

CA PRIMA

ENG INSTRUCTIONS FOR USE.





Devices



CA PRIMA 1:1 REF 1601235-001



CA PRIMA 1:1 L REF 1601236-001



CA PRIMA 1:1 L MS REF 1601237-001



CA PRIMA 1:121 REF 1601267-001

Optional accessories (REF)



MAINT SPRAYNET® (BOX 6 CANS) REF 1600036-006



MAINT LUBRIFLUID® (BOX 6 CANS) REF 1600064-006



MAINT AQUACARE (BOX 6 CANS) REF 1600617-006



IRRIGATION LINE (10/pkg) REF 1500984-010



External Irrigation system (10/pkg) REF 1500431-001

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ENG INSTRUCTIONS FOR USE

1 Symbols

1.1 Description of symbols used

Symbol	Description	Symbol	Description
Syllibot	Description	Symbol	Description
C€ 0123	CE Marking with number of the notified body.	***	Manufacturer.
\triangle	WARNING: hazard that could result in serious injury or damage to the device if the safety instructions are not correctly followed.	i	Consult instructions for use or consult electronic instructions for use.
À	CAUTION: hazard that could result in light or moderate injury or damage to the device if the safety instructions are not correctly followed.		Data Matrix code for product information including UDI (Unique Device Identification).
x: X·	Temperature limitation.	X2	Humidity limitation.
(1) (1) (1) (1) (1) (1) (1) (1) (1) (1)	Atmospheric pressure limitation.	MD	Medical device.
Ť	Keep away from rain.	SN	Serial number.
	Wear rubber gloves.	REF	Catalogue number.
Rx Only	Warning: in accordance with federal law (USA), this device is only available for sale upon recommendation by an accredited practitioner.	EC REP	Authorized EC Representative in the European Community.
	General symbol for recovery/ recyclable.	135°C	Sterilization up to the specified temperature.
「ĭ	Can be processed in an automated washer/disinfector for thermal disinfection.	-\̈́-	Lamp; lighting, illumination.

2 Identification, intended use and notation

2.1 Identification

Medical devices manufactured by Bien-Air Dental SA.

Type

Dental contra-angles handpiece (CA), push-button bur locking, with or without light, with internal, mixed or external sprays.

Description

Bien-Air Dental contra-angle are designed to transmit and apply the mechanical energy produced by an air or electric micromotor.

See the table below for a summary of your contra-angles type.

Instrument	Light		Sprays		Lenght	
Ratio	With li	ght Without light	Internal spray	External spray	Standard	Micro-series
• CA PR	IMA 1:1	•	•		•	
• CA PR	IMA 1:1 L •		•		•	
• CA PR	IMA 1:1 MS		•			•
• CA PR	IMA 1:121	•		•	•	

2.2 Intended Use

Devices intended for use in general dentistry for restorative dentistry, dental prophylaxis, orthodontics and endodontics:

CA PRIMA 1:1, CA PRIMA 1:1 L, CA PRIMA 1:1 L MS, CA PRIMA 1:121

Device intended for use in oral surgery, maxillofacial surgery and in periodontology as well:

• CA PRIMA 1:121

2.3 Intended patient population

The intended patient population for the contra-angles includes any person visiting a dental practitioner's office to receive treatment in line with the intended medical condition. There is no restriction concerning subject age, race, or culture. The intended user is responsible to select the adequate device for the patient according to the specific clinical application.

2.4 Intended user

Product intended for professional use only. Used by dentists and dental professionals.

2.5 Use Environment

Professional healthcare facility environment.

2.6 Intended Medical Conditions

- General dentistry which includes restorative dentistry, dental prophylaxis, orthodontics and addresses the maintenance or reestablishment of dental health.
- Oral surgery treatments include impacted teeth extraction, wisdom teeth extraction, non-salvageable decayed teeth extraction, guided and not-guided bone regeneration, apicoectomy, osteotomy, sequestrectomy and hemi section.

2.7 Patient contra-indication and side effects

No specific patient contra-indication, side effects nor warnings exist for the contra-angle devices when the devices are used as intended.

2.8 In case of accident

If an accident occurs, the device must not be used until repairs have been completed by a qualified, authorized and trained technician in a repair center.

If any serious incident occurs in relation to the device, report it to a competent authority of your country, as well as the manufacturer through your regional distributor. Observe relevant national regulations for detailed procedures.

⚠ WARNING

Any use other than that for which this device is intended is prohibited and may prove dangerous.



3 User and Patient Safety: Warnings & Precautions for use

↑ WARNING

The device must be used by qualified dental professionals in compliance with the current legal provisions concerning occupational safety, health and accident prevention measures, and these instructions for use. In accordance with such requirements, the operators:

- Must only use devices that are in perfect working order; in the event of irregular functioning, coolant failure, excessive vibration, abnormal heating, unusual noise or other signs that may indicate malfunction of the device, the work must be stopped immediately; in this case, contact a repair center that is approved by Bien-Air Dental SA and have the service personnel carry out repair work.
- Must ensure that the device is used only for the purpose for which it is intended, must protect themselves, their patients and third parties from any danger.
- Any modification of the medical device is strictly forbidden.

To prevent any risk of infection, the warnings below must be observed:

⚠ WARNING

- Rest the device on a cleanable support to avoid risks of infection for yourself, the patient or third parties.
- Personal protective equipment is mandatory when operating the devices.
- The device is supplied not sterile. Respect the cleaning, sterilization and maintenance procedure detailed in section 6. Sterilization before first use is mandatory.
- When disposing of the device, the user must return it sterilized to their dealer or contact an authorized body for the treatment and recovery of this type of equipment.
- Medical personnel using or performing maintenance on medical devices that are contaminated
 or potentially contaminated must comply with universal precautions, in particular the wearing of
 personal protective equipment (gloves, goggles, etc.). Pointed and sharp instruments should be
 handled with great care.
- While performing surgical treatment, the handpiece must not receive pressurised cooling air from the unit, to prevent contamination of the area being treated.

To prevent any risk of device overheating, the cautions below must be observed:

A CAUTION

- Respect the cleaning, sterilization and maintenance procedure detailed in section 6.
- The device must not be started without a bur inserted into the chuck.
- To avoid overheating of the push-button, it should not be pressed inadvertently while the instrument is rotating.

To prevent any risk of injury the warnings below must be observed:

⚠ WARNING

- The device is intended for professional use only.
- Respect the cleaning, sterilization and maintenance procedure detailed in section 6.
- In the event of excessive vibrations, abnormal heating, unusual noise or other signs suggesting that the device is malfunctioning, work must be suspended immediately. In this case, contact a repair center approved by Bien-Air Dental SA.
- Never insert or remove a device while the micromotor is rotating.
- Do not touch the dental bur while it is rotating.
- Never push the push-button while the contra-angle handpiece is in operation.
- Each time a bur is inserted, check that the bur is fully inserted to the stop and rotates freely. If it is blocked, contact your usual supplier or Bien-Air Dental SA for repair.
- Always check that the bur is locked by gently pushing and pulling the bur.
- Follow the guidelines for use, according to the bur manufacturer's instructions. Never use a bur if the shaft is not compliant, as there is a risk it can become detached during the procedure and injure the practitioner, the patient or third parties.
- Never partially insert a bur to increase its active length. Comply with maximum lengths by always inserting the bur as far as possible into the locking mechanism. If a bur is operated at high speeds when incorrectly mounted (i.e. not fully inserted into the locking mechanism, or being longer than the values specified in section 4.2) a centrifugal force may be exerted which may bend or break the
- Always ensure that the coolant supply provided by the motor is sufficient and adequate.
- Always ensure that the irrigation flow is sufficient and adequate and that the spray outlets are not
- These medical devices are intended to be used at a maximum height of 1.5 m, to avoid damage in the event of a fall.

To prevent any risk of device malfunction the caution below must be observed:

⚠ CAUTION

- Before performing any clinical application, always test your device to ensure its proper operation.
- Only use original Bien-Air Dental SA devices and accessories or those recommended by Bien-Air Dental SA.
- Respect the cleaning, sterilization and maintenance procedure detailed in section 6.







FIG. 1 FIG. 2 FIG. 3

4 Description

4.1 Overview

FIG. 1 & FIG. 2

- (1) Micromotor connection
- (2) Light output
- (3) Cutting tool connection
- (4) Color coding for transmission ratio
- (5) Push-button with a bur-locking system

FIG. 3

- (1) Micromotor connection
- (2) Cutting tool connection
- (3) Color coding for transmission ratio
- (4) Push-button with a bur-locking system
- (5) External irrigation system
- (6) Plastic holder for the external irrigation system

Note: The technical specifications, illustrations and dimensions contained in these instructions are given merely as an indication. They may not give rise to any claim.

The original language of these instructions for use is English.

For any further information, please contact Bien-Air Dental SA at the address given on the back cover.

4.2 Technical Data

Tanksiani Data	lafa-matica.
Technical Data	Information
Motor coupling compatibility	Coupling according to ISO 3964 - CA PRIMA 1:1 can be coupled to type 2 coupling (air and water) with all coupling dimensions*1 - CA PRIMA 1:1 L can be coupled to type 3 coupling (air, water and light) with all coupling dimensions - CA PRIMA 1:1 L MS can be coupled to type 3 coupling (air, water and light) with a short or extra short dimension - CA PRIMA 1:121 can be coupled to type 1 coupling (no air, no water and no light) with all coupling dimensions*2
Lighting	"L" letter means light CA & PM without L letter means no lighting
Transmission ratio according to ISO 14457	Speed direct ratio 1:1 (blue color)
Motor max speed	40'000 rpm
Bur max speed	40'000 rpm
Irrigation type	Internal Intramatic® irrigation External irrigation for CA PRIMA 1:121
Recommended spray water pressure (for an optimal cooling mist)	100-200 kPa ^{*3}
Recommended spray water pressure (for an optimal cooling mist)	200-400 kPa ^{*4}

^{*1} Can also be coupled to type 3 coupling (air, water, and light)

⚠ WARNING

For devices with external irrigation:

- Always ensure that the coolant supply is sufficient and adequate.
- The minimum irrigation quantity at the entrance of the CA must be set to at least 60mL/min.

^{*2} Can also be coupled to type 4 coupling (only light)

^{*3} The maximum spray water pressure that the contra-angle can withstand is 300 kPa

^{*4} The maximum spray air pressure that the contra-angle can withstand is 600 kPa

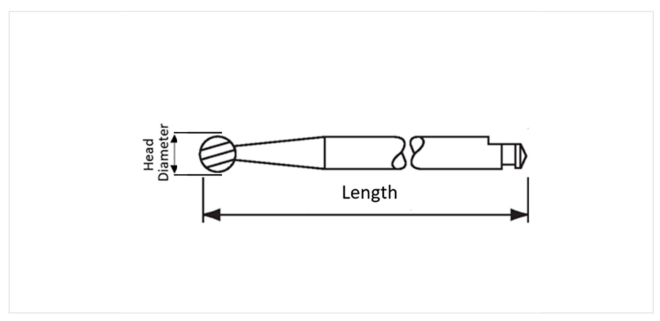


FIG. 4

Cutting tool compatibility	CA PRIMA 1:1 CA PRIMA 1:1 L CA PRIMA 1:1 L MS	CA PRIMA 1:121
Shaft diameter ^{*1} ISO 1797	2.35 mm (Type 1)	2.35 mm (Type 1)
Shaft length ISO 1797	≥ 11 mm	≥ 12 mm
Cutting tool diameter ISO 6360-1	≤ 3 mm	≤ 3 mm
Total length ISO 6360-1	≤ 26 mm (code 4-5)	≤ 34 mm (code 4-5-6 ^{*2})

^{*1} It also covers polishing and prophylaxis tools

^{*2} Intensive use of burs with code 6 can accelerate the wear of the device

4.3 Classification

Class IIa in accordance with European Medical Regulation (EU) 2017/745.

4.4 Performances

Performances	CA PRIMA 1:1 CA PRIMA 1:1 L CA PRIMA 1:1 L MS CA PRIMA 1:121
Speed transmission ratio	1:1

4.5 Operating conditions

Operating conditions			
xx	Temperature range:	[+10°C; +35°C] [+50°F; +95°F]	
X5_X5_X	Relative humidity range:	[30%; 80%]	
(1) (1) (1) (1) (1) (1) (1) (1) (1) (1)	Air pressure range: Barometric pressure range:	[700 hPa; 1060 hPa] [525-795 mmHg]	

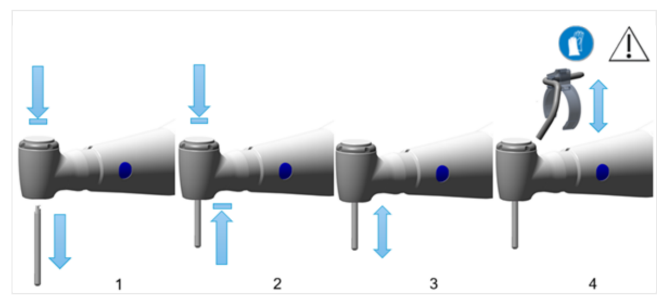


FIG. 5

5 Operation

5.1 Changing the bur

Pictograms used

Sym	Description	Sym	Description
<u> </u>	Movement to the stop in the direction indicated.	1	Back and forth movement.

⚠ CAUTION

- Devices with air-spray must not be used if any open lesions or damaged soft tissue are present or if a recent extraction has taken place. The air flow could propel infected material into the wounds, causing infection and a risk of embolism.
- Never touch soft tissue with the handpiece head. The improper use of the device could lead to burns or injuries.

⚠ CAUTION

• It is essential to use dry, purified compressed air in order to ensure the long working life of the device. Maintain the quality of the air and the water by regular maintenance of the compressor and the filtration systems. The use of unfiltered hard water will lead to early blockage of the tubes, connectors and spray cones.

Contra-angles

FIG. 5

Push-button bur locking system.

- 1. Press the push-button and simultaneously pull out the bur.
- 2. While pressing the push-button insert and rotate the bur inside the chuck system until it fully engages.
- 3. Check that the bur rotates freely and that it is locked by gently pushing and pulling the bur.

For external irrigation version, insert the irrigation system as shown in the picture.

1. To insert and remove the irrigation system, it's recommended to manipulate by holding the plastic clip. Do not pull the metal tube.

6 Cleaning and servicing

6.1 General information

WARNING

- The device is supplied "nonsterile". Clean, dry, lubricate and sterilize the device prior to first use.
- Follow your national directives, standards and guidelines for cleaning and sterilization recommendations.
- Rest the device on a cleanable support to avoid risks of infection for yourself, the patient or third parties.

6.1.1 Precautions for maintenance

- Within a maximum of 30 minutes after each treatment, clean and disinfect the instrument. Observing this procedure eliminates any blood, saliva or residues and prevents the transmission system from being blocked.
- Only use original Bien-Air Dental SA maintenance products and parts or those recommended by Bien-Air Dental SA. For suitable maintenance products refer to section 6.1.2 Suitable maintenance products. Using other products or parts may cause faults during operation and/or void the guarantee.

6.1.2 Suitable maintenance products

Preliminary cleaning:

- Use tap water if the local tap water has pH within the range 6.5 8.5 and chloride content below 100 mg/l. If the local tap water does not meet these requirements, use demineralized (deionized) water instead.
- Aquacare.

Manual cleaning:

• Spraynet®.

Manual disinfection:

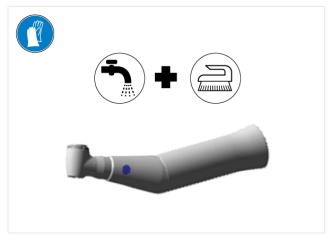
• Alkaline detergent, or detergent-disinfectant (pH 8-11) recommended for cleaning-disinfection of dental or surgical instruments. Disinfectant products composed either of didecyldimethylammonium chloride, quaternary ammonium carbonate or neutral enzymatic product. (e.g. Neodisher® mediclean) are also allowable.

Automatic cleaning-disinfection:

• Use an alkaline or enzymatic product recommended for cleaning in a washer-disinfector for dental or surgical instruments (pH 8-11).

Lubrification:

Lubrifluid®.



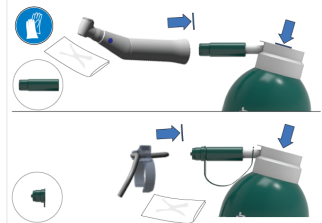


FIG. 6 FIG. 7

6.2 Cleaning

- Do not submerge in physiological liquid (NaCl) nor use saline solution to keep the device moist until it can be cleaned.
- Clean using manual cleaning or automated washer/disinfector only (do not use ultrasonic cleaner).
- Carry out cleaning and sterilization without a bur in the chuck mechanism.
- As with all instruments, following each sterilization cycle, including drying, remove the device to avoid excess exposure to heat which can result in corrosion.
- Use only dynamic sterilizers: do not use a steam sterilizer with a gravity displacement system.

Preparation

- 1. Disconnect the device from the motor and remove the bur (FIG. 5 step 1). Remove the irrigation line then disconnect the device from the electrical motor.
- 2. If the external irrigation system is used, remove the irrigation line.
- 3. If there is a large amount of debris, clean the exterior of the device with disinfectant wipes.

Remove dirt / deposits

- 1. Clean the exterior and interior of the device under tap water at 15°C-38°C (59°F-100°F) provided that the local tap water has a pH within the range of 6.5 8.5 and a chloride content below 100 mg/l. If the local tap water does not meet these requirements, use demineralized (deionized) water instead FIG. 6.
- 2. If external irrigation system is used for surgical procedures, perform preliminary cleaning of the device by using the product Aquacare. Spray the inside and the outside of the device and inside the irrigation tube FIG. 7.

⚠ CAUTION

- Use the Aquacare maintenance pro duct as soon as possible, to remove NaCl deposits.
- If there is a large amount of debris, clean the exterior of the device with disinfectant wipes. Observe the instructions given by the manufacturer.



FIG. 8

6.3 Disinfection

6.3.1 Manual cleaning and disinfection

- 1. Dip the device in a bath containing a cleaning and disinfectant product (e.g. didecyldimethylammonium chloride, quaternary ammonium carbonate or neutral enzymatic product which are allowable chemical agents). Follow the concentration and duration recommended by the fabricant of the disinfection product.
- 2. Brush the device with a smooth, flexible brush (e.g. soft-bristled toothbrush). DO NOT USE a wire brush.
- 3. Optional: perform additional cleaning and disinfection of the external surfaces with non-woven wipes impregnated with a disinfection product (e.g. didecyldimethylammonium chloride).
- 4. Rinse the device twice with running tap water (15°C-38°C) (59°F-100°F) provided that the local tap water has a pH within the range of 6.5 8.5 and a chloride content below 100 mg/l. If the local tap water does not meet these requirements, use demineralized (deionized) water instead.
- 5. After selecting the appropriate nozzle, spray inside the device with Spraynet® (FIG. 8).
- 6. Dry the external surfaces with sterile non-woven compresses (low linting textiles).

6.3.2 Automatic disinfection

Note: The automatic cleaning-disinfection can replace the previous steps 4 to 6.

Washer-disinfector

Carry out automatic cleaning- disinfection using an approved washer-disinfector which complies with ISO standard 15883-1.

Detergent and washing cycle

Use cleaning products (e.g. alkaline detergent or detergent-disinfectant pH 8-11 or neutral enzyme detergent pH 7-8) recommended for washer-disinfector.

Recommended specifications for the thermo-disinfection cycle.

Phase	Parameters
Pre-cleaning	<45°C (113°F); ≥ 2 minutes
Cleaning	45-55°C/113-131°F for enzymatic detergents and 45-65 °C/113-149°F for alkaline detergents ≥ 5 minutes
Neutralization	≥ 2 minutes
Rinsing	Tap water, ≤30°C (86°F), ≥ 2 minutes cold water
Thermal Disinfection	Demineralized water, 90°C — 95°C (194°F — 203°F), 5-10 minutes
Drying	18 — 22 minutes

A CAUTION

Never rinse the devices to cool them.

⚠ CAUTION

If an automatic washer is used instead of the washer/thermo-disinfector, respect the previous program for the Pre-cleaning, Cleaning, Neutralization and Rinsing phases. If the local tap water has a pH outside the range of 6.5-8.5 or if it contains more than 100 mg/l chloride (Cl-ion), do not dry the device inside the automatic washer but dry it manually with low linting textiles.



FIG. 9

6.4 Lubrification

Verifying cleanliness

Visually inspect the device to ensure it is clean. Repeat the cleaning and disinfection procedure if necessary.

Lubrication

Lubricate before each sterilization or at least twice a day. Only the Lubrifluid® spray must be used. FIG. 9

- 1. Place the device in a sterile, non-woven cloth to collect the excess of lubricant.
- 2. Select the appropriate nozzle.
- 3. Insert the nozzle of the Lubrifluid® can in the rear of the device's handle.
- 4. Activate the spray for 1 second and clean the excess oil on the exterior with a sterile, non-woven compress.

6.5 Sterilization

⚠ CAUTION

The quality of the sterilization is highly dependent on how clean the instrument is. Only perfectly clean instruments may be sterilized.

- To improve the effectiveness of the sterilization, make sure the contra-angle is completely dry before and after the sterilization.
- Do not use a sterilization procedure other than the one described below.
- Only use dynamic air removal cycles: pre-vacuum or steam flush pressure pulse (SFPP) cycles.
- If the sterilization is required by national directives, use only dynamic sterilizers: do not use a steam sterilizer with a gravity displacement system.
- As with all instruments, following each sterilization cycle, including drying, remove the device to avoid an excessive exposure to heat which can result in corrosion.
- The bur must be disassembled from the device prior to sterilization.

6.5.1 Procedure

- 1. Pack the device in a packaging approved for steam sterilization.
- 2. Sterilize using steam, following dynamic air removal cycle (ANSI/AAMI ST79, Section 2.19), i.e. air removal via forced evacuation (ISO 17665-1, ISO/TS 17665-2) at 135°C (275°F), for 3 minutes or 132°C (269.6°F) for 4 minutes. In jurisdictions where sterilization for prions is required, sterilize at 135°C (275°F) for 18 minutes.

The device can be used for more than 1000 sterilization cycles.

The recommended parameters for the sterilization cycle are:

- The maximum temperature in the autoclave chamber does not exceed 137°C (278°F), i.e. the nominal temperature of the autoclave is set between 132°C 135.5°C (269.6°F 275.9°F) taking into account the uncertainty of the sterilizer as regards temperature.
- The maximum duration of the interval at the maximum temperature of 137°C (278 °F) is in accordance with national requirements for moist heat sterilization and does not exceed 30 minutes.
- The absolute pressure in the chamber of the sterilizer is 0.07 bar to 3.17 bar (1 PSIA to 46 PSIA/28 Hg to 31 PSIG).
- The rate of change of temperature does not exceed 15°C/min (27°F/min) for increasing temperature and -35°C/min (-63°F/min) for decreasing temperature.
- The rate of change of pressure does not exceed 0.45 bar/min (6.6 psia/min) for increasing pressure and -1.7 bar/min (-25 psia/min) for decreasing pressure.
- No chemical or physical reagents are added to the boiler feedwater.

6.6 Packing and storage

Storage conditions		
x	Temperature range:	[0°C; +40°C] [+32°F; +104°F]
, , , , , , , , , , , , , , , , , , ,	Relative humidity range:	[10%; 80%]
	Air pressure range: Barometric pressure range:	[650 hPa; 1060 hPa] [490 mmHg; 795 mmHg]
*	Keep away from rain	

The device must be stored inside the sterilization pouch in a dry and dust free environment. The temperature must not exceed 55°C (131°F). If the device will not be used for 7 days or more after the sterilization, extract the device from the sterilization pouch and store it in the original package. If the device is not stored in a sterilization pouch or if the pouch is no longer sterile, clean, lubricate and sterilize the device before using it.

↑ CAUTION

If the medical device has been stored refrigerated, allow it to warm up to room temperature prior to removal from its packaging and use.

⚠ CAUTION

Comply with the expiration date of the sterilization pouch which depends on the storage conditions and type of packaging.

6.7 Servicing

Bien-Air Dental SA recommends a regular service for the handpiece after 4000 processing cycles or five years. Never dismantle the device. For all servicing or repair operations, you are advised to contact your usual supplier or Bien-Air Dental SA directly.

7 Transport and Disposal

7.1 Transport

Transpo	ort conditions	
x- X -x-	Temperature range:	[-20°C; +50°C] [-4°F; +122°F]
X5	Relative humidity range:	[5%; 80%]
,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	Air pressure range: Barometric pressure range:	[650 hPa; 1060 hPa] [490 mmHg; 795 mmHg]
*	Keep away from rain	

7.2 Disposal



The disposal and/or recycling of materials must be performed in accordance with the local, national or international regulations.

All contra-angles and handpieces must be recycled. In order to avoid any risk of contamination, the user must return the device sterilized to their dealer or contact an authorized body for the treatment and recovery of this type of equipment.

8 General Information

8.1 Terms of guarantee

Bien-Air Dental SA grants the user a warranty covering any operating fault, or material or manufacturing defect.

The warranty period is 24 months from the date of invoicing.

In the event of a justified claim, Bien-Air Dental SA or its authorized representative will repair or replace the product free of charge.

All other claims of any kind whatsoever, particularly claims for damages, are excluded.

Bien-Air Dental SA cannot be held liable for damage or injury and the consequences thereof, resulting from:

- Excessive wear and tear,
- · Infrequent or improper use,
- Failure to observe the servicing assembly or maintenance instructions,
- Damage caused by unusual chemical, electrical or electrolytic influences.
- Faulty air or water connection.

⚠ CAUTION

The warranty becomes null and void if damage and its consequences result from incorrect servicing or modification by third parties not authorized by Bien-Air Dental SA. Warranty requests will only be taken into consideration if the product is accompanied by a copy of the invoice or delivery note. The following information must be clearly indicated: purchase date, product reference and serial number.

8.2 References

REF	Legend	Ratio
1601235-001	CA PRIMA 1:1	•
1601236-001	CA PRIMA 1:1 L*	•
1601237-001	CA PRIMA 1:1 L MS*	•
1601267-001	CA PRIMA 1:121	•
1600036-006	Spraynet®, cleaning spray 500 ml, box of 6 cans	
1600064-006	Lubrifluid®, lubricant 500 ml, box of 6 cans	
1600617-006	Aquacare, cleaning spray for physio- logical liquid 500 ml, box of 6 cans	
1500431-001	External irrigation system	
1500984-010	Pack of 10 disposable sterile lines	_

^{*}with light



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