antibiotics, kindly refer the patient to a physician or a hospital for the recommended medicine.
- ▶ If the emphysema extends outside the oral cavity, particularly if there is a compression or constriction of airways or patient has difficulties to breath, call the emergency service

3.3.2- AIRFLOW® treatment

3.3.2.1- Recommendations

⚠ Avoid touching hard and soft tissues with the handpiece directly. Keep a minimum distance of 1-2 mm.

⚠ When you remove your foot from the pedal, the air/powder jet continues for a few more seconds, so continue to use the GBT FLOWCONTROL®.

⊘ DO NOT use AIRFLOW® products on patient with uncontrolled acute bronchitis/asthma or upper respiratory tract disease or infection during treatment.

⊘ DO NOT use AIRFLOW® products in specific sites :

- Sites with suppuration on probing, including open wounds

3.3.2.2- Recommended position and movement

Aerosol management:

- ▶ ⚠ Use GBT FLOWCONTROL®
- ▶ Direct the jet projections towards the cannula

For optimal efficiency:

Angle

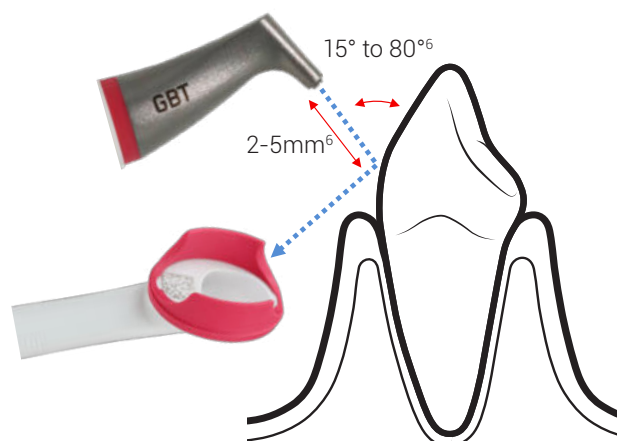
- ▶ Avoid using the AIRFLOW® MAX handpiece at 90°
- ▶ Continuously adapt the angle while working
- ▶ Maximal range of usage is between 15°-80°⁶

Distance

- ▶ General rule: With AIRFLOW® MAX, work closer!
- ▶ Keep the handpiece at 3 to 5 mm⁶ during work
- ▶ In case of heavy stains, keep the handpiece at max 2mm⁶

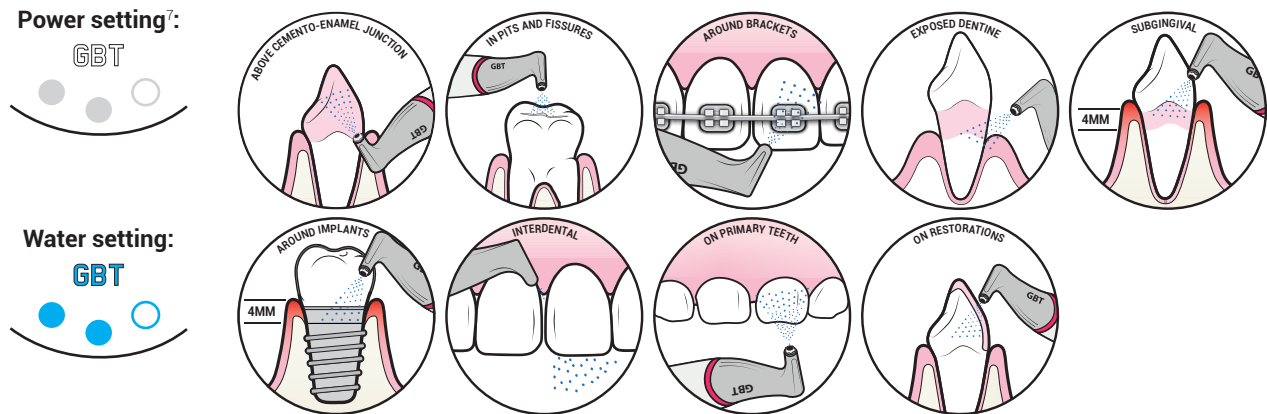
Movement

- ▶ Make continuous semi-circular movement
- ▶ Small smileys mesial to distal
- ▶ Never hold the handpiece stationary



⁶ Settings are for AIRFLOW® MAX handpiece. For the other AIRFLOW® handpieces, the angle must be between 30° to 60° and the distance between 3 and 5 mm.

3.3.2.3- Settings



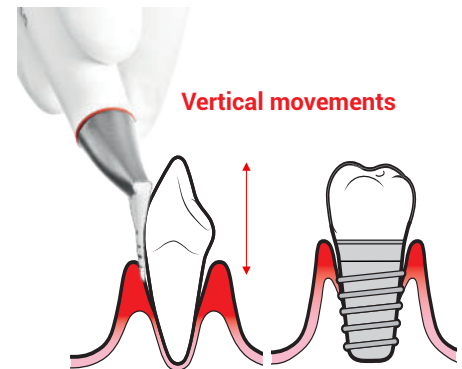
3.3.3- PERIOFLOW® treatment

3.3.3.1- Absolute restrictions

⚠ Please refer to the section "CONTRAINDICATIONS" for information regarding PERIOFLOW®.

3.3.3.2- Recommended use

- ▶ Only use by trained and qualified personnel.
- ▶ Correlate clinical probing pocket depth with radiographs before using the PERIOFLOW® nozzle.
- ▶ Always check the patency of the PERIOFLOW® nozzle before and during use.
- ▶ Single use per patient. DO NOT use the same PERIOFLOW® nozzle in more than 1 patient, PERIOFLOW® nozzle cannot be sterilized or re-used.
- ▶ Use the PERIOFLOW® nozzle between 5-10 secs per site depending on the probing depth.
- ▶ If treating multiple sites in 1 patient, check if the tip of the PERIOFLOW® nozzle is not bent and the quality of the PERIOFLOW® nozzle has not changed.
- ▶ After approx. 20 sites, change the PERIOFLOW® nozzle.
- ▶ In natural teeth, after a 6-point pocket charting, the PERIOFLOW® nozzle is to be used only in sites where probing depths exceed 4 mm.
- ▶ Use your finger and thumb to compress the site.
- ▶ Never push or force the PERIOFLOW® nozzle into the pocket even if the depth is > 4 mm.
- ▶ Use it in a vertical overlapping, repetitive movement. The PERIOFLOW® nozzle must be inside the pocket during the entire cleaning process.
- ▶ Around dental implants, use the PERIOFLOW® nozzle in buccal, lingual, mesial and distal sites - all sites in general.



⚠ PREFERABLY USE the PERIOFLOW® nozzle with tabletop devices.

⚠ ONLY USE AIRFLOW® PLUS or PERIO for subgingival application with PERIOFLOW® nozzle.

⁷ Power adjustment depends on the practitioner's perception and experience.

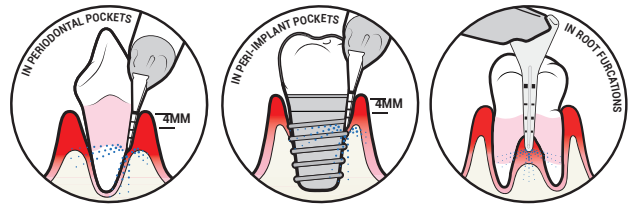
3.3.3.3- Settings

This is dependant on which PERIOFLOW® handpiece being used

Power setting for NEW PERIOFLOW® handpiece (with RFID):



Power setting for OLD PERIOFLOW® handpiece (with no RFID):



Water setting for NEW PERIOFLOW® handpiece (with RFID):



Water setting for OLD PERIOFLOW® handpiece (with no RFID):



3.3.4- How to start AIRFLOW®/PERIOFLOW® treatment

- 1 Connect the WATER bottle (if required).
 - 2 Switch ON the device
 - 3 Position the PLUS powder chamber, filled with PLUS powder, into the device to start GBT treatment
 - 4 Pressurize the chamber
 - 5 Take the AIRFLOW® MAX handpiece or PERIOFLOW® MAX handpiece and nozzle
 - 6 Use the default power GBT settings [or increase the power]
 - 7 Use the default water GBT settings
 - 8 Press the pedal to start treatment.
 - 9 [Step hard on the center of the pedal for BOOST]
 - 10 Release the pedal to stop treatment
 - 11 Put the handpiece back into its holder*
- *Settings come back to default GBT settings
- 12 Depressurize the powder chamber before removing the AIRFLOW® MAX handpiece for sterilization



⚠ 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). Make sure to never point the PERIOFLOW® nozzle toward the patient, during and after operation.

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

3.4- PIEZON® treatment

3.4.1- Recommended use

⚠ Please refer to the section "CONTRAINDICATIONS" for information regarding PIEZON®.

PIEZON® instruments vibrate in a controlled back-and-forth oscillation. During treatment, always hold the instrument parallel to the tooth surface adapting the lateral side of the instrument.

⊘ DO NOT direct the instrument straight to the enamel surface. Never point the tip of the instrument to the tooth surface.

! During PIEZON® treatment, only the last 2mm of the PIEZON instrument, should be in contact with soft tissues in the patient mouth. The metallic part of the PIEZON® handpiece can heat up during a long period of treatment if recommendations are not followed.

! On metal, ceramic restorations, and on prosthetics, ONLY use the PIEZON® PI MAX instrument.

! Listen for a noise change:

▶ For PIEZON® Instruments: a suspicious noise during contactless activation may be a sign of possible damage to the system or improper screwing of the instrument. In this case, check the correct screwing with the CombiTorque®, then the condition of the instrument. If in doubt, contact the support service.

▶ For the PIEZON® Handpiece: it could occur when the instrument is activated in order to detect a possible loosening of the instrument in the handpiece.

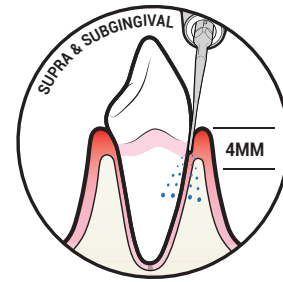
⁸ Power adjustment depends on the practitioner's perception and experience. The PERIOFLOW® MAX handpiece does not have a boost function.

3.4.2- Use and settings

! Always start PIEZON® treatment with default GBT setting and use the BOOST function as necessary (hard calculus).⁹

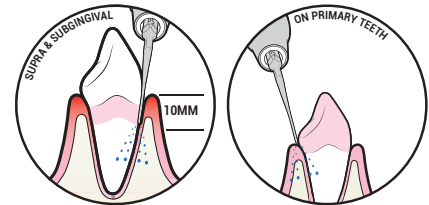
PIEZON® Instrument P

Hard calculus; subgingival up to 4 mm



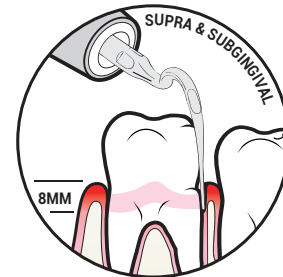
PIEZON® Instrument PS

Supra- and Subgingival up to 10 mm
For 95% of all cases



PIEZON® Instruments PSR/PSL

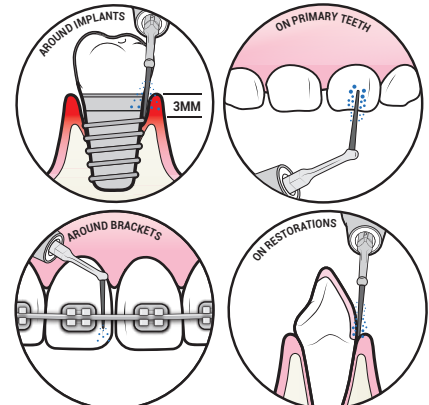
Supra- and Subgingival up to 8 mm



PIEZON® Instrument PI MAX

Subgingival up to 3 mm

⊘ DO NOT use PIEZON® PI MAX with the BOOST mode, it could cause tip breakage.



⚠ Instrument wear goes faster when used on enamel.

⁹ EMS recommends using the GBT setting and the boost if necessary (hard calculus). Power adjustment depends on the practitioner's perception and experience.

3.4.3- How to start PIEZON® treatment

- 1 Connect the WATER bottle (if required).
 - 2 Switch ON the device
 - 3 Take the PIEZON® handpiece.
 - 4 Use the default power GBT settings [or increase the PIEZON® power for supragingival use with PIEZON® PS only]
 - 5 Use the default water GBT settings
 - 6 Press the pedal to start treatment.
 - 7 [Step hard on the center of the Bluetooth pedal for BOOST.]
 - 8 Release the pedal to stop treatment.
 - 9 Put the handpiece back into its holder.*
- * Settings come back to default GBT settings



- ⚠ 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.

3.4.4- End of treatment

After completion of treatment, the patient can do a final rinse with water.

3.4.4.1- Fluoride protection

After the treatment, the teeth are practically free from mucin. It is thus advised to carry out topical fluoride application. It is then important to use a colorless fluoride.

3.4.4.2- Post-treatment recommendations

After completion of the GBT treatment and application of the fluoride protection, patients are recommended not to consume tea, coffee, red wine and/or any food or drink, or smoking, because it could potentially stain the tooth surface for a minimum of 45 minutes.

DURING 45 MINUTES



4- CLEANING AND REPROCESSING

4.1- Water lines cleaning



Keeping the device's water lines clean is recommended to prevent microbial contamination.

A regular cleaning and maintenance protocol should be adopted to clean and protect dental unit waterlines. EMS recommends using EPA-registered dental unit waterline cleaners, VistaClean™ Irrigant Solution Concentrate and VistaTab™ Dental Unit Waterline Cleaner by Hu-Friedy or Monarch Lines Cleaner by Air Techniques.

ⓘ Follow the instructions for use of the product.

The instructions for use should be followed to ensure the appropriate water quality to help protect patients, staff and equipment.

ⓘ Both handpieces should be removed prior to using VistaClean™ /VistaTab™ or Monarch Lines Cleaner.

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

Initial start-up treatment and routine treatment

ⓘ Follow the instructions for use of the product.

For daily water line cleaning: Add drops¹⁰ of VistaClean™ Irrigant Solution Concentrate in the water bottle fully filled (800ml) and use it with patients.

Dental unit waterline cleaner and GBT Machine AIRFLOW® Prophylaxis Master

1- Waterlines cleaning with VistaTab™ Dental Unit Waterline Cleaner or Monarch Lines Cleaner.

ⓘ Follow the instructions for use of the product.

¹⁰ Follow cleaning products manufacturer instructions. Please refer to your public health guidelines.



Place the CLEANER bottle fully filled, switch ON the device and lift both handpieces from their holders. The 3 blue LED lights are ON and a sonar sound will start playing.

❗ Before placing, remove CLIP+CLEAN from the device. Each cleaning consumes 30ml of CLEANER.

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



Remove both handpieces from their cord connectors. Connect the CLIP+CLEAN on the cord connectors and put them over a sink.

Contamination prevention: ❗ DO NOT make any contact between the sink and the handpiece cords.

⚠️ CLIP+CLEAN shall be reprocessed after each use.



Press the pedal, then wait for 20 seconds.

The 3 buttons will blink to indicate the progress. Cleaning can be paused and restarted by pressing the pedal.



Disconnect the CLEANER bottle and release the pressure by unscrewing the cap.

Then screw the cap back on and place the bottle upside down on the device to indicate that the device needs to be rinsed before the next use.

The CLEANER bottle **must be left upside down** on the system overnight, to remind the user that the CLEANER has to be completely flushed out of the system in the morning before treatment.

Every morning before the first patient: Water lines rinse



Place the water bottle fully filled.

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



Hold both handpiece cords with CLIP+CLEAN over a sink.

Contamination prevention: ❗ DO NOT make any contact between the sink and the handpiece cords.



Press the pedal, then wait 20 seconds.

The 3 buttons will blink to indicate the progress. Rinsing can be paused and restarted by pressing the pedal.

Remove the water bottle and rinse well prior to filling with water for the first patient of the day.

- ⚠ DO NOT leave water in your water bottles overnight. Allow the bottles to air with the lids off overnight.
- ⚠ Risk of ingestion of the cleaning agent. Check that no more residue of CLEANER is flushing out of the handpiece 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.

4.2- Device cleaning and parts reprocessing

After every patient

Overall cleaning and disinfection



Use a cleaning and disinfecting wipe (less than 35% alcohol) compatible EPA-registered intermediate level surface disinfectant or that complies with the standards applicable in the country (e.g. MICROKLEEN or Opti-Cide³ wipes).

Reprocess handpieces and instruments. Please refer to FB-358/NA "reprocessing instructions".

- ❗ Wipe and dry the device after disinfection and/or decontamination.
- ❗ Once a week use a damp microfibre cloth with warm water to remove the residue build up that all wipes can leave on your GBT AIRFLOW[®] Prophylaxis Master, handpiece cords and handpieces followed by a dry off with a soft non lint cloth.
- ⊘ DO NOT use paper towels as these can be abrasive.
- ⚠ Follow carefully the instructions provided by the wipes manufacturer.
- ⊘ DO NOT use Advantaclear wipes. It damages EMS products
- ⚠ Risk of contamination. Always disinfect the bottom and top areas of device air connections.
- ⚠ Please remember to dry your AIRFLOW[®] MAX and PIEZON[®] handpiece cord connections and ensure they are cooled and dried prior to each use. Excess moisture can cause blockages.
- ⚠ Follow the recommendations of the "Reprocessing Instructions" manual (FB-358/NA) regarding procedure for cleaning and sterilizing the components, and the present-day regulations on reprocessing in effect in your country.

5- MAINTENANCE

Scan for support



5.1- AIRFLOW® MAX/PERIOFLOW® MAX handpiece daily maintenance

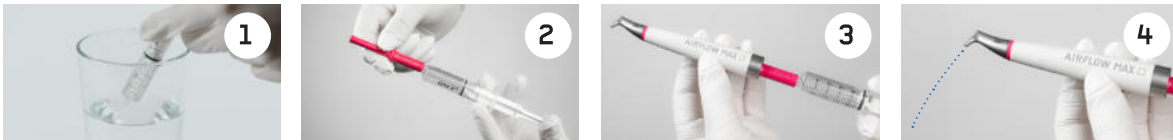
! Before the reprocessing or in case of a clogged AIRFLOW® MAX and PERIOFLOW® MAX handpieces: use Easy Clean provided in your AIRFLOW® Application box.

1- Clean the water channel:



Clear the water channel with air

2- Clean the air-powder channel with Easy Clean:

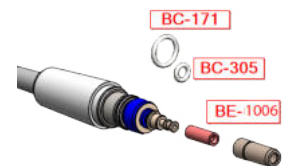


- ▶ Fill the syringe with distilled water
- ▶ Connect the Easy Clean
- ▶ Connect the handpiece
- ▶ Rinse the air-powder channel

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

5.2- AIRFLOW® MAX/PERIOFLOW® MAX handpiece leakage

In case of leakage between handpiece and handpiece cord connector, replace the cord gaskets with the spare parts provided in the EL-1211 kit located in the GBT Machine AIRFLOW® Prophylaxis Master Maintenance Kit. Contact EMS aftersales service in Southlake if needed.



5.3- PIEZON® handpiece leakage

In case of leakage between handpiece and handpiece cord connector please contact EMS aftersales service to replace the BC-1039 PIEZON® connector o-ring.

5.4- Light guide check & replace¹¹

The light guide loses its transparency after undergoing repeated reprocessing cycles. Check the transparency of it every month and proceed as follows:



1. Remove the PIEZON® Instrument and unscrew the handpiece nose cap by hand.
2. Take off the light guide and inspect it.
3. Place in a new light guide .
4. Screw the nose cap on again, by hand only.

¹¹ Only valid for PIEZON® handpiece LED

5.5- Wear



For PIEZON® instruments, check regularly instrument length and tip using the PIEZON® instrument check tool. Ultrasound instruments wear during use and become shorter. Worn instruments are ineffective and are a cause of discomfort to the patient. If PIEZON® instrument extremity reaches or is shorter than the limit indicated by the GBT logo, it can have excessive and uncontrolled vibration. Replace the PIEZON® instrument.

Scan for support



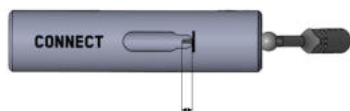
For the PIEZON® PI MAX instrument, check the tip length with the PI TOOL or the PIEZON® instrument check tool.

Remove the SCREW from the PI TOOL base.

Gently place the TIP in the hole marked CHECK up to STOP.

If the TIP sticks out, it can continue to be used.

If the TIP does not stick out = TIP too short. Replace the PIEZON® PI MAX instrument



! For precautionary reasons, do not exceed the validated lifespan of the components (see the "TECHNICAL DESCRIPTION" section).

Always use EMS original products. Using non-original components may damage the equipment, and practitioner or patient may be injured.

5.6- Handpiece cord system replacement

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

! Depressurize the powder chamber before disconnecting the AIRFLOW® MAX handpiece cord system.

In case of persistent malfunction or damage to the PIEZON® or AIRFLOW® MAX 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 system disconnecting procedure:

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

5.7- Monthly check

Each month check for cleanliness of both air and water filters.

! 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 filters cleanliness

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

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

Air filters usually remain cleaner for longer periods of time. Replace once a year. (The yearly maintenance service includes the replacement of the 3 filters.)




Good

Worn-out




To change the water and air filters from your device on a GBT Station or other support, remove the device from the support via the centre screw. Then follow the procedure below:

1.  Disconnect the power cord from the grid first.
2. Disconnect the central water and air hoses by pulling it off the connector.
3. Pull the filter off by hand or by using a small flat screwdriver. To remove the water bottle filter (3), press the clip (shown in the picture on the left) with your finger.
4. Replace with a new filter and reconnect the hose.

5.8- Preventive maintenance and repair

 EMS devices and products must only be maintained and/or repaired by EMS service center in Southlake.

 A preventive maintenance each 2000 hours (this is when the wrench symbol on your device is ON prompting you to contact your EMS service provider), 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.

The remote maintenance mode allows EMS aftersales service to use your device informations in order to resolve certain failures more quickly and easily. If the failure cannot be resolved remotely by the 1st level support, you will need to return your device.



When returning the device for service, it is recommended that you ship the device with its pedal, powder chamber, bottles and handpiece 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 section 6.4).

 Reprocess your device before returning it to EMS aftersales service in Southlake.

5.9- Pairing a (new) pedal

If bluetooth symbol is solid orange or if you want to pair a new pedal, follow the procedure below:



1. Place the two handpieces in their holders.
2. Press briefly left and right buttons simultaneously (3 seconds). A sonar sound will start playing (if not, repeat step 3).



3. Remove one battery from the pedal (no need to remove both). While the sonar sound plays, replace the battery into the wireless pedal.
4. Within a short time (less than 15 seconds), the pairing will be complete, the 3 white LEDs will blink for a while, and the bluetooth logo will switch to white and then disappear ; 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, turn the machine OFF, wait 10 seconds, then turn it ON again and redo the procedure from the beginning.

6- TROUBLESHOOTING

For each troubleshooting procedure of the section "6- Troubleshooting", follow the steps one by one until your problem is solved. Once you have tried all the steps and if the problem persists, please contact your local EMS aftersales service/dealer.

6.1- For AIRFLOW®/PERIOFLOW® products

No powder/air jet coming from the handpiece

- 1° Depressurize/repressurize the powder chamber
- 2° If applicable, change the PERIOFLOW® nozzle
- 3° Depressurize the powder chamber and unclog AIRFLOW® MAX/PERIOFLOW® MAX handpiece, as described in the section "5.1- AIRFLOW® MAX/PERIOFLOW® MAX handpiece daily maintenance" and clean and dry the handpiece cord connector

No water spray is coming from the PERIOFLOW® nozzle

- 1° Check that water setting is on 3 and put power setting to 3
- 2° Change the PERIOFLOW® nozzle
- 3° Unclog PERIOFLOW® MAX handpiece, as described in the section "5.1- AIRFLOW® MAX/PERIOFLOW® MAX handpiece daily maintenance" and clean and dry the handpiece cord connector


Water leakage between the handpiece and the PERIOFLOW® nozzle (unusual leak of droplets)

- 1° Check for powder residue between the PERIOFLOW® nozzle and the handpiece
- 2° Check the alignment/position of the PERIOFLOW® nozzle on the handpiece
- 3° Change the PERIOFLOW® nozzle

Water leakage between the AIRFLOW® MAX/PERIOFLOW® MAX handpiece and the handpiece cord.

- 1° Check that the handpiece is correctly connected to the handpiece cord
- 2° Unclog AIRFLOW® MAX/PERIOFLOW® MAX handpiece, as described in the section "5.1- AIRFLOW® MAX/PERIOFLOW® MAX handpiece daily maintenance" and clean and dry the handpiece cord connector
- 3° Replace the AIRFLOW® cord gaskets as described in the section "5.2- AIRFLOW® MAX/PERIOFLOW® MAX handpiece leakage"

Insufficient or no water from AIRFLOW® MAX/PERIOFLOW® MAX handpiece

- 1° Check that your water setting is on 3
- 2° If applicable, change the PERIOFLOW® nozzle
- 3° Unclog AIRFLOW® MAX/PERIOFLOW® MAX handpiece, as described in the section "5.1- AIRFLOW® MAX/PERIOFLOW® MAX handpiece daily maintenance" and clean and dry the handpiece cord connector
- 4° Check your water filter cleanliness and replace it if necessary, as described in the section "5.7- Monthly check"
- 5°  Disconnect the mains plug before servicing any filter
- 5° Make sure you have correctly connected your water supply and with sufficient pressure

The unit efficiency decreases

- 1° Depressurize the powder chamber
- 2° Check the powder chamber level, as described in section 3, "Filling the powder chambers"
- 3° Check the air pressure of the dental unit
- 4° If applicable, change the PERIOFLOW® nozzle
- 5° Unclog AIRFLOW® MAX/PERIOFLOW® MAX handpiece, as described in the section "5.1- AIRFLOW® MAX/PERIOFLOW® MAX handpiece daily maintenance" and clean and dry the handpiece cord connector

If the handpiece falls

- | | |
|---|---|
| AIRFLOW® MAX handpiece: | PERIOFLOW® MAX handpiece: |
| 1° Check the state of the nozzle | 1° Check you can replace the PERIOFLOW® nozzle |
| 2° Check if the handpiece is working properly | 2° Check the handpiece is still working properly |
| 3° Replace the handpiece by a new/another handpiece | 3° Replace the handpiece by a new/another handpiece |

6.2- For PIEZON® products

The PIEZON® handpiece heats up

- 1° Check that power setting is on 2 and water setting is on 3
- 2° Take off the handpiece from the handpiece cord connector and wait 5 minutes for it to cool

White LED PIEZON® is not working¹²

- 1° Clean and dry the PIEZON® handpiece and the handpiece cord connector
- 2° Put back the PIEZON® handpiece into the holder and wait 1 minute

Insufficient LED PIEZON® lighting⁶

- 1° Replace the light guide
- 2° Replace the handpiece by a new/another handpiece

Water leakage between the PIEZON® handpiece and the handpiece cord

- 1° Check that the handpiece is correctly connected to the handpiece cord
- 2° Clean and dry the PIEZON® handpiece and the handpiece cord connector
- 3° Replace the PIEZON® connector o-ring as described in the section "5.3- PIEZON® handpiece leakage"

¹² Only valid for PIEZON® handpiece LED

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

- 1° Make sure that the PIEZON® instrument is correctly screwed on (use the CombiTorque® tool)
- 2° Check the wear of the PIEZON® instrument with the check tool (see section "5.5- Wear"), and replace it if necessary
- 3° Clean and dry the PIEZON® handpiece and the handpiece cord connector

No or low water flow

- 1° Check that your water setting is on 3
- 2° Blow compressed air through the PIEZON® instrument
- 3° Replace the PIEZON® instrument
- 4° Clean and dry the PIEZON® handpiece and the handpiece cord connector
- 5° Check your water filter cleanliness and replace it if necessary, as described in the section "5.7- Monthly check"

 Disconnect the mains plug before servicing any filter

- 6° Make sure you have correctly connected your water supply and with sufficient pressure

Suspicious noise

- 1° Check the correct screwing with the CombiTorque®
- 2° Check the wear of the PIEZON® instrument with the check tool (see section "5.5- Wear"), and replace it if necessary

If the handpiece falls

- 1° Check the state of the PIEZON® instrument, and replace it if necessary
- 2° Check if the handpiece is working properly
- 3° Replace the handpiece by a new/another handpiece


6.3- For the device

6.3.1- Device troubleshooting


The device is whistling or making strange noises

 Risk of bottle explosion



- 1°  Disconnect the mains plug and stop using your device immediately
- 2° Check the water bottle for crack or any damage and, if the case, replace it with a new one
- 3° Check the supplied air pressure: it must be between 4.5 bar and 7 bar
- 4° If 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

The device is making smoke (and fire)

 Risk of fire and electric shock




- 1°  Disconnect the mains plug and stop using your device immediately

Handpiece cord or device leakage

 Risk of fire and electric shock



- 1°  Disconnect the mains plug
- 2° If the leak come from the AIRFLOW® handpiece, go to "6.1- For AIRFLOW®/PERIOFLOW® products", if the leak come from the PIEZON® handpiece, go to "6.2- For PIEZON® products"
- 3° If the leak come from AIRFLOW® MAX/PIEZON® handpiece cord system, disconnect it from the device and clean the male and female connectors of the handpiece cord system

Water filter leakage

- 1° Check your water filter cleanliness and replace it, if necessary, as described in section "5.7- Monthly check"



 Disconnect the mains plug before servicing any filter

Bottle/bottle connection leakage

- 1° Ensure the bottle cap has been correctly closed
- 2° Clean both connections: cap and device sides
- 3° Replace the bottle




Cleaning liquid remaining after water lines rinsing

- 1° Make sure your water setting is on 3
- 2° Make sure you have correctly connected your water supply and with sufficient pressure
- 3° Perform a second rinsing phase before treatment



The unit does not start

- 1° Check the power grid is correctly connected
- 2°  Disconnect the mains plug and remove the power cord from the device
- 3° With the help of a small flat screwdriver, open the fuse-holder cover
- 4° Replace fuses only with the exact type required, as described in section "TECHNICAL DESCRIPTION"





Wireless pedal does not work

- 1° Release the pedal and wait for 1 minute
- 2° Disconnect and reconnect both PIEZON® and AIRFLOW® MAX handpiece cord systems, as described in the section "2.8- Check the handpiece cord system connections"
- 3° Pair the wireless pedal, as described in section "5.9- Pairing a (new) pedal"
- 4° Replace both batteries from the wireless pedal with new high-quality AA lithium batteries



No pressurization of the powder chamber

- 1° Check that your device is ON: GBT symbol is lighted (white, or pink for GBT certified practitioners)
- 2° Check that AIRFLOW® MAX handpiece cord system symbol is OFF and that it is well connected, as described in the section "2.8- Check the handpiece cord system connections"

Powder chamber white light is BLINKING during pressurization attempt

- 1° Make sure you have correctly connected your air supply and with sufficient pressure
- 2° Check your air filter cleanliness and replace it if necessary, as described in section "5.7- Monthly check"
- ⚡ Disconnect the mains plug before servicing any filter

Powder chamber white light is BLINKING at depressurization

- 1° Unclog AIRFLOW® MAX/PERIOFLOW® MAX handpiece, as described in the section "5.1- AIRFLOW® MAX/PERIOFLOW® MAX handpiece daily maintenance" and clean and dry the handpiece cord connector

Powder sprays out of chamber at depressurization

- 1° Check the powder chamber level, as described in section 3 "Filling the powder chambers"

Powder leaks under the AIRFLOW® MAX handpiece cord system

- 1° Switch OFF the device and disconnect AIRFLOW® MAX handpiece cord system, as described in the section "2.8- Check the handpiece cord system connections"
- 2° Clean and dry the male and female electrical contacts (mini-jack) of the AIRFLOW® MAX handpiece cord system
- 3° Replace your AIRFLOW® MAX handpiece cord with a new one



Powder chamber is leaking

- 1° Clean and dry the powder chamber bottom and cap, including the o-ring, as well as the connection points on the caliper
- 2° Replace the powder chamber



6.3.2- Symbols troubleshooting

REMINDER: For each troubleshooting procedure of the section "6- Troubleshooting", follow the steps one by one until your problem is solved. Once you have tried all the steps and if the problem persists, please contact your local EMS aftersales service/dealer.



Wrench symbol is ON

⚠ Maintenance indicator. It is time to send your device to yearly maintenance service. Quickly contact EMS aftersales service in Southlake.



Wrench symbol is BLINKING

⚠ Permanent or transitory hardware fault condition detected

⚠ Risk of fire and electric shock

1° ⚡ First disconnect the mains plug and stop using your device immediately

2° Wait for 30 seconds, then plug it back again and restart the device

3° ⚡ Disconnect again the mains plug and wait for 1 hour, then plug it back again and restart the device



Battery symbol is ON

1° Replace both batteries from the wireless pedal with new high-quality AA lithium batteries



Bluetooth symbol is ON

1° Pair the wireless pedal, as described in section "5- Maintenance" --> "5.9- Pairing a (new) pedal"



AIRFLOW® MAX handpiece cord system symbol is ON

1° Switch OFF the device and disconnect AIRFLOW® MAX handpiece cord system, as described in the section "2.8- Check the handpiece cord system connections"

2° Clean and dry the male and female electrical contacts (mini-jack) of the AIRFLOW® MAX handpiece cord system



PIEZON® handpiece cord system symbol is ON

- 1° Switch OFF the device and disconnect PIEZON® handpiece cord system, as described in the section "2.8- Check the handpiece cord system connections"
- 2° Clean and dry the male and female electrical contacts (mini-jack) of the PIEZON® handpiece cord system



AIRFLOW® MAX/PERIOFLOW® MAX handpiece symbol is ON

- 1° Check that AIRFLOW® MAX/PERIOFLOW® MAX handpiece contains the RFID/NFC logo.
- 2° If it is the case, check that this handpiece is correctly connected to the handpiece cord connector.
- 3° If the logo RFID/NFC is missing, you can use your EMS handpiece but you will be restricted in connectivity. Contact your EMS representative for further assistance and information.



PIEZON® handpiece symbol is ON

- 1° Check that PIEZON® handpiece contains the RFID/NFC logo.
- 2° If it is the case, check that this handpiece is correctly connected to the handpiece cord connector.
- 3° If the logo RFID/NFC is missing, you can use your EMS handpiece but you will be restricted in connectivity. Contact your EMS representative for further assistance and information.

Wifi symbol is ON

(when switching ON the device)



- 1° To connect your GBT Machine AIRFLOW® Prophylaxis Master to the wifi, press simultaneously the buttons 2 and 3, as described in the section "2.5.1- Connecting / Disconnecting from the Wi-Fi"
 - 2° Open your EMS interface to select the right network and enter the password
- Please refer to the the section "2.5.1- Connecting / Disconnecting from the Wi-Fi" for more information.
 NB: If you want to skip this step, touch the ON/OFF button on the caliper

Upgrade software symbol is ON

(when switching ON the device)



- To launch the software upgrade, press simultaneously the buttons 2 and 3 during 2 seconds.
- The maintenance key symbol will light up and fade several times. This step may take between 10 seconds and 3 minutes, depending on the update performed.
 NB: If you want to skip this step, touch the ON/OFF button on the caliper. The next time you turn ON your device, this update will be requested again.

Scan for support



6.4- Contact EMS aftersales service

ELECTRO MEDICAL SYSTEMS L.L.C.
 2150E Continental Boulevard
 Southlake, Texas, 76092, US
 T. +1 972 690 8382
 E-mail: custsvc@ems-na.com
 Aftersales: emsrepairs@ems-na.com

6.5- 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: vigilancemailbox@ems-ch.com

By fax: +41 (0) 22 99 44 701

By post: E.M.S. Electro Medical Systems S.A., Chemin de la Vuarpillière 31, 1260 Nyon – Switzerland



SUSTAINABILITY

1- DISPOSAL OF WASTE



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.



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

2- SUSTAINABLE DESIGN



The device, on a voluntary basis, respects the latest Eco design low energy standby and off mode consumption regulation¹³. Packaging cardboards are recycled and recyclable.



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

WARRANTY

The warranty is valid if you complete maintenance 2000 hours after purchase and then 2000 hours after each last maintenance. 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.

TECHNICAL DATA COLLECTION AND PRIVACY POLICY

During maintenance and/or repair of the device, EMS or any authorized EMS repair center will have access to certain technical information such as usage statistics (hereinafter "Technical Data"), collected during the device service.

Such technical data shall be analyzed and used by EMS in its legitimate interest, e.g. to carry out statistical analysis and to improve its customer service and/or its Research and Development processes.

EMS may also use such technical data along with your personal details in order to be able to understand your personal usage of the device and offer you a better customer experience and tailored service. However, you can unsubscribe from this process at any time, by simply sending us an email at privacy@ems-ch.com.

Rest assured that these activities will be carried out in compliance with applicable data protection laws. For any questions regarding your personal data, please consult our privacy policy at www.ems-company.com or send an email to privacy@ems-ch.com.




























¹³ 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.

TECHNICAL DESCRIPTION

Manufacturer	E.M.S. Electro Medical Systems S.A., Chemin de la Vuarpillière 31, 1260 Nyon, Switzerland																																											
Model	GBT Machine AIRFLOW® Prophylaxis Master, product code FT-300																																											
Classification IEC 60601-1	Electrical Insulation Class-I Applied part Type B IP21 Control unit IP21 Foot pedal																																											
Essential Performance	This medical device has no Essential Performance in the meaning of the IEC 60601-1 :2020 - The device has no function whose absence or degradation would result in an unacceptable risk.																																											
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	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)																																											
Filters	Water filter: 50 micrometers Air filter: 14 micrometers																																											
Wireless communication module	<table border="1"> <thead> <tr> <th></th> <th>Radio Interface n°1</th> <th>Radio Interface n°2</th> <th>Radio Interface n°3</th> <th>Radio Interface n°4</th> </tr> </thead> <tbody> <tr> <td>Type of Radio:</td> <td>WIFI</td> <td>LTE¹⁴</td> <td>RFID</td> <td>Bluetooth Low Energy</td> </tr> <tr> <td>Reference of Radio chipset-module:</td> <td>ESP32-S3-WROOM-1</td> <td>ME310G1-WW</td> <td>STR25R3911B-AQTF</td> <td>BGM220SC22WGA2R</td> </tr> <tr> <td>Frequency Bands:</td> <td>2412 to 2484 MHz</td> <td>B1, B3, B8, B20, B28</td> <td>13.56 MHz</td> <td>2402 MHz to 2480 MHz</td> </tr> <tr> <td>Voltage Range:</td> <td>3.0 to 3.6 V</td> <td>3.2 to 4.2 V</td> <td>2.4 to 5.5 V</td> <td>1.8 to 3.8 V</td> </tr> <tr> <td>Temperature Range:</td> <td>-40 to +85°C</td> <td>-40 to +85°C</td> <td>-40 to +125°C</td> <td>-40 to +85°C</td> </tr> <tr> <td>Antenna:</td> <td>PCB Gain 3.26 dBi</td> <td>Embedded Gain 5 dBi</td> <td>0 dBi</td> <td>Integral antenna Gain 2.3 dBi</td> </tr> <tr> <td>Maximum Output Power:</td> <td>< 20 dBm</td> <td>< 23 dBm</td> <td>< 42 dBµA/m</td> <td>< 10 dBm</td> </tr> </tbody> </table>					Radio Interface n°1	Radio Interface n°2	Radio Interface n°3	Radio Interface n°4	Type of Radio:	WIFI	LTE ¹⁴	RFID	Bluetooth Low Energy	Reference of Radio chipset-module:	ESP32-S3-WROOM-1	ME310G1-WW	STR25R3911B-AQTF	BGM220SC22WGA2R	Frequency Bands:	2412 to 2484 MHz	B1, B3, B8, B20, B28	13.56 MHz	2402 MHz to 2480 MHz	Voltage Range:	3.0 to 3.6 V	3.2 to 4.2 V	2.4 to 5.5 V	1.8 to 3.8 V	Temperature Range:	-40 to +85°C	-40 to +85°C	-40 to +125°C	-40 to +85°C	Antenna:	PCB Gain 3.26 dBi	Embedded Gain 5 dBi	0 dBi	Integral antenna Gain 2.3 dBi	Maximum Output Power:	< 20 dBm	< 23 dBm	< 42 dBµA/m	< 10 dBm
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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 3000m Pressure: 700hPa to 1060hPa Under certain operating conditions, applied parts may exceed 41°C of temperature and reach a maximum temperature of 48 °C.																																											
Storage conditions	<p>Device only (with no water inside): Temperature: 5°C to 40°C Humidity: up to 85% Pressure: 500hPa to 1060hPa Accessories: see individual packaging</p> <p>Device with accessories, in packaging: Temperature: 5°C to 25°C Humidity: up to 85% Pressure: 500hPa to 1060hPa</p>																																											
Transport conditions	<p>Device only (with no water inside): Temperature: -29°C to 38°C Humidity: up to 85% Pressure: 500hPa to 1060hPa Accessories: see individual packaging</p> <p>Device with accessories, in packaging: Temperature: -20°C to 38°C Humidity: up to 85% Pressure: 500hPa to 1060hPa</p>																																											

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. Note: The pH and particle size information are not relevant. Air: pressure 4.5-7bar, dry-only (humidity 1.032g/m3 max.), oil filtered 0.1mg/m3 max., minimum flow-rate 20 l/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	PIEZON® and CLEANER bottles: 5 years
Validated lifetime	Handpieces (main bodies): 1000 sterilization cycles PI MAX instrument Tip: 20 sterilization cycles PI MAX instrument Holder: 1000 sterilization cycles Instrument PI: 100 sterilization cycles Others instruments and their CombiTorque®: 1000 sterilization cycles
Lifespan	PIEZON® instruments: Check PIEZON® instrument length and tip thread using the PIEZON® instrument check tool. PIEZON® PI MAX instrument: Use the PI MAX instrument Tool or the PIEZON® instrument check tool. The tip mustn't be shorter than 3mm.
Expected service life	Device: 7 years, having regular recommended maintenance (every 2000 hours) Handpiece cord systems : 2 years

SYMBOLS


	General Warning
	Warning Electricity
	Non-ionizing radiation (radio communication)
	Read the operation instructions
	Device requiring protective earth
	Disconnect the mains plug for purposes of maintenance and in case of malfunction
	Electronic instructions for use
	Mandatory action
	Wear personal protective equipment
	Expiration date
	Single use. Do not re-use.
	Do not do.
	Disposal of old electronic equipment (European Union & other countries with separate collection systems)
	Sterilizable at up to 135°C in the autoclave
	Thermal disinfection
	Input
	Output
	Fuse
IP ...	Protection against water permeability
	Applied part, type B
	Manufacturer
	EU Authorized Representative
	Manufacturing date
	Serial number
	Catalog number / Product reference
	Medical device
	Medical Device compliant with EU Regulation 2017/745 Number of the Notified Body
	CAUTION: Federal (US) law restricts this device to sale by or on the order of a physician

ELECTROMAGNETIC COMPATIBILITY

This device has been designed and tested for compliance with IEC 60601-1 and IEC 60601-4-2 requirements for electromagnetic immunity and emissions. Proper installation and use according to these instructions are essential to maintain Basic Safety.

1- INTENDED ELECTROMAGNETIC ENVIRONMENT

The GBT Machine AIRFLOW® Prophylaxis Master is intended for use in dental offices with typical mains power quality and controlled electromagnetic conditions.

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

The use of parts other than those supplied or listed as accessory may negatively affect EMC performance.

The customer or the user of this product should assure that it is used in such an environment.

2- DEVICE PERFORMANCE AND EMC DISTURBANCE EFFECTS

This device has no Essential Performance as defined by IEC 60601-1:2020. Basic Safety is maintained even if electromagnetic disturbances occur. However, EMC interference may temporarily affect normal operation.

If electromagnetic interference occurs, the operator may observe:

- Interruption or irregularity in powder spray or water flow
- Reduced ultrasonic power or unexpected variation in scaling performance
- Display malfunctions or unexpected shutdown
- Delayed response to pedal activation

3- CLINICAL IMPACTS

These effects may lead to treatment interruption or delay but do not create an unacceptable risk to patient safety.


4- OPERATOR ACTIONS

- Stop treatment immediately.
- Verify cabling and connections.
- Move portable RF communications equipment (e.g., mobile phones, Wi-Fi routers) at least 30 cm (12 inches) away from the device.
- If interference persists, disconnect the device and contact EMS technical support.

5- COMPLIANCE LEVELS

The device was tested according to IEC 60601-4-2 and related standards. Immunity and emissions compliance levels are summarized below:

Immunity test	IEC 60601 test level	Compliance level	Electromagnetic environment - guidance
Electrostatic discharge (ESD) IEC 61000-4-2	± 8 kV contact ± 15 kV 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 lines 100 kHz repetition frequency ± 1 kV for input/output lines 100 kHz repetition frequency		Mains power quality should be that of a typical commercial or hospital environment.
Surge IEC 61000-4-5	± 1 kV line(s) to line(s) ± 2 kV line(s) to earth		Mains power quality should be that of a typical commercial or hospital environment.
Voltage dips IEC 61000-4-11	70 % UT (30 % dip in UT) for 25/30 cycles 0 % UT for 0,5 cycle at 0°, 45°, 90°, 135°, 180°, 225°, 270° and 315° 0 % UT for 1 cycle 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	0% UT for 5 s 0% UT for 250/300 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 150 kHz to 80 MHz 6V in ISM bands 150kHz and 80 MHz 80 % AM at 1 kHz	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 GBT Machine AIRFLOW® Prophylaxis Master, including its cables. Otherwise, degradation of the performance of this equipment could result. Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey ¹⁵ , should be less than the compliance level in each frequency range ¹⁶ . Interference may occur in the vicinity of equipment marked with the following symbol:  or 
Radiated RF IEC 61000-4-3	3 V/m 80 MHz to 6 GHz 80% AM at 1 kHz	3 V/m 80 MHz to 6 GHz 80% AM at 1 kHz	
Proximity fields from RF wireless communications equipment IEC 61000-4-3	See Table below		
IEC 61000-4-39	134.2 kHz / Modulation pulsée 2.1 kHz / 65A/m 13.56 MHz / Modulation pulsée 50 kHz / 7.5A/m	134.2 kHz / Modulation pulsée 2.1 kHz / 65A/m 13.56 MHz / Modulation pulsée 50 kHz / 7.5A/m	

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.

6- PROXIMITY FIELDS FROM RF WIRELESS COMMUNICATIONS EQUIPMENT

IEC 61000-4-3

Test Frequency (MHz)	Modulation	Required level (V/m)	Immunity test level (V/m)
385	Pulse Modulation 18Hz	27	27
450	Pulse Modulation 18Hz	28	28
710 745 780	Pulse Modulation 217 Hz	9	9
810 870 930	Pulse Modulation 18 Hz	28	28
1720 1845 1970	Pulse Modulation 217 Hz	28	28
2450	Pulse Modulation 217 Hz	28	28
5240 5500 5785	Pulse Modulation 217 Hz	9	9

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.

7- 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.

¹⁵ 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.

¹⁶ Over the frequency range 150 kHz to 80 MHz, field strengths should be less than 3 V/m.

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 network that supplies buildings used for domestic purposes.
Harmonic emissions IEC 61000-3-2	Class A	
Voltage fluctuations / flicker emissions IEC 61000-3-3	Complies	

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) , but not limited to the following standards and/or normative documents:

- EMC
- ETSI EN 301 489-1 V2.2.3
- ETSI EN 301 489-17 v3.2.4
- ETSI EN 301 489-3 v2.3.2
- ETSI EN 301 489-52 v1.2.1
- ETSI EN 300 330 v2.1.1
- EN 301 908-1 v15.1.1
- EN 301 908-13 v13.2.1
- SPECTRUM
- ETSI EN 300 328 v2.2.2

1- FCC STATEMENTS

Any changes or modifications not expressly approved by Electro Medical Systems for compliance could void the user's authority to operate this equipment.

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 circuit different from that to which the receiver is connected.
- Consult the dealer or an experienced radio/TV technician for help

1.1- RF Exposure mobile Device

This device complies with FCC and IC radiation exposure limits set forth for general population. This device must be installed to provide a separation distance of at least 20cm from all persons and must not be co-located or operating in conjunction with any other antenna or transmitter.

This device complies with Part 15 of the FCC Rules. Operation is subject to the following two conditions: (1) this device may not cause harmful interference, and (2) this device must accept any interference received, including interference that may cause undesired operation.

1.2- RF Exposure portable device only for RFID

This device complies with FCC radiation exposure limits set forth for general population. This device must not be co-located or operating in conjunction with any other antenna or transmitter.

2- ISED STATEMENTS

This device contains licence-exempt transmitter(s)/receiver(s) that comply with Innovation, Science and Economic Development Canada's licence-exempt RSS(s). Operation is subject to the following two conditions: (1) this device may not cause interference, and (2) This device must accept any interference, including interference that may cause undesired operation of the device.

2.1- RF Exposure mobile Device

This device complies with ISED radiation exposure limits set forth for general population. This device must be installed to provide a separation distance of at least 20cm from all persons and must not be co-located or operating in conjunction with any other antenna or transmitter.

2.2- RF Exposure portable device only for RFID

This device complies with ISED radiation exposure limits set forth for general population. This device must not be co-located or operating in conjunction with any other antenna or transmitter.

3 - WIRELESS COMMUNICATION MODULE, ONLY FOR COSTA RICA

	Radio Interface n°1	Radio Interface n°2	Radio Interface n°4
Type of Radio:	WIFI	RFID	Bluetooth Low Energy
Reference of Radio chipset-module:	ESP32-S3-WROOM-1	STR25R3911B-AQTF	BGM220SC22WGA2R
Frequency Bands:	2412 to 2484 MHz	13.56 MHz	2402 MHz to 2480 MHz
Voltage Range:	3.0 to 3.6 V	2.4 to 5.5 V	1.8 to 3.8 V
Temperature Range:	-40 to + 85°C	-40 to + 125°C	-40 to + 85°C
Antenna:	PCB Gain 3.26 dBi	0 dBi	Integral antenna Gain 2.3 dBi
Maximum Output Power:	17.17 dBm	-4.43 dB μ A/m	9.97 dBm

► EMS powders are much less abrasive than traditional mechanical debridement techniques such as rubber cups and handscaling.

We recommend to use AIRFLOW® and PERIOFLOW® first, followed by PIEZON®. GBT: the minimally invasive way for Professional Mechanical Plaque Removal (PMPR).

► As part of the GBT disclosing phase, coloration also minimizes AIRFLOW® abrasion. No more color signifies that all biofilm has been removed: the objective of AIRFLOW® is achieved, to preserve the integrity of dental tissues.

⚠ EMS devices have been designed and tested for exclusive use with EMS prophylaxis powders.

! ONLY USE EMS POWDERS



So-called "EMS compatible" powders in the marketplace may cause damage to oral tissues and the device.

See below a damaged AIRFLOW® nozzle after use with a "compatible" powder. If EMS products are damaged by using "compatible" powders the product Warranty will be lost.



Damaged
handpiece



ONLY USE THE SWISS ORIGINAL EMS INSTRUMENTS!

The EMS Instrument, the Handpiece and the electronic module have to vibrate in harmony like in a trilogy.

Using so-called "compatible" and copy tips from third parties can damage your patients' teeth as well as EMS handpieces.

They were developed and made for each other by EMS.

One PS Instrument is designed for up to 1000 treatments, thus reducing the cost of one treatment to only a few cents.

Therefore, there is absolutely no reason to "save" money by using so-called "compatible" tips. Moreover, if the Piezon Handpiece gets damaged (the thread) it will lose its warranty so at the end of the day, "compatible" tips will always cost you more.



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