

Chiropro Plus 3rd Gen ENG INSTRUCTIONS FOR USE.







Set Chiropro Plus 3rd Gen KM REF 1700739-001

















REF 1600994-001

REF 1303393-001

- REF 1600755-001 F
 - REF 1601069-001
- REF 1600631-001
- REF 1501635-010

REF 1307727-010



REF 1301575-001

REF 1502329-002

Set Chiropro Plus 3rd Gen KM CA 20:1L KM REF 1700738-001

-Ď



REF 1700739-001

Set Chiropro Plus 3rd Gen CA 20:1 L KM JAPAN REF 1700773-001



Set Chiropro Plus 3rd GEN CA20:1 L KMWL JAPAN REF 1700908-001





*For instructions for use of the wireless foot control, please refer to the Quick Guide REF 2100443.

Table of content

1.	Sy	rmbols6
	1.1	Description of symbols for Chiropro Plus 3rd
	Gen	units 6
	1.2	Description of symbols for Chiropro Plus 3rd
	Gen	accessories 7
2.	lde	entification, Intended Use and
N	otat	ion 8
	2.1	Identification 8
	2.2	Intended use 8
	2.3	Intended patient population 8
	2.4	Intended User 8
	2.5	Intended medical conditions 8
	2.6	Patient contra-indications and warnings 8
	2.7	In case of accidents
	2.8	Notation and chapter links
3.	Us	ser and patient safety: Warnings &
Ρı	reca	utions of Use 10
4.	De	escription
	4.1	Chiropro Plus 3rd Gen system overview 13
	4.2	Sets supplied14
	4.3	Options 17
	4.4	Technical data
	4.5	Performance 22
	4.6	Environmental protection and information
	for o	disposal
	4.7	Electromagnetic compatibility (technical
	des	cription)-Emissions & Immunity 24
5.	Ins	stallation 27
	5.1	Install the Chiropro Plus 3rd Gen system . 28
	5.2	On/off procedure 29
6.	Int	terface overview

	6.1	Chiropro Plus 3rd Gen modes	30
	6.2	Rotating knob functions overview	31
	6.3	Sound alerts	32
7.	Op	peration	33
	7.1	Operation screen description	33
	7.2	Perform an operation, steps P1 and P2	34
	7.3	Perform an operation, steps P3, P4 and	
	P5.		35
8.	Op	eration – Surgery mode	37
	8.1	Operation screen description	37
	8.2	Perform an operation	37
9.	Se	ttings	39
	9.1	Operation mode	39
	9.2	MX-i LED micromotor speed	39
	9.3	MX-i LED micromotor torque	41
	9.4	MX-i LED micromotor rotation direction	41
	9.5	Irrigation level	42
	9.6	Contra-angle ratio	42
	9.7	Luminosity level	42
1(). Sp	ecial modes	44
11	I. Lis	st of errors & Troubleshooting	47
	11.1	Alert notifications (operating)	47
	11.2	Device operating error	48
12	2. M a	aintenance	49
	12.1	Servicing	49
	12.2	Sterilization	50
	12.3	Important	51
	12.4	Replacement of fuses	52
13	3. Gu	arantee	53
	13.1	Terms of guarantee	53

ENG INSTRUCTIONS FOR USE

1 Symbols

1.1 Description of symbols for Chiropro Plus 3rd Gen units

Sym	Description	Sym	Description
CE 0123	CE Marking with number of the notified body.	R P	General symbol for recovery/ recyclable.
\bigcirc	OFF (power).	X	Separate collection of electric and electronic equipment.
	ON (power).	***	Manufacturer.
⊕	Fuse.	-¤̈́-	Lamp; lighting; illumination.
\sim	Alternating current.		Sound alerts.
((•))	Non-ionizing electromagnetic radiation.	Rx Only	Warning: in accordance with federal law (USA), this device is only available for sale upon recommendation by an accredited practitioner.
Â	CAUTION! hazard that could result in light or moderate injury or damage to the device if the safety instructions are not correctly followed.	, [®] ^s	CSA marking - Complies with U.S and Canadian standards.
	WARNING! hazard that could result in serious injury or damage to the device if the safety instructions are not correctly followed.	SN	Serial number.
6	Refer to instruction manual/booklet (<u>https://dental.bienair.com/fr_ch/</u> <u>support/download-center/</u>).	REF	Catalogue number.
EC REP	Authorized EC Representative in the European Community.	MD	Medical Device.
	Data Matrix code for product information including UDI (Unique Device Identification).	Å	Equipotentiality.

1.2 Description of symbols for Chiropro Plus 3rd Gen accessories

Sym	Description	Sym	Description
CE	CE Marking with number of the notified body.	Ă]	Thermo washer disinfectable.
\square	Expiration date.	R A	General symbol for recovery/ recyclable.
2	Do not reuse.	X	Separate collection of electric and electronic equipment.
para	Sterilized with Ethylene Oxyde.	135℃ ∭	Sterilizable in autoclave up to the specific temperature.
¥	Electrical safety. Applied part type B.	••••	Manufacturer.
REF	Catalogue number.	SN	Serial number.
DEHP	Does not contain DEHP.	LOT	Batch code.
	Do not use if package is damaged.		

2 Identification, Intended Use and Notation

2.1 Identification

The Chiropro Plus 3rd Gen device encompasses a table-top system for dental implantology and oral surgery allowing to control a dental micromotor which drives a dental handpiece. A peristaltic pump conveys the physiological liquid via a sterile single-use irrigation line. The console includes a single knob control to set the parameters and a foot control used to turn on/off the pump, to navigate through the various steps of the selected procedure and to control the rotation direction of the motor. The device's LCD display shows many parameters of the operation, such as the handpiece gear ratio, bur speed, torque value and irrigation flow setting.

2.2 Intended use

All Chiropro Plus 3rd Gen devices are intended to be used in dental implantology and oral surgery. The consoles are designed to operate a specific dental micromotor that drives dental handpieces fitted with appropriate tools to cut hard and soft tissues in the mouth and to screw dental implants. The intended electromagnetic environment (per IEC 60601-1-2 ed. 4.0) is Professional healthcare facility environment.

2.3 Intended patient population

The intended patient population of the Chiropro Plus consoles includes any person visiting a dental practitioners' 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

The Chiropro Plus 3rd Gen is meant to be used only by dentists and dental surgeons in dental offices and hospitals.

2.5 Intended medical conditions

Dental implantology is the elective treatment to replace one or more missing teeth. Teeth can be missing for various reasons, such as traumas, partial or total edentulism, and advanced decay that leads to tooth sacrifice because restorative treatments are no longer possible.

Dental implantology requires to prepare jawbone to accommodate a dental implant, which is typically a titanium screw fitted with an abutment and a prosthetic crown made of ceramic material mimicking the natural missed tooth.

Multi-teeth prosthetic solutions are also available, usually supported by more than one single implant.

2.6 Patient contra-indications and warnings

No specific contra-indication exists for the Chiropro Plus device family when the device is used as intended.

2.7 In case of accidents

If an accident occurs, the Chiropro Plus 3rd Gen must not be used until repairs have been completed by a qualified and trained technician authorized by the manufacturer.

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.

2.8 Notation and chapter links

• A, B, C, etc.

Text preceded by a letter indicates a procedure to be carried out step-by-step.

✤ Indicates a procedure result.

• (1), (2), (3), etc.

Text preceded by a number indicates text used in

conjunction with an illustration.

• OK , Settings, etc.

Text in bold italic font style indicates, on-screen elements such as buttons, menus, menu items, screen areas, values, fields when they are named and screen names.

In order to simplify the notation, in this manual:

- "Clockwise" is referred to as "CW";
- "Counterclockwise" is referred to as "CCW";
- Forward micromotor rotation mode is referred to as "FWD";
- Reverse micromotor rotation mode is referred to as "REV";
- Rotational speed unit "revolutions per minute" is referred to as "rpm";
- Torque unit "newton centimetre" is referred to as "Ncm";
- Micromotor control unit is referred to as "DMX".

D Z Z

3 User and patient safety: Warnings & Precautions of Use

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

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

A WARNING

According to IEC 60601-1:2005+A12012/AnnexG, electrified devices (motors, control units, couplers and attachments), can be safely used in a medical environment in which potentially explosive or flammable mixtures of anaesthetic substances are delivered to the patient only if:

- The distance between the motor and the anaesthetic breathing circuit exceeds 25 cm.
- The motor is not used simultaneously to the administration of the anaesthetic substances to the patient.

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

- The device must be connected only to a supply main with a protective earth.
- Always ensure that there is no water under the unit before switching it on.
- The cleaning procedure defined in chapter 11.2 Cleaning & Sterilization on page 35 must be followed.
- All connectors must be dry before use. Ensure the absence of residual moisture due to cleaning.
- Never simultaneously touch the patient and the electrical connection of the unit. The system must never be touched by the patient.
- Never attempt to open the device while it is connected to the electrical mains.
- The power plug must be always easily accessible as it may be used for disconnection in case of problems.

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

\triangle warning

- To avoid any risk of contamination, only control the device via the foot control during surgical procedures. Never touch the device during a clinical operation.
- The cleaning procedure of the device defined in chapter 11.2 Cleaning & Sterilization on page 35 must be followed.
- The cleaning and sterilization procedure of the knob defined in chapter 11.2 Cleaning & Sterilization on page 35 must be followed.
- Always refer to the accessories IFU for dedicated maintenance procedure.
- Always replace the sterile protective sheet after an operation as they are single use only.
- Always replace the irrigation line after an operation as they are single use only.
- Always ensure that the irrigation line package is intact before use.

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

⚠ WARNING

- If the irrigation pump is used and regulated by the device, verify that the pump is working properly before starting the treatment as well as during the treatment. The device has neither been conceived for controlling the working status of the pump nor for detecting possible failures of the pump.
- There is no detection of empty physiological liquid flask. Always check the content of the flask before operating.
- Never run the pump without the irrigation line being securely fastened.

To prevent any risk of console or motor overheating, the cautions below must be observed:

\triangle caution

- Always ensure that both the cable and the motor are in good condition.
- Ensure that the micromotor hose is not bent.
- Do not use the device outside the range of operating temperature.
- Let the system cool down when the motor overheating alert notification is on display.
 See chapter 10.1 Alert notifications on page 33 for more details.

To prevent any risk of injury (damage to bone, teeth, tissue) the cautions below must be observed:

\triangle caution

- The predefined settings contained in the device are indicative only. Bien-Air Dental SA cannot be held liable for them.
- The predefined torque and speed values are only intended as a guide. The drill values used must be adapted according to the implant manufacturer instructions. Always refer to the implant manufacturer specifications to set up the console settings.
- Always verify that the configured parameters correspond to your medical application. The predefined parameters may be subject to modification without notice.

To prevent any risk of adverse tissue reaction, the caution below must be observed:

\triangle caution

• If the irrigation pump is used, only use biocompatible irrigation line recommended by the manufacturer and refer to the recommendation of the pump manufacturer.

To prevent any wireless connection loss, the cautions below must be observed:

\triangle caution

- Ensure that there are no obstructions, such as clutter, furniture, or other items, between the foot control and the console. The foot control and the console must always be in the same room.
- The foot control emits a red signal when the battery is low. In this case, it is recommended to complete the ongoing operation and change the batteries before starting a new operation.

To prevent any risk of injury and/or material damage the warnings/ cautions below must be observed:

- Place the device on a flat surface capable of bearing its weight. It may be positioned on a table, on a trolley or any other surface but in no circumstances on the floor.
- Always use Bien-Air Dental SA accessories or those recommended by Bien-Air Dental SA.

\triangle caution

- Never connect a handpiece on a running micromotor.
- Do not switch off the device while the motor is running.
- Always check that the lid is not opened when running the irrigation pump.
- Beware of the risk of pinching when closing the irrigation valve.
- Only use original Bien-Air Dental maintenance products and parts or those recommended by Bien-Air Dental. Using other products or parts may cause operational failure and/or void the guarantee.

ENG

To avoid any risk of electromagnetic interference that could affect active implantable medical devices, and sustainable life devices, the warnings below must be observed:

- The device must not be placed in the vicinity (30cm) of other sustainable life devices.
- Dental professionals need to be aware of potential electromagnetic interference between electronic dental devices and active implantable medical devices and should always inquire about any devices implanted in the patient.
- 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 device is not intended to be used in the vicinity of High Frequency surgical equipment.

To avoid any risk of electromagnetic interference that could affect the performance of the device the warnings below must be observed:

\triangle warning

- Since compliance with the international standard IEC 60601-1-2 does not guarantee immunity against 5G worldwide (due to the different frequency bands used locally), avoid the presence of devices equipped with 5G broadband cellular networks in the clinical environment or ensure that the network functionality of these devices is disabled during the clinical procedure.
- Radio transmitting equipment, cellular phones, etc., should not be used in the immediate vicinity of the device, since this could affect its operation. Special precautions should be taken when using strong emission sources such as highfrequency surgical equipment and other similar devices, to ensure that HF cables are not routed above or near the device. If in doubt, please contact a qualified technician or Bien-Air.
- 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 specified by the manufacturer. Otherwise, degradation of the performance of this equipment could result.
- The use of accessories, transducers and cables other than those specified, with the exception of transducers and cables sold by Bien-Air, may result in increased emissions or decreased immunity.

4 Description

4.1 Chiropro Plus 3rd Gen system overview



FIG. 1

- (1) Peristaltic pump lid
- (2) Foot control connector
- (3) Marking
- (4) Bracket support
- (5) Main switch
- (6) Fuse box
- (7) Mains connector
- (8) Potential equalization connector
- (9) MX-i LED micromotor

- (10) Button to start/stop irrigation
- (11) Button to reverse the rotation of the MX-i LED micromotor
- (12) "Program" button to go to next operation step
- (13) Motor start
- (14) MX-i LED micromotor connector
- (15) Control knob
- (16) LCD control screen

4.2 Sets supplied

Chiropro Plus 3rd Gen set REF 1700710-001

Designation	REF number
Chiropro Plus 3 rd Gen unit (1x)	1600994-001
MX-i LED micromotor (1x)	1600755-001
3-button foot control (1x)	1600631-001
Cable MX-i LED (2m) (1x)	1601069-001
Sterile protective sheet (2x)	1502329-002
Pack of 5 disposable sterile irrigation lines	1500984-005
Pack of 10 attachment collars for fastening the sterile irrigation line to a cable	1307727-010
Bracket for fluid bottle (1x)	1303393-001
Handpiece support (1x)	1301575-001

Chiropro Plus 3rd CA 20:1 L WL set REF 1700891-001

Designation	REF number
Chiropro Plus 3 rd Gen unit (1x)	1600994-001
MX-i LED micromotor (1x)	1600755-001
Wireless foot control +Dongle (1x)	1601192-001
Contra-angle handpiece CA 20:1 L Micro- Series (light) (1x)	1600692-001
Cable MX-i LED (2m) (1x)	1601069-001
Sterile protective sheet (2x)	1502329-002
Pack of 5 disposable sterile irrigation lines	1500984-005
Pack of 10 attachment collars for fastening the sterile irrigation line to a cable	1307727-010
Bracket for fluid bottle (1x)	1303393-001
Handpiece support (1x)	1301575-001

Chiropro Plus 3rd Gen CA 20:1 L set REF 1700709-001

Designation	REF number
Chiropro Plus 3 rd Gen set (1x)	1700710-001
Contra-angle handpiece CA 20:1 L Micro- Series (light) (1x)	1600692-001

Chiropro Plus 3rd Gen CA 1:2.5 L set REF 1700751-001

Designation	REF number
Chiropro Plus 3 rd Gen set (1x)	1700710-001
Contra-angle handpiece CA 1:2.5 L Micro- Series (light) (1x)	1601055-001

Chiropro Plus 3rd Gen KM set REF 1700739-001

Designation	REF number
Chiropro Plus 3 rd Gen unit (1x)	1600994-001
MX-i LED micromotor (1x)	1600755-001
3-button foot control (1x)	1600631-001
Cable MX-i LED (2m) (1x)	1601069-001
Sterile protective sheet (2x)	1502329-002
Kirschner/Meyer pack of 10 disposable sterile lines	1501635-010
Pack of 10 attachment collars for fastening the sterile irrigation line to a cable	1307727-010
Bracket for fluid bottle (1x)	1303393-001
Handpiece support (1x)	1301575-001

Chiropro Plus 3rd Gen CA 20:1 L KM set REF 1700738-001

Designation	REF number
Chiropro Plus 3 rd Gen KM set (1x)	1700739-001
Contra-angle handpiece CA 20:1 L KM Micro- Series (light) (1x)	1600786-001

Chiropro Plus 3rd Gen CA 20:1 L KM JAPAN set REF 1700773-001

Designation	REF number
Chiropro Plus 3 rd Gen set (1x)	1700710-001
Contra-angle handpiece CA 20:1 L KM Micro- Series (light) (1x)	1600786-001

Chiropro Plus 3rdCA20:1 L KMWL JAPAN set REF 1700908-001

Designation	REF number
Chiropro Plus 3 rd Gen unit (1x)	1600994-001
MX-i LED micromotor (1x)	1600755-001
Wireless foot control +Dongle (1x)	1601192-001
Contra-angle handpiece CA 20:1 L KM Micro- Series (light) (1x)	1600786-001
Cable MX-i LED (2m) (1x)	1601069-001
Sterile protective sheet (2x)	1502329-002
Pack of 5 disposable sterile irrigation lines	1500984-005
Pack of 10 attachment collars for fastening the sterile irrigation line to a cable	1307727-010
Bracket for fluid bottle (1x)	1303393-001
Handpiece support (1x)	1301575-001

4.3 Options

Designation	REF number
3-button foot control	1600631-001
Wireless foot control +Dongle <u>*</u>	1601192-001
Sterile protective sheet	1502329-002
Pack of 10 disposable sterile lines 3.5 m	1501738-010
Kirschner/Meyer pack of 10 disposable sterile lines	1501635-010
Kirschner/Meyer type detachable irrigation set for CA 20:1 L KM Micro- Series, comprising 10 rings and 10 tubes	1501621-010
Pack of 10 disposable sterile lines	1500984-010
Bracket for fluid bottle	1303393-001
Handpiece support	1301575-001
Cable MX-i LED (2m)	1601069-001
Pack of 10 attachment collars for fastening the sterile irrigation line to a cable	1307727-010
Pack of 10 fuses T4.0AH 250 VAC high breaking capacity	1307312-010
Knob	1307031-001

4.4 Technical data

Dimensions L x W x H	
Chiropro Plus 3 rd Gen unit	240 x 240 x 102 mm
Chiropro Plus 3 rd Gen unit (with bracket)	240 x 240 x 482 mm
Foot control (without handle)	206 x 180 x 60 mm
Foot control (with handle)	206 x 200 x 155 mm
Wireless Foot control Width x Height x Depth (without hook)	206 x 180 x 60 mm
Wireless Foot control Width x Height x Depth (with hook)	206 x 200 x 155 mm
Motor cable (REF 1601069)	L 2.0 m
Foot control cable	L 2.9 m
MX-i LED micromotor	23 x 91 mm

Weight	
Chiropro Plus 3 rd Gen unit	2.44 kg
Foot control (without handle and cable)	830 g
Foot control (with handle and cable)	877 g
Wireless Foot control (without hook, two batteries included)	934 g
Bracket	115 g
Cable	105 g
MX-i LED micromotor	110 g

Electrical data	
Voltage	100 – 240 VAC
Frequency	50-60 Hz

Environmental conditions

Storage	
Temperature range:	0°C/+40°C
Relative humidity range:	10% - 80%
Air pressure range:	650 hPa – 1060 hPa
Transport	
Temperature range:	-20° C / + 50° C
Relative humidity range:	5% - 80%
Air pressure range:	650 hPa – 1060 hPa
Operating temperature	
Temperature range:	+ 5° C / + 35° C
Relative humidity range:	30% - 80%
Air pressure range:	700 hPa – 1060 hPa

${\rm \ensuremath{\underline{\wedge}}}$ caution

Do not use Chiropro Plus 3rd Gen outside the range of operating temperature.

Classification

Class IIa in accordance with European Regulation (EU) 2017/745 concerning medical devices.

Electric insulation class

Class I per IEC 60601-1 (apparatus protected against electric shocks).

\bigtriangleup caution

Wireless foot control

The device must be only used by the operator.

Applied parts (per IEC 60601-1):	
MX-i LED micromotor REF 1600755-001	
CA 20:1 L Micro-Series REF 1600692-001	
CA 20:1 L KM Micro-Series REF 1600786-001	
Irrigation lines REF 1500984-010	
KM Irrigation lines REF 1501635-010	
Degree of ingress protection	
Unit	IP 41 (protection against insertion of objects larger than 1 mm and dripping water (vertically falling drops)).
Foot control	IP X8

IP X6

Memory

Memory storage of 5 steps settings including adjustment of speed, torque, rotation direction, irrigation, and contra-angle ratio for each step.

Languages

English.

Bracket for physiological liquid flask

Stainless steel.

Peristaltic pump		
Pump delivery From 30 to 130 ml/min. (5 levels		
Irrigation line	External Ø 5.60 mm	
	Internal Ø 2.40 mm	
Wall thickness	1.60 mm	
Intended for use with:	See instructions for use	
MX-i LED micromotor	REF 2100245	
Cable MX-i LED	REF 2100163	
Contra-angle CA 20:1 L Micro-Series, light	REF 2100209	
Contra-angle CA 20:1 L KM Micro-Series, light	REF 2100209	
Micro-Series, light	REF 2100337	

\triangle caution

The use of the system with other handpieces, motors or cables has not been validated/certified (speed and torque values are not guaranteed in this case).

List of errors & Troubleshooting

See chapter "10 List of errors & Troubleshooting".

4.5 Performance

Performance	REF 1600994	
Motor speed regulation	Accuracy ± 5% in the speed range 100 - 40'000 rpm (*)	
Motor torque regulation	Torque adjustable from 10% to 100% of the maximum torque	
Maximum motor torque	5 (±5%) Ncm (*)	
Maximum motor power	95 (±10%) W (*)	
Max motor LED current	250 (± 10%) mA rms	
Max motor LED current range	Not adjustable, always at full intensity	
Power supply output limitation	< 300 W	
Irrigation flow	5 levels:	
	1 drop = 30ml/min	
	2 drops = 60ml/min	
	3 drops = 90ml/min	
	4 drops = 120ml/min	
	5 drops = 150ml/min	

(*) Measurement realized in combination with motors MX-i LED 3rd Gen 1601008 and MX-i LED 1600755, contraangle CA 20:1 L Micro Series 1600692 and/or handpiece PML 1121 1600156. The maximum torque is measured at 1000 rpm with irrigation stopped and it corresponds to a maximum torque of 80 Ncm at the rotative tool if the motor is combined with the contra-angle CA 20:1 L Micro Series 1600692.

In accordance with IEC 80601-2-60, no essential performance is linked to this dental equipment

4.6 Environmental protection and information for disposal



The disposal and/or recycling of materials must be performed in accordance with the legislation in force.



Separate collection of electric and electronic equipment and accessories in view of recycling. Electrical and electronic equipment may contain dangerous substances which constitute health and environmental hazards.

The user must return the device to its dealer or establish direct contact with an approved body for treatment and recovery of this type of equipment (European Directive 2012/19/EU).

4.7 Electromagnetic compatibility (technical description)-Emissions & Immunity

This electronic control is in compliance with electrical safety standards in line with standard IEC 60601-1-6, third edition, and those governing electromagnetic compatibility in line with standard IEC 60601-1-2, fourth edition.

Guidance and manufacturer's declaration – Electromagnetic emissions

The Chiropro Plus 3rd Gen is intended for use in the electromagnetic environment specified below. The customer or the user of the Chiropro Plus 3rd Gen must ensure that it is actually used in such an environment.

Emissions test	Compliance	Electromagnetic environment - guidance
RF emissions CISPR 11	Group 1	The Chiropro Plus 3 rd Gen uses RF energy for its internal operation only. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.
RF emissions CISPR 11	Class B	The Chiropro Plus 3 rd Gen is
Harmonic emissions IEC 61000-3-2	Class A	suitable for use in any building including residential buildings and those directly connected to the
Emissions due to voltage fluctuations IEC 61000-3-3	Conforming	public low-voltage power supply network that supplies buildings used for residential purposes.

Guidance and manufacturer's declaration – Electromagnetic immunity

The Chiropro Plus 3rd Gen is intended for use in the electromagnetic environment specified below. The customer or the user of the Chiropro Plus 3rd Gen must ensure that it is actually used in such an environment.

Immunity test	IEC 60601 test level	Compliance level	Electromagnetic environment - guidance
Electrostatic discharge (ESD) IEC 61000-4-2	±8 kV contact ±2 kV air ±4 kV air ±8 kV air ±15 kV air	±8 kV contact ±2 kV air ±4 kV air ±8 kV air ±15 kV air	Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30%.
Electrical fast transient/burst IEC 61000-4-4	±2 kV for power supply lines ±1 kV for other lines	±2 kV for power supply lines N.A.	Mains power quality should be that of a commercial or hospital environment.
Surge IEC 61000-4-5	±0.5 kV line to line ±1 kV line to line ±0.5 kV line to earth ±1 kV line to earth ±2 kV line to earth	±0.5 kV line to line ±1 kV line to line ±0.5 kV line to earth ±1 kV line to earth ±2 kV line to earth	Mains power quality should be that of a commercial or hospital environment.
Voltage dips, short interruptions and voltage variations on power supply input lines IEC 61000-4-11	0% UT for 0.5 cycle, at 0°, 45°, 90°, 135°, 180°, 225°, 270° and 315° 0% UT for 1 cycle and 70% UT for 25/30 cycles at 0° 0% UT for 250 cycles at 0°	0% UT for 0.5 cycle, at 0°, 45°, 90°, 135°, 180°, 225°, 270° and 315° 0% UT for 1 cycle and 70% UT for 25/30 cycles at 0° 0% UT for 250 cycles at 0°	Mains power quality should be that of a commercial or hospital environment. If the user of the Chiropro Plus 3 rd Gen requires continued operation during mains power interruptions, it is recommended that the Chiropro Plus 3 rd Gen be powered from an uninterruptible power supply or a battery.
Magnetic field due to mains frequency (50/60 Hz) IEC 61000-4-8	30 A/m	30 A/m	Magnetic fields generated by the mains frequency should be at levels characteristic of a typical location in a typical commercial or hospital environment.
Conducted disturbances induced by RF fields IEC 61000-4-6	3 VRMS 0,15 MHz – 80 MHz 6 VRMS in ISM bands 0,15 MHz – 80 MHz 80% AM at 1 kHz	3 VRMS 0,15 MHz – 80 MHz 6 VRMS in ISM bands 0,15 MHz – 80 MHz 80% AM at 1 kHz	Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey ^a should be less than the compliance level in each frequency range. Interference may occur in the vicinity of
Radiated RF EM fields 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	symbol:

Immunity test	IEC 60601 test level	Compliance level		Electromagnetic environment - guidance
Proximity fields from RF wireless communications equipment IEC 61000-4-3	Test freq. [MHz]	Max. power [W]	Immunity test level [V/m]	
	385	1.8	27	Distance: 0.3 m
	450	2	28	
	710,745,780	0.2	9	
	810,870,930	2	28	
	1720,1845,1970	2	28	
	2450	2	28	
	5240,5500,5785	0.2	9	
NOTE: UT is the AC mains voltage prior to application of the test level.				

a. Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and mobile field radios, amateur radios, AM and FM radio broadcasts and TV broadcasts 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 Chiropro Plus 3rd Gen is used exceeds the RF compliance level mentioned above, the Chiropro Plus 3rd Gen should be observed to verify that it is operating normally. If abnormal operation is observed, additional measures may be necessary, such as reorienting or relocating the Chiropro Plus 3rd Gen.

5 Installation









FIG. 2

FIG. 3

FIG. 4

FIG. 5



FIG. 6



FIG. 8







FIG. 10



FIG. 11



FIG. 12

28

5.1 Install the Chiropro Plus 3rd Gen system

FIG. 1

A. Place the Chiropro Plus 3rd Gen on a flat surface capable of bearing its weight.

FIG. 2

B. The fuse box may be opened with a screwdriver. 100 - 240 VAC = fuse T4.0AH 250 VAC REF 1307312-010.

To replace a fuse, see chapter "11.4 Replacement of fuses". **C.** Connect the power cable (1) to the connector (2).

Note :

FIG. 3

D. Connect the foot control cable to the input provided on the rear panel, guiding the connector and plug by means of the index pin on the connector.

\triangle caution

- Do not lift the foot control holding the connection cable.
- To disconnect the foot control cable, pull the cable socket connector (1). Do not pull the cable (2) without disconnecting the cable socket before.
- If a wireless foot control is used, please follow the IFU/Quick Start Guide REF. 2100443.

FIG. 4

E. Connect the MX-i LED micromotor cable to the motor output, guiding the connector and plug by means of the index pin on the connector.

FIG. 5

F. Align and attach the bracket to the housing provided on the rear of the console and suspend the flask or bottle.

G. Check the packaging integrity, as well as the expiry date of the irrigation line on the label (1).

H. Remove the single-use sterile irrigation line (2) from its pouch.

FIG. 7

I. Connect the flexible hose of the irrigation line to the spray tube of the handpiece or contra-angle.

FIG. 8

J. Install the peristaltic cassette (1) in the peristaltic pump (2).

Check that the cassette is clipped correctly.

FIG. 9

K. Close the pump lid (3). If there is resistance to closing, open the lid again and check the correct positioning of the cassette. When the lid is correctly closed, the user should hear a click sound.

FIG. 10

L. Perforate the cap of the physiological liquid flask with the pointed end of the irrigation line after removing the protective cap.

FIG. 11

M. Attach the irrigation line on the motor cable using the 3 attachment collars REF 1307727-010.

5.2 On/off procedure

The device can be switched on and off in complete safety using the main switch on the Chiropro Plus 3rd Gen.

Note : The equipment is powered by the mains power supply (100 - 240 VAC / 300VA / 50-60Hz).



FIG. 2

6 Interface overview

6.1 Chiropro Plus 3rd Gen modes

The Chiropro Plus 3rd Gen allows to visualize and control operation parameters by the means of the LCD display.

A unique screen allows to use the following modes: FIG. 1

• Operation mode (to perform an operation in 3 steps) See chapter "7 Operation" on page 26 for details. FIG. 2

• Settings mode (to set up operation parameters)

See chapter "9 Settings" on page 29 for details. FIG. 3

• Special modes (to test system and reset settings)

See chapter "10 Special modes" on page 31 for details. FIG. 4

A. Long press on the rotating knob (1) to switch between Operation and Settings modes.

Note : See instructions for use

See chapter "6.2 Rotating knob functions overview" below for details. See chapter "10 Special modes" on page 31 for entering special modes.

6.2 Rotating knob functions overview

Note : Any knob or foot control action will be ignored when the motor is running.		
Knob action	Description	
CW rotation	Increase current value, go to the element on the right	

Note : Any knob or foot control action will be ignored when the motor is running

CW rotation	Increase current value, go to the element on the right
CCW rotation	Decrease current value, go to the element on the left
One short press (Operation mode)	Go to the next programmed step, acknowledge error messages
One short press (Settings mode)	Enter selected setting, validate and store the current setting value, exit the current setting, acknowledge error messages
One long press	Switch between Operation and Settings modes
Double short press	Enter special modes (only when gear ratio is selected in settings mode)





FIG. 4

6.3 Sound alerts

Sound alert	Description
One short beep	Activating irrigation, going to next step, and switching rotation direction to FORWARD
Two short beeps	Deactivating irrigation, and switching rotation direction to REVERSE
Two long beeps	Switching from low speed to high speed programmed step
Alternate short beeps	Alert notifications
Alternate medium beeps	Micromotor REVERSE running indicator
Alternate long beeps	System failure notification

Note : The Operation mode is the default startup mode.

Any knob or foot control action will be ignored when the motor is running.

ENG



FIG. 2

7 Operation

7.1 Operation screen description

FIG. 1

The Operation screen differs whether the micromotor is stopped or running and depending on the active step.

It allows to perform an operation in 3, 4 or 5 predefined steps P1, P2, P3, P4, P5 (which can respectively be used to program settings for the bone preparation, drilling, threading and implant insertion phases), and displays the following information:

(1) Step P1 (inactive step, in black)

(2) Step P2 (inactive step, in black)

(3) Step P3 (active step, in green)

P4 and P5 steps are disabled by default, see " Number of steps", chapter "10 Special modes" on page 31 for enabling them.

(4) Speedometer

Note : Real-time speed value is displayed in black when the MX-i LED micromotor is running. Maximum reachable speed value stored is displayed in cyan when the MX-i LED micromotor is not running, in steps P1 and P2

(5) Torquemeter

Note : Torquemeter is only displayed when micromotor speed is below 100 RPM in steps P1 and P2. (6) Contra-angle ratio

Note : The contra-angle ratio is cyan-colored for direct-drive and green-colored for reduction gears. (7) Bar graph for torque

Note : Torque bar graph is only displayed when micromotor speed is below 100 RPM.
(8) Operation settings symbols
See chapter "9 Settings" on page 29 for details on adjusting settings.

7.2 Perform an operation, steps P1 and P2

FIG. 2

A. Operate by pressing the foot control to adjust the MX-i LED micromotor speed.

- \mathbf{k} Inactive steps symbols turn off when the motor is running.
- Speedometer displays real-time speed value in black.

Note : Each step settings are restored from the corresponding step last used settings, excluding quick settings made directly in the Operation mode.

In REVERSE mode, the rotation direction symbol 🖤 blinks and there is a sound alert (alternate medium beeps). The Torque value is automatically increased in REVERSE mode when torque-meter is displayed. The torque value can be increased from 0 to 10 Ncm, see "Reverse torque boost value", chapter "10 Special modes" on page 31 to adjust it.

Actions on foot control's buttons have no effect when the micromotor is running. FIG. 3

B. If necessary, release the foot control to perform the following actions:

- Speedometer (1) displays the set micromotor maximum reachable speed in cyan.
- Turn the knob CW or CCW to respectively increase or decrease the micromotor maximum reachable speed (quick setting mode).

Note : Changes made in this mode (either by rotating the knob or by changing parameters through the foot control buttons) are considered as temporary settings and are never saved.

🔖 The speedometer is cyan and displays the set micromotor maximum reachable speed (1).

Note : Changing the torque in steps P1 or P2 can only be performed through the Settings mode

• Long press on the knob to change operation settings.

🔖 The Settings mode is displayed.

See chapter "9 Settings" on page 29 for details.

• Long press on the orange button to activate the 5 Ncm torque boost.

Note : The torque boost can only be activated when the torquemeter is displayed in Operation mode, in low speed steps (<100 RPM).

- C. Short press on the foot control's orange button or on the knob to go to the next step.
- 🔖 The next step symbol turns green and the step's last used settings are restored.

Note : Actions on foot control's buttons have no effect when the micromotor is running.

Changing the torque in steps P1 or P2 can only be performed through the Settings mode.

The torque boost can only be activated when the torquemeter is displayed in Operation mode, in low speed steps (<100 RPM).

For safety reasons, the speed setting icon turns red and blinks together with the speedometer for 2 seconds when switching from low speed to high speed (=100 RPM) step.

7.3 Perform an operation, steps P3, P4 and P5

FIG. 4

- A. In steps P3 (1), P4 and P5, operate by pressing the foot control to adjust the MX-i LED micromotor speed.
- 🔖 All inactive steps symbols turn off when the motor is running.
- 🥾 Speedometer (2) displays real-time value.
- 🌜 Torquemeter (3) displays real-time value.
- The torque bar (5) displays ratio between the real-time torque value (represented by cyan dots when the micromotor is running) and the maximum reached torque (represented by green dot).

Note : Each step settings are restored from the corresponding step last used settings, excluding quick settings made directly in the Operation mode.

In REVERSE mode, the rotation direction symbol 🖤 blinks and there is a sound alert (alternate medium beeps). The Torque value is automatically increased in REVERSE mode when torque-meter is displayed. The torque value can be increased from 0 to 10 Ncm, see "Reverse torque boost value", chapter "10 Special modes" on page 31 to adjust it.

Actions on foot control's buttons have no effect when the micromotor is running.

B. If necessary, release the foot control to perform the following actions:

- Y Torquemeter (3) displays maximum reached value together with the Max symbol (4).
- Solution Torque bar (5) dots that were displayed in cyan turn black, except for the maximum value dot which turns green.

Note : Changes made in this mode (either by rotating the knob or by changing parameters through the foot control buttons) are considered as temporary settings and are never saved.



FIG. 4

• Turn the knob CW or CCW to respectively increase or decrease the micromotor maximum reachable torque (quick setting mode).

The torquemeter (3) turns cyan and displays the set micromotor maximum reachable torque.

Note : Changing the speed in steps P3, P4 and P5 can only be performed through the settings mode.Long press on the knob to change operation settings.

See chapter "9 Settings" on page 29 for details.

• Long press on the orange button to activate the 5 Ncm torque boost.

Note : The torque boost can only be activated when the torquemeter is displayed in Operation mode, in low speed steps (<100 RPM).

C. Short press on the foot control's orange button or on the knob to go to the next step.

🔖 The next step symbol turns green and the step's last used settings are restored.

Note : Actions on foot control's buttons have no effect when the micromotor is running.

Changing the torque in steps P1 or P2 can only be performed through the Settings mode.

The torque boost can only be activated when the torquemeter is displayed in Operation mode, in low speed steps (<100 RPM).

For safety reasons, the speed setting icon turns red and blinks together with the speedometer for 2 seconds when switching from low speed to high speed (=100 RPM) step.



FIG. 2

8 Operation – Surgery mode

8.1 Operation screen description

FIG. 1

The Operation screen differs whether the micromotor is stopped or running and depending on the active step.

It allows to perform an operation in 3, 4 or 5 predefined steps P1, P2, P3, P4, P5, and displays the following information:

(1) Step P1 (active step, in green)

(2) Step P2 (inactive step, in black)

(3) Step P3 (inactive step, in black)

P4 and P5 steps are disabled by default, see "Number of steps", chapter "10 Special modes" on page 33 for enabling them.

(4) Speedometer

Note : Real-time speed value is displayed in black when the MX-i LED micromotor is running. Maximum reachable speed value stored is displayed in cyan when the MX-i LED micromotor is not running, in steps P1 and P2.

(5) Contra-angle ratio

Note : The contra-angle ratio is cyan-colored for direct-drive, green colored for reduction gears and redcolored for multiplication gears.

(6) Operation settings symbols

See chapter "9 Settings" on page 30for details on adjusting settings.

8.2 Perform an operation

FIG. 2

A. Operate by pressing the foot control to adjust the MX-I LED micromotor speed.

🥾 Inactive steps symbols turn off when the motor is running.

Speedometer displays real-time speed value in black.

Note : Each step settings are restored from the corresponding step last used settings, excluding quick settings made directly in the Operation mode.

In REVERSE mode, the rotation direction symbol 🖤 blinks and there is a sound alert (alternate medium beeps).

Actions on foot control's buttons have no effect when the micromotor is running.



FIG. 3

B. If necessary, release the foot control to perform the following actions:

- Speedometer (1) displays the set micromotor maximum reachable speed in cyan.
- Turn the knob CW or CCW to respectively increase or decrease the micromotor maximum reachable speed (quick setting mode).

Note : Changes made in this mode (either by rotating the knob or by changing parameters through the foot control buttons) are considered as temporary settings and are never saved.

🦠 The speedometer is cyan and displays the set micromotor maximum reachable speed (1).

Note : Changing the torque can only be performed through the Settings mode.

- Long press on the knob to change operation settings.
- 🔖 The Settings mode is displayed.

See chapter "9 Settings" on page 30on page 18_ for details.

C. Short press on the foot control's orange button or on the knob to go to the next step.

🔖 The next step symbol turns green and the step's last used settings are restored.

Note : Actions on foot control's buttons have no effect when the micromotor is running.



FIG. 2

9 Settings

FIG. 1

The Settings mode allows changing all parameters of each step. It is accessed by long pressing the knob from the Operation mode and leaved by also long pressing the knob or by running the motor. All changes made in this mode are automatically saved for the corresponding step.

Note : The rotation direction and the irrigation level symbols differ depending on the actual settings.

- A. From the Settings mode menu, navigate through the operation parameters by turning the knob CW or CCW.
- 🔖 The selected parameter symbol (1) is encased in a cyan square and an arrow points on it.
- B. If necessary, short press on the foot control's orange button to go to the next step without going back to the Operation mode.
- The Settings mode is still displayed, the next step symbol turns green and the step's last used settings are restored.
- C. Short press on the knob to change the selected parameter setting (setting sub-mode).
- 🔖 The selected setting sub-mode is displayed.

9.1 Operation mode

A. From the Settings mode menu, select the symbol and short press on the knob to change operation mode.

Note : The operation mode, rotation direction, irrigation level and the luminosity level symbols differ depending on the actual settings.

- B. Turn the knob CW or CCW to alternatively toggle between IMPLANTOLOGY IM and SURGERY IM mode.
- C. Short press on the knob to exit operation mode setting.
- Solution mode is saved and the Settings mode menu is displayed again, FIG. 1.

9.2 MX-i LED micromotor speed

A. From the Settings mode menu, select the 🎇 symbol and short press on the knob to change maximum reachable speed.

FIG. 2

B. Turn the knob CW or CCW to respectively increase or decrease micromotor maximum reachable speed.

- 🔖 The speedometer (1) displays the set maximum reachable speed.
- C. Short press on the knob to exit speed setting.
- New maximum reachable speed is saved and the Settings mode menu is displayed again, FIG. 1.

ENG A

9.3 MX-i LED micromotor torque

A. From the Settings mode menu, select the 🔍 symbol and short press on the knob to change maximum reachable torque.

FIG. 3

- B. Turn the knob CW or CCW to respectively increase or decrease micromotor maximum reachable torque.
- 🥾 The torquemeter (1) displays the set maximum reachable torque.
- C. Short press on the knob to exit torque setting.
- New maximum reachable torque is saved and the Settings mode menu is displayed again, FIG. 1.

9.4 MX-i LED micromotor rotation direction

A. From the Settings mode menu, select the 😉 symbol and short press on the knob to change rotation direction.

Note : The rotation direction and the irrigation level symbols differ depending on the actual settings.

- B. Turn the knob CW or CCW to alternatively toggle between FORWARD 🚱 and REVERSE 🙂 micromotor rotation.
- C. Short press on the knob to exit rotation direction setting.
- 🔖 Rotation direction is saved and the Settings mode menu is displayed again.

Note : The Torque value is automatically increased in REVERSE mode when torquemeter is displayed. The torque value can be increased from 0 to 10 Ncm, see chapter 9, "Reverse torque boost value" 31on page 31 to adjust it.



FIG. 2

9.5 Irrigation level

A. From the settings mode menu, select the 👀 symbol and short press on the knob to change irrigation level.

Note : The rotation direction and the irrigation level symbols differ depending on the actual settings. FIG. 4

B. Turn the knob CW or CCW to set up the irrigation level (1).

5 levels of adjustment are possible:

30ml/min, 60ml/min, 90ml/min, 120ml/min, 130ml/min.

Note : When setting the irrigation level to OFF, all dots (1) are displayed in black. Irrigation level is off when the irrigation is completely turned off by means of the foot control's blue button, regardless of the active step. In this case, the OFF symbol is displayed in Operation mode. The irrigation is considered as a quick setting and therefore is turned ON when starting again from step P1.

C. Short press on the knob to exit irrigation level setting.

& Irrigation level is saved and the Settings mode menu is displayed again.

9.6 Contra-angle ratio

- A. From the Settings mode menu, select the **O**[•] symbol and short press on the knob to change the contra-angle ratio.
- B. Turn the knob CW or CCW to change the contra-angle ratio.

Note : The contra-angle ratio is cyan-colored for direct-drive, green colored for reduction gears and redcolored for multiplication gears.

The contra-angle labelled "125L" corresponds to a multiplication ratio of 1:2.5.

9.7 Luminosity level

A. From the Settings mode menu, select the symbol and short press on the knob to change luminosity level.

Note : The operation mode, rotation direction, irrigation level and the luminosity level symbols differ depending on the actual settings.

A. Turn the knob CW or CCW to set up the luminosity level. 10 levels of adjustment are possible.

- B. Short press on the knob to exit luminosity level setting.
- & Luminosity level is saved and the Settings mode menu is displayed again.
- C. Short press on the knob to exit the contra-angle ratio setting.

🔖 The contra-angle ratio is saved and the Settings mode menu is displayed again.





10 Special modes

The special modes allow to, in the following order:

- Display software version;
- Test LCD display;
- Define number of steps (3, 4 or 5);
- Define reverse torque boost value;
- Restore factory settings.

Note : Pressing the foot control has no effect in the Special modes

A. From the Operation mode, long press on the rotating knob to enter Settings modes.

🌭 The Settings mode is displayed.

FIG. 1

B. Turn the knob CW or CCW to select the contra-angle ratio symbol (1).

🔖 The contra-angle ratio symbol is encased in a cyan square and an arrow points on it.

Software version

FIG. 2

C. Double short press on the knob to enter special modes.

🔖 The contra-angle ratio symbol (3) turns blue to differentiate it from the ratio change cyan symbol.

- ✤ The software version is displayed as following:
- (1) Major version

(2) Minor version

LCD display test

FIG. 3

D. Short press on the knob to test LCD display.

All dots are displayed in black, except for the contra-angle ratio symbol 오 (1).

Number of steps

E. Short press on the knob to define the number of steps.

✤ The step number screen is displayed.

F. Turn the knob CW or CCW to alternatively display the 3, 4 or 5 text.

G. Short press on the knob to define the number of steps.

Reverse torque boost value

Reverse torque boost allows an automatic increase of torque value when in REVERSE mode, in order to ease bur rotation when stuck.

H. Short press on the knob to define reverse torque boost value.

🔖 The reverse torque boost screen is displayed.

I. Turn the knob CW or CCW to alternatively display the 0, 5 or 10 text.

J. Short press on the knob to define no boost value when **0** is displayed, or short press on the knob to respectively define 5 Ncm or 10 Ncm boost value when **5** or **10** is displayed.

Settings reset

FIG. 4

K. Short press on the knob to display factory settings reset screen.

✤ The factory settings reset screen is displayed.

L. Turn the knob CW or CCW to alternatively display the reset yes or reset no text (1).

The **reset** no text is displayed by default.

M. Short press on the knob to restore factory settings when the **reset yes** text is displayed, or short press to go back to the Settings mode when the **reset no** text is displayed.

\$

Reset can take up to 2 seconds. Meanwhile, the a symbol is displayed, and the **yes** text is turned off. When reset is done, the Settings mode is displayed again.



FIG. 4

Note : Pressing the foot control has no effect in the Special modes. Go through all the special modes to display the Settings mode again. The reset no text is displayed by default.

11 List of errors & Troubleshooting

11.1 Alert notifications (operating)

Alert description	Message	Cause of warning	Action
Motor overheating	*	Excessive power demand of the MX-i LED micro- motor.	Avoid extended use. Let system cool down.
Release pedal [foot control]	▲ ≟	 Foot control is pressed when accessing settings submodes. Foot control is pressed during device startup. Foot control is pressed after recovering from an error. 	 Confirm setting by pressing the knob. Release foot control and press it again. Release foot control and press it again.
Low to high speed step tran- sition	▲ ₩	User switches from low speed to high speed (= 100 RPM) step.	No action needed, the alert disappears after 2 seconds.
Motor jammed	A	Motor is jammed for more than 2 seconds. Motor power supply is cut to avoid overheating.	Release the foot control, release the bur and press foot control again.
Footpedal [foot control] not connected	•• ••	Foot control is not connected to device.	Connect the foot control to the device
Motor not connected	A	Motor is not properly connected to device, Motor hardware is damaged.	 Acknowledge error. (Re)connect the motor cable. If problem persists, contact Bien-Air Dental SA.

11.2 Device operating error

Error description	Caus	e of error	When	Acti	ion	
ERROR 1						
Motor short- circuit	Electrical failure: short- circuit between motor phases.		ln running mode.	Rep	Replace motor and/or cable.	
ERROR 2						
Main controllerOtherfaultconditionerrordetected by software.		Any time.	 Switch off system. Contact Bien-Air Dental SA. 			
ERROR 3						
Motor driver communication timeout error	driver cation rror 232. Failure of main controller		ln running mode.	1. Switch off system. 2. Contact Bien-Air Dental SA.		
ERROR 4						
Invalid EEPROM memory	Failure of EEPROM memory.		Any time.	Con Ack norr save or re	Contact Bien-Air Dental SA. Acknowledging this error allows the operator to work normally but it will not allow settings to be saved or restored. This error will appear at each save or restoration attempt.	
Error descripti	on	Cause of error			When	Action
ERROR 5						
Motor drive temperature	over	Motor overload in a high temperature environment. Failure of DMX controller.		ature	Any time.	 Wait for system cooling. If problem persists, contact Bien-Air Dental SA.
ERROR 6						
Motor driver voltage error	under	Motor overload in a high temperature environment. Failure of power supply.		Any time.	 Acknowledge error. If problem persists, contact Bien-Air Dental SA. 	
ERROR 7						
Motor driver voltage error	over	Failure of power supply. Tool used has a too high inertia.		Any time.	 Acknowledge error. If problem persists, contact Bien-Air Dental SA. 	
ERROR 8						
Irrigation general failure	pump	Electrical failure: short-circuit to ground or to supply. Electrical failure: short-circuit between motor phases.		ln running mode.	1. Switch off system. 2. Contact Bien-Air Dental SA.	
ERROR 9						
Knob failure	re Electrical failure of knob encoder			Any time.	 Switch off system. Contact Bien-Air Dental SA. 	



12 Maintenance

12.1 Servicing

\triangle caution

- Never disassemble the device. For all servicing or repair operations, you are advised to contact your regular supplier or Bien-Air Dental SA directly.
- In order to avoid any risk of contamination, the knob must be sterilized prior to servicing. See chapter 11.2 Cleaning & Sterilization below for details.

Service period

The device was tested by simulating 10,000 clinical procedures (corresponding to a service period of 6 to 10 years). If the actual use of the device exceeds the tested service period, preventive maintenance of the device is recommended.

12.2 Sterilization

\triangle caution

- Do not immerse in disinfectant solution.
- Not designed for an ultrasonic bath.

FIG. 1

Cleaning

(A) Remove the knob (1) and rinse it twice with running tap water (15° C-38° C) 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.

Note : The knob is held magnetically. There is no need to preserve its angular position when removing it or putting it back in place.

(B) Clean the unit including the bracket, the foot control and the external and internal surfaces of the knob by gently rubbing it with a clean cloth soaked in a suitable product (i.e. Bien-Air Dental Spraynet or isopropyl alcohol for about 15 sec.).

Sterilization of the knob

For an efficient sterilization process of the knob, the warnings and cautions below must be observed:

- Before using it for the first time clean and sterilize the knob.
- Do not use a sterilization procedure other than the one described below.

\triangle caution

- The quality of the sterilization is highly dependent on how clean the instrument is. Only perfectly clean instruments should be sterilized.
- Pack the knob in a packaging approved for steam sterilization.
- Only use dynamic air removal cycles: pre-vacuum or steam flush pressure pulse (SFPP) cycles.

Sterilize the knob 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. In jurisdictions where sterilization for prions is required, sterilize at 135° C for 18 minutes.

The recommended parameters for the sterilization cycle are:

- The maximum temperature in the autoclave chamber does not exceed 137° C, i.e. the nominal temperature of the autoclave is set at 134° C, 135° C or 135.5° C taking into account the uncertainty of the sterilizer as regards temperature.
- The maximum duration of the interval at the maximum temperature of 137° C 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 comprised in the interval between 0.07 bar to 3.17 bar (1 psia to 46 psia).
- The rate of change of temperature does not exceed 15° C/min for increasing temperature and -35° C/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 water steam.

12.3 Important

For maintenance:	See instructions for use
MX-i LED micromotor	REF 2100245
Cable for micromotor	REF 2100163
Contra-angle CA 20:1 L Micro-Series, light	REF 2100209
Contra-angle CA 20:1 L KM Micro-Series, light	REF 2100209
Micro-Series, light	REF 2100337



12.4 Replacement of fuses

- A. Switch off the Chiropro Plus 3rd Gen unit.
- B. Disconnect the mains cable.

\triangle caution

The power cable must be disconnected at least 10 seconds before opening the fusebox.

FIG. 2

C. Remove the fuse box (1) with a flat screwdriver.

FIG. 3

D. Replace the fuses (2) by the new ones and put the fuse box back (1) in place.

${\rm \ensuremath{\underline{\wedge}}}$ caution

Only use fuses T4.0AH 250 VAC REF 1307312-010.

13 Guarantee

13.1 Terms of guarantee

Bien-Air Dental SA grants the user a guarantee covering all functional defects, material or production faults.

The device is covered by this guarantee from the date of invoicing for:

- 12 months for the motor cable;
- 24 months for the Chiropro Plus 3rd Gen unit and CA 20:1 L Micro-Series;
- 36 months for the MX-i LED micromotor.

In case of justified claim, Bien-Air Dental SA or its authorized representative will fulfill the company's obligations under this guarantee by repairing or replacing the product free of charge.

Any other claims, of whatever nature, in particular in the form of a claim for damages and interest, are excluded.

Bien-Air Dental SA shall not be held responsible for damage or injury and the consequences thereof, resulting from:

- excessive wear and tear
- improper use
- non-observance of the instructions for installation, operation and maintenance
- unusual chemical, electrical or electrolytic influences
- poor connections, whether of the air, water or electricity supply.

The guarantee does not cover flexible optical fiber type light conductors, or any parts made of synthetic materials.

The guarantee shall become null and void if the damage and its consequences are due to improper manipulation of the product, or modifications to the product carried out by persons not authorized by Bien-Air Dental SA.

Claims under the terms of the guarantee will be considered only on presentation, together with the product, of the invoice or the consignment note, on which the date of purchase, the product reference and the serial no. should be clearly indicated.

Please refer to the General Terms and Conditions of Sale on www.bienair.com.



Bien-Air Dental SA Länggasse 60 Case postale 2500 Bienne 6 Switzerland Tel. +41 (0)32 344 64 64 Fax +41 (0)32 344 64 91 dental@bienair.com

Other addresses available at **www.bienair.com**

EC REP Bien-Air Europe Sàrl 19-21 rue du 8 mai 1945 94110 Arcueil France