



MANUAL PRESENTATION

Technical Name: Equipment for Dental Prophylaxis Sodium Bicarbonate/Ultrasound Trade Name: Equipment for Prophylaxis by Ultrasound / Bicarbonate Jet Models: Profi Neo / Profi Neo LED / Profi Class / Profi Class LED Brand: Dabi Atlante

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77000001308 - Rev.: 11 - May/22

Document originally written in Portuguese.

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GENERAL INFORMATION

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1. GENERAL INFORMATION

1.1.DEAR CUSTOMER

Congratulations on your excellent choice. When you buy ALLIAGE quality equipment, you can be sure of purchasing products with technology compatible with the best in the world in their class. This manual provides you with a general presentation of your equipment, describing important details that can guide you in its correct use, as well as in solving small problems that may eventually occur. This manual must be read in full and kept for future reference.

1.2.INDICATIONS FOR USE

The Ultrasound / Bicarbonate Jet Prophylaxis Equipment is intended to assist in dental treatments such as removing plaque and residual stains; tartar removal; periodontal treatment; endodontic treatment; micro retro surgery; cavity preparations for restorations; amalgam condensation, inlays and on lays and gutta percha; removal of pins and crowns, among other procedures related to dental treatments.

1.3.CONTRAINDICATIONS

This equipment is contraindicated for use in patients who have serious respiratory, renal or hemodialysis changes, these cases must have medical monitoring. We recommend the use of a mask and glasses to apply the bicarbonate jet.

1.4.SYMBOLOGY

The following symbols are used both throughout this manual and on the product. Make sure you fully understand each symbol and follow the instructions that come with it.



REF	Catalog number	135℃ ∭	Sterilizable in a steam sterilizer (autoclave) at specified temperature
ҟ	Type B applied parts	X	Indicates that the product must be taken to a special waste collection site at the end of its useful life. Applies to both the device and accessories
\triangle	Attention		Electrostatic sensitive devices (ESD)
	Protective ground wire		General warning
~	Alternating current	SN	Serial number
0	Turn Off (Power: Disconnects from the main switch)		Mandatory action
Ι	Turn On (Power: Connects from the main switch)		Follow the instructions for use

Profi Line

***	Manufacturer	\sim	Manufacturing date
MODEL	Model	~	Periodontal ultrasound
	Bicarbonate jet	<u>+</u> -■	Endo ultrasound
\bigcirc	Peristaltic pump	-	Power down and language selection
+	Power increase and language selection	1	Access to programming memory 1
2	Access to programming memory 2	3	Access to programming memory 3
4	Access to programming memory 4	Φ	Start operation key



Choosing a Dental Tip

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WARNINGS, CAUTIONS AND RECOMMENDATIONS

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2. WARNINGS, CAUTIONS AND RECOMMENDATIONS

General warnings

	Read and understand all instructions contained in these instructions for use before installing or operating this equipment.
	Use only the equipment in perfect condition and protect yourself, patients and others against possible dangers.
\triangle	The Ultrasound / Bicarbonate Jet Prophylaxis Equipment has 4 different interactions with the user, they are: - Identification label: Located on the side of the equipment; - Safety symbols: Located at risk locations and on their identification tag; - Central panel; - Pedal.

During transportation

The equipment must be transported and stored, observing the following:

- Handle with care to avoid falls, excessive vibrations and impacts;
- The arrows on the packaging must be pointing upwards;
- To handle the package as a single unit, consider the center of gravity indicator;
- Do not stack above the quantity indicated on the packaging;
- Do not walk or stand above the package
- Protect against sunlight, moisture, water and dust;
- Observe the temperature, pressure and relative humidity limits.

During the installation of the equipment

The installation procedure must be carried out by an authorized technician. Instructions for installing the equipment are found in this manual.
To avoid the risk of electric shock, this equipment must only be connected to a mains supply with a protective ground.



Before turning on the equipment, make sure that it is connected to the correct voltage.

• The equipment must be installed only by authorized technical assistants.

• The service manual's recommendations regarding the mandatory existence of protective earth must be followed.

• The recommendations in the service manual regarding the mandatory existence of a protective circuit breaker must be followed.

• Install the equipment in a place where it will not be in contact with moisture, water, plants and animals.

• Install the equipment in a location where it will not be damaged by pressure, temperature, humidity, direct sunlight, dust, salts or corrosive products.

 \bullet The equipment must be correctly attached according to the service manual and must not be subjected to an inclination greater than 5 °. Risk of tipping.

• This equipment is not designed for use in the presence of vapors from flammable anesthetic mixtures or nitrous oxide.

• Place any other external devices at least 1.5 meters away from the equipment, so that the patient cannot touch any other external devices while they are being attended to.

• The recommendations in this manual for EMC should be followed. Communications equipment and RF generating sources can affect the operation of the equipment.

• Equipment may cause radio interference or interrupt the operation of nearby equipment, making it necessary to take mitigating measures, such as reorientation, relocation of equipment or shielding the location.

Before using the equipment

To help ensure proper hygiene and protect against infectious diseases, before using for the first time, the equipment must be cleaned and disinfected according to the instructions contained in this manual.

While using the equipment

- Under no circumstances can the patient operate the equipment.
- The patient must not t ouch other parts than those specific t o be treated.
- The equipment must be oper ated only by qualified health pr ofessionals.
- To operate the equipment, operating personnel must:
- Read and understand the user manual
- Be familiar with the basic structure and functions of this equipment.
- Be familiar with the emergency situation protocols for this equipment.

- Be able to recognize irregularities in the operation of the equipment and implement the appropriate measures, when necessary.

• The equipment has been designed according to the electromagnetic compatibility standards, but in very extreme conditions, it can cause interference with other equipment. Do not use this equipment in conjunction with other devices that are extremely sensitive to interference or with devices that create high electromagnetic disturbances.

• Do not position the patient on the equipment while starting it, as the patient may be injured if the equipment does not work properly. In the event of an error that requires turning the equipment off and on, remove the patient before turning it on again.

• In case of risk to the patient, press the emergency button immediately located on the side of the equipment.

• If this product is exposed to water, moisture or foreign substances, turn it off immediately and contact an Alliage Authorized Service Center.

• In case of damage or defect, do not use the equipment and contact an Authorized Alliage Service Center.

• Do not use the equipment if any of its compartments or parts are damaged, loose or have been removed. Contact an Alliage Authorized Service Center and request repair or replacement of any

Profi Line

damaged, loose or removed enclosures or parts of the equipment before using the equipment again. • Do not touch the equipment or use it if it is being repaired or if the equipment's cabinets have been removed.

• Do not open or remove any of the equipment's cabinets. No internal parts can be repaired by the user.

• In case of falling or impact of moving parts causing it to break, be careful when handling them, there may be sharp parts.

- The operator cannot come into contact with the patient when in contact with accessible connectors.
- The operator cannot use tools to open the equipment.

When using the product, the dental tip can reach the normal use temperature of 114.2 ° C. That if you are in contact for more than 1 min there may be a risk of slight superficial burns or irritation.
If there are obstructions or blockages in the cooling system, the tip can reach a maximum rate of temperature increase of 63% (179.7 ° C). That if you are in contact for more than 1 min there may be a risk of slight superficial burns or irritation.

Cross contamination prevention



Adequate cleaning and disinfection / sterilization measures should be taken to avoid cross-contamination between patients, users and others.

• For each new patient, perform cleaning, disinfection / sterilization procedures and according to the instructions contained in this manual.

After using / operating the equipment

- Turn off the equipment if it is not in use for a long time
- All parts that have been in contact with the patient must be cleaned and disinfected / sterilized with each new patient to prevent the transmission of infectious agents that can cause serious illness.
- Perform cleaning and disinfection / sterilization according to instructions contained in this manual.
- Do not unplug the cable or other connections unnecessarily.
- Do not modify any part of the equipment.

Precautions in case of alteration of the equipment operation

If the equipment shows any abnormality, check if the problem is listed in any item listed in the "Problem diagnosis" topic of this user manual.

If it is not possible to solve the problem, turn off the equipment, contact an Alliage Authorized Technical Assistance.



Precautions for reducing environmental impact

Alliage S / A aims to achieve an environmental policy to promote the supply of environmentally conscious medical and dental products that continually minimize environmental impact and are more friendly to the environment and human health.

To maintain a minimal impact on the environment, observe the recommendations below:

• After installation, send recyclable materials for recycling process.

• During the life cycle of the equipment, turn it off when it is not in use.

• To prevent environmental contamination, the disposal of waste and consumables must follow the normal procedure for biomedical waste.

Biomedical waste includes non-acute materials that may cause disease or suspicion of harboring pathogenic organisms that must be stored in a yellow bag properly labeled with a biohazard symbol, stored in a puncture-resistant, watertight container until collection and incineration.



The equipment packaging consists of cardboard and polyethylene that are 100% recyclable materials.

DIMENSIONS:

380 x 380 x 270mm /MASS: Approximately: 4,2 Kg

Precautions in case of unusable equipment

To avoid environmental contamination or improper use of the equipment, when it is unusable, it must be disposed of (in accordance with current legislation) in an appropriate place, as the materials inside can contaminate the environment.

For the European Economic Area (EEA), this product is subject to Directive 2012/19 / EU, as well as the corresponding national laws. This directive requires that the product must be taken to a special waste collection site at the end of its useful life. Applies to both device and accessories Contact your dealer if final product placement is required.



This equipment must not be disposed of as household waste

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SYSTEM OVERVIEW

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3.SYSTEM OVERVIEW

3.1.SYSTEM DESCRIPTION

Equipment for prophylaxis by ultrasound and Bicarbonate Jet, modern and bold design, assembled together composed of body and cover made of ABS (Acrylonitrile, butadiene, styrene) and digital control panel in polycarbonate. Available in cart and bench models.

Synchronized electropneumatic system, with valves that provide cuts and aspirations of water instantly, thus avoiding water contact with bicarbonate at the tip of the handpiece.

It has Piezoelectric ultrasound activated through ceramic tablets in which it allows use in operations without the use of water.

Removable bicarbonate jet pen, with concentric diffuser that mixes air, water and bicarbonate a short distance from the tip.

Transducer cover made of rigid and self-cleaning thermoplastic resin.

Function selector key with 3 programmable options, P (periodontal), E (endo) and S (scaling).

Fine adjustment potentiometer for precise regulation of ultrasonic power, suitable for each type of procedure.

Internal depressurization through automatic scanning of the bicarbonate, from the valves to the handpiece.

Bicarbonate container with easy access, transparent and removable that allows its removal without the need to turn all the equipment to remove the leftover bicarbonate powder.

Bicarbonate jet interruption system with an anti-agglutination module that prevents clogging in the valves.

It has a contamination-free system for feeding the ultrasound pens and the bicarbonate jet, through the peristaltic pump with antiseptic liquid, water, serum or similar. The peristaltic system has the function of pulsating the liquid from the reservoir to the tips (ultrasound and bicarbonate jet).

3.2.APPLICATION SPECIFICATION

The Dental Prophylaxis by Sodium Bicarbonate / Ultrasound project is intended for prophylaxis with ultrasound and bicarbonate jet, which was developed to be used in various dental practices such as: periodontics, endodontics, prosthesis, surgery and others.

3.2.1.Principles of operation

Ultrasound is derived from physical vibrations of matter particles, similar to sound waves, with a frequency higher than the level of human perception, which produces a frequency of up to 30,000 vibrations per second.

The Bicarbonate Jet (Prophylaxis) comes from the release under pressure of sodium bicarbonate particles that, together with the water, mix in the tip of the tip forming a jet in the form of a concentrated spray.

3.2.2.Significant physical characteristics

Ultrasound

A generator of ultrasonic waves (with frequency from (24,000 to 30,000 Hz) which transmit mechanical vibration to active instruments in the same frequency by means of a transducer using ceramic pieces. Helps dentists perform different procedures in various dental fields such as periodontics, endodontics, prosthetics, surgery and others.

Sodium Bicarbonate Jet

It is a pressurized mixture of air, water and sodium bicarbonate. Such mixture is conducted through

ducts to the tip of the handpiece, where it forms a uniform jet used to remove bacterial plaque, materia alba and dental stains.

3.2.3.User profile

The user to operate and handle the Ultrasound and Bicarbonate Jet must be aged between 18 and 70 years old, both sexes, with the ability to read and understand images, symbols, icons, western Arabic characters (Arial font), alpha numeric characters, to know distinguish intraoral part of the human body, not being able to present a degree of visual imperfection for reading or seeing and an average degree of impairment of recent memory, not being clearly able to perform the activities and functions of the product in a correct manner to the profession.

The user needs to be a qualified health professional and trained to perform the activities, functions frequently used in the application of Dental Prophylaxis by Sodium Bicarbonate / Ultrasound and their primary operations functions.

3.3. MAIN PRODUCT COMPONENTS







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LEGEND

- 1 Sodium bicarbonate jet handpiece (fig. A and B).
- **2** Sodium bicarbonate jet handpiece holder (fig. A and B).
- **3** Ultrasound handpiece (fig. A and B).
- 4 Ultrasound handpiece holder (fig. A and B).
- 5 Control panel (fig. A and B).
- 6 Foot control (fig. A, B e C) Optional: round or square.
- 7 Sodium bicarbonate reservoir (fig. C).
- 8 Fluid reservoir (fig. C and L).*
- 9 Hose (fig. C and L).
- 10 Electric power cable (fig. C).
- 11 Compressed air input (fig. C).
- 12 Fluid reservoir cover (fig. C).
- 13 Air filter (fig. C).
- 14 On/Off main switch (fig. C).
- 15 Endo/ Perio/ Retro tips installation wrench (fig. F and G).
- 16 Handpiece protective cover (fig. G).
- 17 Dentistry Tip adapter (fig. H).
- 18 Tightening nut (fig. H).
- 19 Dentistry Tip (fig. H).
- 20 Endo L and Dentistry tips installation wrench (fig. K).
- 21 Reservoir fastening columns (fig. C e L).
- 22 Peristaltic pump (fig. C and L).
- 23 Peristaltic Pump arm (fig. C and L).
- 24 Hose connection (fig. L).
- 25 LED (fig. G).**

^{*} Optional Reservoir Heater (Subject to commercial availability).

^{**} Handpiece with exclusive LED lighting for the Profi Neo Led e Profi Class Led models.

CONTROL PANEL (PROFI NEO / PROFI NEO LED)

- 1 On/Off switch and selection of ultrasound and sodium bicarbonate jet functions.
- **2** Ultrasound power selection switch.
- 3 Orange range indicator of recommended power limit for Endo Tips.
- 4 Orange+brown range indicator of Perio Tips power limit.
- 5 Peristaltic pump switch (on/off and fluid flow rate control).

CONTROL PANEL (PROFI CLASS / PROFI CLASS LED)

- 1 'ON/OFF' key (fig. E)
- 2 Key 🖃 (fig. E)
- 3 Key 📑 (fig. E)
- 4 'Peristaltic pump' key (fig. E) 🔘
- 5 'Bicarbonate jet' key (fig. E) 🚈
- 6 'Endo ultrasound' key (fig. E) 📩
- 7 'Perio ultrasound' key (fig. E) 🛹
- ${\bf 8}$ Access to memory programming (fig. E)
- 9 Display (fig. E)
- 10 'Dentistry' key (fig. E)

FUNCTIONS OF CONTROL PANEL KEYS (PROFI CLASS / PROFI CLASS LED)

Use the icons below to identify the control panel keys and their respective functions (fig. E):

- ON/OFF key (fig. E, item 1).
- 🖂 'Perio ultrasound' key and power setting (fig. E, item 7).
- 🕒 'Endo ultrasound' key and power setting (fig. E, item 6).
- 'Sodium bicarbonate' jet function (fig. E, item 5).
- O 'Peristaltic pump' function and irrigant flow rate setting (fig. E, item 4).
- Power lowering and language selection (fig. E, item 2).
- + Power increasing and language selection (fig. E, item 3).
- **1 2 3 4** Access to memory programming (fig. E, item 8).
- Dentistry Tip selection (fig. E, item 10).

3.4.SETS AND ACCESSORIES



All parts, accessories and options described in the owner's manual are for exclusive use.

The use of any parts, accessories or materials not specified in this manual is the sole responsibility of the user.

Accessories that accompany the product



The Kit consists of:

01. Tip Periodontal Supra (1 units)

02. Tip Periodontal Sub (2 units)

03. Nozzle cleaning needle (1 unit)

04. O-ring Int Ring 11,17 (1 unit)

05. O-ring Int Ring 12.49 (1 unit)

Accessories that do not accompany the product



Profi Line



Parts and pieces accompanying the product



01 - Silicone hose

09. Tip Retro A5

- 02 AR hose
- 03 Air tee for connection
- 04 Transducer cover
- 05 Power input cable
- 06 Tips fixing key

Consumables



01. Sodium bicarbonate

3.5.APPLIED PARTS

The following item is used in the treatment of the patient

Туре с	of parts	Contact type	Contact duration	Classification
TIPs	Removable	Mucous membrane/ Bone structure	<60s	Type B
Transducer cover Bicarbonate Jet	Removable	Mucous membrane	<60s	Туре В
Transducer cover Ultrasound	Removable	Mucous membrane	<60s	N/A
Plastic Covers	Fixed	Mucous membrane	<60s	N/A

3.6.LABEL POSITIONING

The following figure illustrates the location of the labels on the equipment components.



- A. Identification label
- B. Security label
- C. Consultation instructions manual

Alliage S/A Indústrias Médico Odontológica Roda Abrão Assed, Km 53 + 450m - CEP 14097-500 Ribeirão Preto - SP - Brazil - Tél. : +56 (16) 3512-1212 PRODUCT / PRODUTO EQUIPMENT FOR PROPHLAXIB SV ULTRASSOM / JATO DE BICARBONATE JET EQUIPMENT FOR PROPHLAXIB ULTRASSOM / JATO DE BICARBONATO MODEL SN O.P M POWER INPUT / POTENCIA DE ENTRADA 127/220 V - 50/60 Hz BIVOLT BIFÁSICO 42VA - 55VA OPERATION / OPERAÇÃO T.on: 1min / T.off: 4min

Demonstrative image Real dimensions 63 x 48 mm



Demonstrative image Real dimensions 50 x 23 mm





Demonstrative image Real dimensions diameter. 10 mm

3.7.SYSTEM REQUIREMENTS

3.7.1.Compressor requirements

The compressor is required to provide compressed air for clinical and laboratory use, having stable performance and flow capacity in accordance with the minimum requirements for the installation of the Ultrasound / Bicarbonate Jet Prophylaxis Equipment, in addition to being oil and emission free. smoke, vapors or unpleasant odors.

It must have a safety system with a valve that goes into operation to release the pressure in case the pressure switch fails and also an overload protector in order to protect the equipment from overheating. The location of your installation should be an airy place, preferably outside the office and should not be installed in sanitary facilities such as bathrooms and toilets, in order to minimize the contamination of the air used in the offices.

For the safety of the patient, the operator and the perfect functioning of the product, the installation of the compressor must respect the following recommendations:

Install a pressure relief device next to the compressor;

Install an air filter with pressure regulator, thus preventing oil, moisture and solid particles from entering the office and subsequently reaching its vital parts, such as; valves, pen, etc.;

Install the compressor close to the supply point to avoid losses;

In installations, preferably use rigid copper tubes. The pipes can also be made with galvanized steel, stainless steel, nylon or polyethylene tubes.

Pressure limit of 80 psi;

Flow rate limit \geq 47 Nl / min;

Humidity limit between 40 and 60%;

Oil contamination limit of 0.5 mg / m^3 ;

Particle contamination limit <100 particles / m³ (particles between 1 and 5µm in size);

Air quality regulations are in accordance with the laws of each country.

3.7.2.System layout



* Do not accompany the product



To meet safety standards, do not operate non-medical equipment within the patient's area.

Outside the patient's area, the presence of non-medical equipment is acceptable, provided approved and certified equipment is used.



It cannot be possible to connect any component of the system with other equipment not recommended by the manufacturer.



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4.OPERATION

4.1.INITIAL PREPARATION



To isolate the equipment from the mains, use the general switch.

To start operating the equipment, follow the instructions below.

4.1.1.Profi Class / Profi Class LED

1. Switch on the device by pressing the on / off switch.

2. Select the function Ultrasound or Bicarbonate Jet.

3. If the ultrasound function is selected, select the operation mode - periodontics, endodontics and scaling.

4. Operate the pedal.

4.2. USE OF EQUIPMENT

4.2.1. Ultrasound Operation

Use of ultrasound

Attach the protective cover to the handpiece (fig. G, item 16).

Endo / Perio / Retro Tips

Insert the selected tip into the wrench (fig. F, item 15) and then screw it onto the handpiece (fig. G).

Dentistry Tips

Screw the adaptor (fig. H, item 17) onto the handpiece and tighten it firmly with the aid of an installation wrench.

Screw the nut (fig. H, item 18) a little loosely and insert the desired Dentistry Tip, pushing it flush with the stop. The set must be like in (fig. I).

Tighten the nut with the installation wrench.

To change the Tip, loosen the nut (fig. H, item 18), remove the Tip, insert the desired Tip and tighten the nut again.



We recommend that after use, do not leave the handpiece with the Tip in the tip holder in order to avoid accidents.

Remove the reservoir cover (fig. C, item 12) and fill it up with your preferred fluid, without exceeding the maximum volume indication line (check section "Reservoir replenishment"). Turn the equipment on by setting the main switch (fig. C, item 14) to the 'ON' position.

Profi Neo / Profi Neo LED - Set the selection switch (fig. D, item 1) to the "Ultrasound" function. **Profi Class / Profi Class LED** - Press the ON/OFF key (fig. E, item 1) and then key or b. Set the ultrasound power and the irrigant flow rate (check the corresponding sections in this Manual). Press the foot control to start the procedure (fig. A and B, item 6)..

Upon finishing the procedure, release the foot control, place the handpiece into the holder and

turn the equipment off.

4.2.2. Bicarbonate jet operation

Use of the sodium bicarbonate jet

The bicarbonate jet removes dark stains from the teeth, caused by cigarettes, coffee, tea, etc., associated with bacterial plaques and not the stone.

To obtain the best result from the bicarbonate jet, we recommend respecting the distance of the handpiece in relation to the tooth (5mm), with an inclination of 30 ° to 45 ° describing small circular movements over the teeth.

To avoid unpleasant sensations in patients, the jet of bicarbonate should be directed to the occlusal edge and not to the gingival sulcus.

This equipment is contraindicated for use in patients who have serious respiratory or renal disorders or who are undergoing hemodialysis, these cases must have medical monitoring. We recommend the use of a mask and glasses to apply the bicarbonate jet.

1 - Remove the sodium bicarbonate reservoir cover (fig. C, item 7) and if there are residues on the bottom, clean with the aspirator.

2 - Rub the Dabi Atlante sodium bicarbonate packet for a while to homogenize it.

3 - Pour one Dabi Atlante sodium bicarbonate packet inside the reservoir. Never pour more than the indicated by the volume level mark.

4 - Replace the cover. Make sure the reservoir is tightly closed.

5 - Remove the fluid reservoir cover (fig. C, item 12) and fill it up with your preferred fluid (check section "Reservoir replenishment").

- 6 Turn the equipment on by setting the main switch (fig. C, item 14) to the 'ON' position.
- 7 Profi Neo/ Profi Neo LED Set the selection switch (fig. D, item 1) to the bicarbonate jet function.
- 8 Profi Class/Profi Class LED Press the ON/OFF key (fig. E, item 1) and then key 🔄 .
- 9 Set the irrigant flow rate (check section "The irrigant flow rate setting").



To speed up such setting, we recommend the use of the memory keys in Profi Class/Profi Class LED.

Press the foot control to start the procedure (fig. A and B, item 6).

Upon finishing the procedure, release the foot control, place the handpiece into the holder and turn the equipment off.



Do not add more than 40g of bicarbonate to the container as this will cause the powder to become blocked.



The effectiveness depends on the dosage of the water volume and the amount of powder.



In order to activate only the peristaltic pump in Profi Class/Profi Class LED (fig. E, item 4), disable the last function "sodium bicarbonate jet" or "ultrasoud" by pressing the corresponding key.

4.2.3. Techniques and applications

All the tips of the ultrasound have the peculiarity of vibrating in a single plane (vibrations from front to back, and on the tip axis).

The lateral vibrations common to other scalers do not exist, the straight displacement favors a more precise approximation of the tooth and gum.

Enamel and cement are protected from useless shock.

Within this main plane of vibration, the tip of each tip is driven by small vibratory movements.

To obtain the maximum performance of the ultrasound, the operator must consider the vibration settings, specific to each Tip.





The shape and weight of each Tip are determining factors to obtain maximum performance of the ultrasound generator, the operator's attention to these two characteristics, will ensure the maintenance of the best performance of the unit, however, we recommend that the structure of the Tip is not altered (by filing or twisting it), in the same way the aging of a Tip leads to a change of its original characteristic, making it ineffective. Any Tip that has been damaged by use or by accidental impact must be replaced.

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4.2.4.Supply of the bicarbonate jet

To supply your equipment, we recommend the use of "Clean Okta" sodium bicarbonate (Reg. ANVISA 80339810002) or one that has similar characteristics:

Product composition: Sodium Bicarbonate (99.6%), Silicic Anhydrous, Essence - 105 microns spherical pattern.

For more information about the product, see the manufacturer's instructions on the product.

4.2.5.Filling the peristaltic pump reservoir



Before filling the tank, check that the water to be used does not contain impurities or contaminants.

1 - Remove the fluid reservoir cover (fig. C, item 12). Pull the hose (fig. L, item 9) until it detaches from the trough inside the reservoir.

2 - Pull the reservoir up until it detaches from the two body columns of the unit (fig. L, item 21). Press the pump side rods (Fig. L, item 23) and pull the pump to detach it from the unit body. Disconnect the hose (fig. L, item 24).

3 - Connect the new pump hose and attach it to the unit body by pressing the side rods. Insert the hose into the irrigant reservoir trough and attach it to the body fastening columns, making sure the hose is not kinked.



After you finish using the product, turn it off using the main switch and remove it from the outlet.

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CLEANING, DISINFECTION AND STERILIZATION

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5.CLEANING, DISINFECTION AND STERILIZATION



Before starting the cleaning and disinfection procedure, turn off the main switch of the equipment to avoid permanent damage.



For your protection, during the cleaning and disinfection process of the equipment use PPE such as disposable gloves and goggles.

5.1.EQUIPMENT

The cleaning and disinfection process must be performed at each patient change.

When starting the process, check for visible dirt, such as blood or saliva.

Thoroughly clean the patient's entire contact area.

For cleaning use a clean, soft cloth moistened with mild soap and then dry with a clean, soft cloth or paper towel.

For the disinfection process of the equipment, use foam disinfectant detergent that has active components based on didecyldimethylammonium chloride.

Apply the disinfectant detergent foam on the surface or on a clean cloth and spread it on the surface to be treated. Respect the antimicrobial contact time specified by the manufacturer.

After application, allow to dry. Do not rinse.

Some of the removable parts that come in contact with the patient can be autoclaved. These parts are: Ultrasound cover, Tips, Bicarbonate jet covers and Insert grip wrench.

5.1.1.Ultrasound

Remove the Tip of the transducer, and then remove the cover by means of pressure, "do not try to rotate", then take it for autoclaving (packed).



5.1.2.Bicarbonate Jet

Unthread the handpiece of the bicarbonate jet (A) and then remove the hose (B), as it cannot be autoclavable and take it for sterilization in an autoclave (packed).



Profi Line

After each day of use, place the equipment on a dry piece of cloth, incline it sideways at an angle of approximately 45°. Drain the air filter (fig. C, item 13) by pressing the depressurization pin until all the water has been eliminated.





Minimum sterilization time 30 minutes at 121°C or minimum 3 minutes at 135°C, and equivalent temperature and time parameters can be used within this range. Maximum sterilization temperature 135 °C.

If these items are autoclaved, disinfection by alternative methods is not necessary. There is no limit on cycles or application time that the equipment and its parts can tolerate during the cleaning, disinfection and / or sterilization process, following the instructions in this manual.



Do not spill liquid disinfectant on the equipment.



Do not use organic solvents, for example, thinner, to clean the equipment. In the event that the developer solution is spilled on the panel, clean it immediately, as these solutions may compromise the equipment's paint.



Sterilization parameters must always be followed. Accessories that are not properly sterilized can cause disease in patients.

5.2. BICARBONATE CONTAINER

Locate the bicarbonate container through the side access, remove it by turning it counterclockwise and clean it with a dry cloth.

Check that the thread is completely free of dust and replace it by turning it clockwise.

TROUBLESHOOTING

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6.TROUBLESHOOTING

6.1.SOLUTION OF PROBLEMS

Occasionally, malfunctions may occur during use. In the event of an error, restart the equipment and resume operation. If the problem persists, follow the instructions below.

Failures	Probable causes	Solutions
	Not plugged in the power soure outlet	Plug it in
Equipment will not start	Fuse is burned out	Replace fuse
	Power failure	Await normalization
	Circuit breaker is off	Turn it on
System is not pressurized	Compressor is off and/or valve is closed	Turn compressor on and/or open valve
Tip will not oscillate but LED goes on	Tip is not correctly fixed	Fix tip appropriately or try another tip
Water leakage in the handpiece	Tip is incorrectly screwed on	Screw tip on properly or replace O-ring
	Sodium bicarbonate jet handpiece terminal is not properly screwed on	Properly tighten the sodium bicarbonate jet handpiece terminal
Sodium bicarbonate will not	Tip is clogged	Unclog tip
come out	No sodium bicarbonate in the reservoir	Supply the reservoir
	Sodium bicarbonate excess in the reservoir	Remove excess
Water will not come out	No irrigating solution in the reservoir	Supply the reservoir
	Peristaltic pump is not enabled	Activate the peristaltic pump
Perception of electric shock on tip	No or ineffective grounding	Ground equipment adequately

If problems persist, contact the Alliage Service Department.

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INSPECTION AND MAINTENANCE

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7.INSPECTION AND MAINTENANCE



Maintenance or service procedures may only be carried out by a technical service authorized by the manufacturer.

All instructions for using the equipment as intended are provided in this user guide. If a problem is detected and cannot be corrected with the instructions in the problem diagnostics section, contact the Alliage Service Department.

7.1.PERIODIC INSPECTION

It is imperative that this equipment be regularly inspected to ensure operational safety and functional reliability. This inspection must be carried out by personnel familiar with the necessary precautions to avoid exposing the patient to risk.

Periodic inspection should be carried out at regular intervals (at least once a year) to ensure that the product is permanently safe and operational. All components subject to normal wear and tear should be checked and, if necessary, replaced.

The manufacturer and the assembler / installer are exempt from responsibility for the standard results not being compliant in cases where the user does not perform the maintenance recommended by the manufacturer.

Neither inspection nor service is part of the equipment warranty.

Maintenance performed must be documented and maintained with the equipment.

The table below gives a description of the main inspection items and recommended frequency.

Item	Inspection description	Recommended frequency
Operation / Security System	Foot pedal, tip power, water flow, dust flow, jet pressure (Auditory and visual).	Diary
Electrical parts	Overheating / Noise / Burning smell (Auditory and visual).	Monthly
Parts and pieces	Operation / Noise / Vibration (Auditory and visual).	Yearly
Pedal and Controls	Operation / Damage (Auditory and visual).	Yearly

If problems are detected during the inspection, contact the Alliage Service Department.

7.2.PREVENTIVE MAINTENANCE

In addition to the annual inspection, to ensure a long service life and smooth operation of your equipment, it is important to carry out preventive maintenance for a maximum period of three (3) years.

Contact the Alliage Service Department about our periodic review and preventive maintenance program.



Profi Class/Profi Class LED: O sistema identifica o desgaste da bomba e quando necessário sugere a troca da mesma por meio de um aviso no display.

7.3.CORRECTIVE MAINTENANCE



The corrective maintenance that can be performed by the user Ultrasound / Bicarbonate Jet Prophylaxis equipment is limited to unblocking the bicarbonate jet pen.



Do not open the equipment or attempt to repair it yourself or with the help of someone without training or authorization. This can aggravate the problem or produce a failure that can compromise the safety of the equipment.



The power cables and electronic boards can be changed only by the authorized technician.



The equipment or any of its parts cannot be maintained or serviced during use with a patient.



The equipment contains parts under high voltage. Risk of electric shock. Turn off the main switch before servicing.



The service manual is only available for Authorized Technical Assistance.

7.3.1.Equipment

Alliage declares that the provision of circuit diagrams, component lists or any other information that provides technical assistance on behalf of the user, can be requested as long as previously agreed between the user and Alliage.

The warranty will be void if the original parts are removed / replaced by unauthorized service technicians.

7.3.2. Bicarbonate Jet Pen

The "Jet of bicarbonate" pen is equipped with an automatic system for depressurization and internal cleaning of hoses and handpiece. With the function key positioned in Bicarbonate Jet, when the command pedal is stopped, there will be an internal sweeping air jet of the entire system, however, if there is blockage in the system, proceed as follows:

a. Remove the hose (20) from the spout (21), direct the tip to a suitable location (spit, sink bowl, etc.) and operate the pedal to make sure that the clog is in the spout (21).

b. Clean the hole with the plunger (22), inserting it until it crosses completely several times.

c. Replace the hose (20) nozzle (21). If necessary, replace the hose (20).

d. Remove the adapter (23) from the bicarbonate jet tip (24) by unscrewing it counterclockwise and take the bicarbonate jet tip for autoclaving (packaged).



7.4.ALLIAGE AUTHORIZED SERVICE NETWORK

All services performed on the Alliage equipment must be performed by an Authorized Technical Assistant, as otherwise they will not be covered by the warranty.

If you need to request electrical diagrams and or specification of components that is not stated in the user manual, use the Alliage Customer Service to make the request.

Telephone: +55 (16) 3512-1212

Address: Rodovia Abrão Assed, Km 53 - Recreio Anhanguera – Ribeirão Preto -SP/ Brazil ZIP CODE 14097-500



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8.WARRANTY

This equipment is covered by the warranty periods, terms and conditions contained in the Warranty Certificate that comes with the product.

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STANDARDS AND REGULATIONS

Alliage S/A Industrias Médico Odontológica Telefone: +55 (16) 3512-1212 Rodovia Abrão Assed, Km 53 – CEP 14097-500 –Ribeirão Preto – SP –Brasil

9.STANDARDS AND REGULATIONS

This equipment has been designed and manufactured to meet the following rules:

ABNT NBR IEC 60601-1:2010 Amendment 1:2016	Medical Electrical Equipment - Part 1: General requirements for basic security and essential performance.
ABNT NBR IEC 60601-1-2:2017	Medical Electrical Equipment, Part 1-2: General basic safety requirements and essential performance - Collateral standard: Electromagnetic interference - Requirements and tests.
ABNT NBR IEC 80601-1-60:2015	General requirements for basic safety and essential performance of dental equipment.
ABNT NBR 60601-1-6:2011	Electromedical equipment - Part 1-6: General requirements for basic safety and essential performance - Collateral standard: Usability.
ABNT NBR IEC 62366:2010	Health products - Application of usability engineering to health products.
IEC 60601-1-9:2007+AMD1:2013	Medical electrical equipment - Part 1-9: General requirements for basic safety and essential performance - Collateral standard: Requirements for environmentally conscious design.
IEC 62304:2006	Medical device software - Software lifecycle processes.
ISO 10993-1:2018	Biological assessment of medical devices - Part 1: Assessment and testing.
ABNT NBR ISO 14971:2009	Medical devices - Application of risk management to medical devices.
ABNT NBR ISO 13485:2016	Quality management systems - Requirements for regulatory purposes.

10 TECHNICAL SPECIFICATIONS

Alliage S/A Industrias Médico Odontológica Telefone: +55 (16) 3512-1212 Rodovia Abrão Assed, Km 53 – CEP 14097-500 –Ribeirão Preto – SP –Brasil

10. TECHNICAL SPECIFICATIONS

10.1 EQUIPMENT CLASSIFICATION

Characteristics	Profi Neo / Profi Neo LED	Profi Class / Profi Class LED
Ultrasound Handpiece	х	х
Sodium Bicarbonate Jet Handpiece	х	х
2 Handpieces: Ultrasound and Bicarbonate Jet	х	х
Injected ABS Framework	х	х
Frequency Stabilizer Electronic Circuit	х	×
Piezoeletric Ceramic Transdutor	х	×
Transdutor Removable Caps for sterilization	х	×
Keys with protection for tips installation	х	×
Sodium Bicarbonate Reservoir in injected ABS	х	×
Transparent lid for the sodium bicarbonate reservoir	х	х
Air filter	х	х
Unique Foot Control	х	х
Handpiece with smooth and flexible hoses	х	×
Peristaltic Pump Irrigation	х	×
Control Panel with Knobs	х	
Endodontics function	0	×
Dentistry function	0	×
Periodontics function	х	х

X = Serial characteristics and technical features 0 = Optional characteristics and technical features

Product classification according to ANVISA	
Class of framing	Class II (medium risk)
Product classification according to NBR IEC 60601-1 (GENERA	L STANDARD)
Type of protection against electric shock	Class I
Degree of protection against electric shock	Type B equipment
Protection against harmful water penetration	IPX0 - Cabinet IPX1 - Pedal
Degree of safety of application in the presence of a flammable anesthetic mixture with air, oxygen or nitrous oxide	Not suitable
Operation mode	Non-continuous (ON: 1min. / OFF: 4min)



The pedal cannot be installed in an emergency room.

10.2.APPLIANCE INFORMATION (GENERAL)

Connection to suitable power supply

External power supply

Power supply voltage

127 / 220 V~ (Bivolt)

Power supply frequency

50 / 60 Hz

Allowable fluctuation

+/- 10 %

Number of phases

Biphasic

Stand-by rated current

2,6 mA - 127 V~ 1,5 mA - 220 V~

Rated current during load

3,5A (máx) - 127 V~ 2,5A (máx) - 220 V~

Master key

Contact resistance: maximum of 20 milliohms with application of 1A in VCC; Electrical characteristics: 10A / 120 VAC; Insulation resistance: minimum of 1,000 megaohms;

Power consumption

42VA-55VA

Maximum grid impedance

0,2Ω

Net weight

Profi Neo: 3,2 kg Profi Neo LED: 3,2 kg Profi Class: 3,2 kg Profi Class LED: 3,2 kg

Gross weight

Profi Neo: 3,2 kg Profi Neo LED: 3,2 kg Profi Class: 3,2 kg Profi Class LED: 3,2 kg

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10.3.SPECIFIC INFORMATION

Air pressure

80 PSI (5,52 BAR)

Maximum air consumption

80 l/min

Water tank capacity

1000 ml

10.4.ULTRASOUND SPECIFICATIONS

Ultrasound Vibration Frequency

30.000 Hz

Consumption of irrigating liquid

28 ml/min

Power consumed

15 VA

Transducer system

Electric piezo ceramic

Maximum tip temperature in normal use

114,2°C

Maximum rate of temperature rise of tip

179,7°C

10.5.ENVIRONMENTAL CONDITIONS

Environmental conditions of transport and storage

Transport or storage ambient temperature range

-29°C to +60°C

Transport and storage relative humidity range

20% to 90%

Atmospheric pressure range

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500 hPa to 1060 hPa
(375 mmHg to 795 mmHg)
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Environmental conditions of installation and operation

Ambient operating temperature range

+10°C to +34°C

Profi Line

Recommended ambient temperature range

+21°C to +26°C

Operating relative humidity range (non-condensing)

30% to 75%

Atmospheric pressure range

700 hPa to 1060 hPa (525 mmHg to 795 mmHg)

Operating altitude

≤ 2000 m

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ELECTROMAGNETIC COMPATIBILITY

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11.ELECTROMAGNETIC COMPATIBILITY

The Equipment for Prophylaxis by Ultrasound / Bicarbonate Jet is intended for use in the electromagnetic environment specified below. The buyer or user should ensure that it is used in such an environment.

The Equipment for Prophylaxis by Ultrasound / Bicarbonate Jet is suitable for use in a professional health care environment, not including areas where sensitive equipment or sources of intense electromagnetic disturbances are present, such as the RF shielded room of a magnetic resonance imaging system. , in operating rooms close to active AF surgical equipment, electrophysiology laboratories, armored rooms or areas where short wave therapy equipment is used.

The following tables provide information on the equipment's compliance with the ABNT NBR IEC 60601-1-2: 2017 standard.

11.1.GUIDANCE AND DECLARATION FOR ELECTROMAGNETIC EMISSIONS

Emissions Tests	Compliance	Electromagnetic Environments - guidelines		
RF emissions CISPR 11	Group 1	The Equipment for Prophylaxis by Ultrasound / Bicarbonate Jet uses RF energy only for its internal functions. Therefore, its RF emissions are extremely low and are unlikely to cause any interference in nearby electronic equipment.		
RF emissions CISPR 11	Class A	The Equipment for Prophylaxis by		
Harmonic emissions IEC 61000-3-2	Class A	Ultrasound / Bicarbonate Jet is suitable for use in all establishments, except households and those directly connected		
Voltage fluctuation / Scintillation emissions IEC 61000-3-3	Compliant	to the public low voltage power supply network that powers buildings used for domestic purposes.		

Note: The emission characteristics of this equipment make it suitable for use in industrial areas and hospitals (IEC / CISPR 11, Class A). If used in a residential environment (for which IEC / CISPR 11, Class B is normally required), this equipment may not provide adequate protection for radio frequency communication services. The user may need to take mitigation measures, such as relocating or redirecting equipment.

11.2. ORIENTATION AND DECLARATION FOR ELECTROMAGNETIC IMMUNITY

Phenomenon	Basic EMC standard or test method Immunity test level		Compliance level	
Electrostatic discharge	IEC 61000-4-2	±8 KV contact ± 2 KV, ± 4 KV, ± 8 KV, ± 15 KV air	±8 KV contact ± 2 KV, ± 4 KV, ± 8 KV, ± 15 KV air	
EM fields of radiated RF	IEC 61000-4-3	3 V/m 80 MHz – 2,7 GHz 80 % AM at 1 kHz	3 V/m 80 MHz – 2,7 GHz 80 % AM at 1 kHz	
Fields in the vicinity from RF wireless communications equipment	ity from IEC 61000-4-3 See table nications		See table	
Fast / saved electrical	IEC 61000-4-4 alternating current power input	± 2 kV 100 kHz frequência de repetição	± 2 kV 100 kHz frequência de repetição	
transients	IEC 61000-4-4 signal input / output	± 1 kV 100 kHz repetition frequency	± 1 kV 100 kHz repetition frequency	
Outbreak Line by line	IEC 61000-4-5	± 0,5 kV, ± 1 kV	± 0,5 kV, ± 1 kV	
Outbreak Ground line	IEC 61000-4-5	± 0,5 kV, ± 1 kV, ± 2 kV	± 0,5 kV, ± 1 kV, ± 2 kV	
Conducted disorders induced by RF fields	IEC 61000-4-6	3 V 0.15 MHz - 80 MHz 6 V in ISM bands between 0.15 MHz and 80 MHz 80% AM at 1 kHz	3 V 0.15 MHz - 80 MHz 6 V in ISM bands between 0.15 MHz and 80 MHz 80% AM at 1 kHz	
Magnetic fields at the stated feed frequency	IEC 61000-4-8	30 A/m 50 Hz or 60 Hz	30 A/m 50 Hz or 60 Hz	
Voltage drops	IEC 61000-4-11	0 % UT; 0,5 cycle At 0°, 45°, 90°, 135°, 180°, 225°, 270° and 315° 0 % UT; 1 cycle and 70 % UT; 25/30 cycles Single phase: at 0°	0 % UT; 0,5 cycle At 0°, 45°, 90°, 135°, 180°, 225°, 270° and 315° 0 % UT; 1 cycle and 70 % UT; 25/30 cycles Single phase: at 0°	
Voltage interruptions	IEC 61000-4-11	0 % UT; 250/300 cycles	The device will shut down and / or reset if the power is interrupted for five seconds.	

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NOTE 1 At 80 MHz and 800MHz, the higher frequency range is applicable.

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

NOTE 3 UT is the AC mains voltage before applying the test level.

Test frequency (MHz)	Band (MHz)	Service	Modulation	Maximum power (W)	Distance (m)	Immunity test level (V/m)
385	380-390	TETRA 400	18 Hz pulse modulation	1,8	0,3	27
450	430-470	GMRS 460,FRS 460	FM deviation of ± 5 kHz 1kHz sinusoidal	2	0,3	28
710	704-787	Band LTE	217 Hz pulse	0,2	0,3	9
745		13, 17	modulation			
7480						
810	800-960	GSM	18 Hz pulse	2	0,3	28
870		800/900, TETRA 800.	00/900, modulation TRA 800, EN 820, MA 850, nd LTE 5			
930		iDEN 820, CDMA 850, Band LTE 5				
1720	1700 -1990	GSM 1800;	217 Hz pulse	2	0,3	28
1845		CDMA 1900: GSM	modulation			
1970		1900; DECT; Band LTE 1, 3, 4, 25; UMTS				
2450	2400-2570	Bluetooth, W L A N 8 0 2 . 1 1 b/g/n, RFID 2450, Band LTE 7	217 Hz pulse modulation	2	0,3	28
5240	5100 - 5800	WLAN	217 Hz pulse	0,2	0,3	9
5500		802.11 a/n	modulation			
5785						

Proximity fields from wireless RF communications equipment

List of used cables

Cables		Description	Length
Power		Tripolar Power Cable 3x Gauge 2.50 mm², 250V AC, Male Plug 20A NBR 14136 2P + T, with female plug, INMETRO.	3 m
	The Ed assist of EM betwe	quipment for Prophylaxis by Ultrasound / Bicarbonate J the health professional, and it is for exclusive dental u IC disturbances, the operator may experience loss of een the equipment and controls.	et is intended to se. In the event communication
	Compl that h equipr	iance with EMC and EMI standards cannot be guaranteed have been altered or that do not comply with the same s ment has been validated.	by using cables tandards as the
	Use of result the ot	this equipment adjacent to other equipment should be a in improper operation. If this use is necessary, it is advisa ther equipment be observed to verify that they are oper	voided as it may ble that this and rating normally.
	Do no cables result in imp	t use accessories, transducers, internal parts of compoi s other than those previously specified by the manufa in increased emission or decreased electromagnetic imm roper operation.	nents and other cturer. This can unity and result
<u>^</u>	Portal cables the Pr specif equipr	ble RF communication equipment (including peripherals s and external antennas) should not be used within 30 c rofi Neo / Profi Neo LED / Profi Class / Profi Class LED, i ied by the manufacturer. Otherwise, performance deg ment may occur.	such as antenna m of any part of ncluding cables radation of this
	To mai life, al and fo	intain basic safety from electromagnetic disturbances dur lways use the equipment in the specified electromagne ollow the maintenance recommendation described in thi	ng the expected tic environment s manual.
	The pi must i	ins, connector sockets or elements that carry the ESD not be touched or connected without ESD protection me	warning symbol easures.

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