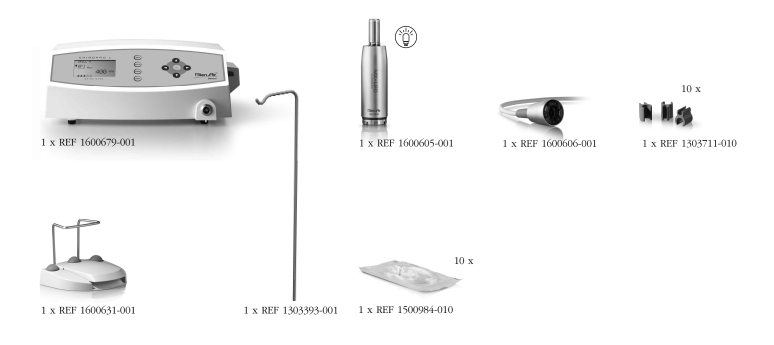


CHIROPRO L

ENG Instructions for use



Set REF 1700349-001



Options



REF 1600632-001



REF 1600605-001











REF 1600631-001



REF 1303711-010





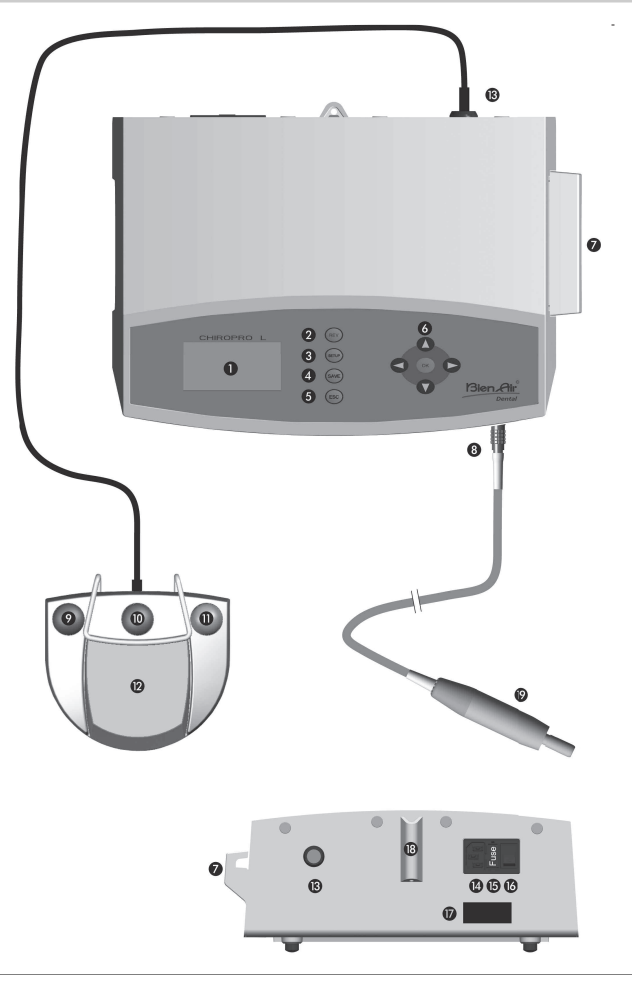
REF 1301560-010

REF 1500984-010









Summary

Starting display



Available values

MAIN MENU	Steps	Ratio	Speed in rpm	Torque in Ncm	Irrigation in ml/min
Implantalagu	Round bur 1	128:1	100 40/000 rpm	0.48 - 4.8 Ncm	30 ml/min 20%
Implantology		128.1	100 - 40′000 rpm	0.48 - 4.8 NCM	30 mi/min 20%
Endodontics	Round bur 2	64:1	with a CA 1 : 1	with a CA 1 : 1	60 ml/min 40%
Surgery	Drill 1	30:1			90 ml/min 60%
	Drill 2	27:1	Depends on the CA	Depends on the CA	120 ml/min 80%
	Drill 3	20:1			150 ml/min 100%
	Drill 4	16:1			
	Tapping	10:1			
	Tap unscrewing	1:1			
	Implant screwing	1:2]		
	Unscrewing	1:5]		

Implantology	Open pulp chamber	128:1	100 - 40'000 rpm	0.48 - 4.8 Ncm	30 ml/min 20%
Endodontics	endo file 1	64:1	with a CA 1 : 1	with a CA 1 : 1	60 ml/min 40%
Surgery	endo file 2	30:1			90 ml/min 60%
	endo file 3	27:1	Depends on the CA	Depends on the CA	120 ml/min 80%
	endo file 4	20:1			150 ml/min 100%
	endo file 5	16:1			
	endo file 6	10:1			
	endo file 7	1:1			
	endo file 8	1:2			
	endo file 9	1:5			

Implantology	Procedure 1	128:1	100 - 40'000 rpm	0.48 - 4.8 Ncm	30 ml/min 20%
Endodontics	Procedure 2	64:1	with a CA 1 : 1	with a CA 1 : 1	60 ml/min 40%
Surgery	Procedure 3	30:1]		90 ml/min 60%
	Procedure 4	27:1	Depends on the CA	Depends on the CA	120 ml/min 80%
		20:1]		150 ml/min 100%
		16:1]		
		10:1]		
		1:1			
		1:2			
		1:5			

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16+34

1 Meaning of symbols

ENG

C	E
01	23

CE Marking with number of the notified body.



Caution: Federal law (USA) restricts this device to sale by or on the order of a licensed healthcare practitioner.



Protective earth (ground).



Main switch ON - The instrument is energized. OFF - The instrument is de-energized.

orr - me instrument is de-



Alternating current.

Device of type B.

Fuse Ø 5 x 20 mm.



CAUTION ! Dangerous voltage.



Element sensitive to electrostatic discharges.



CAUTION! Refer to the accompanying documents.



Danger of pinching. Do not put your fingers in rotating parts.



Variability in steps.



Symbol for «Water-cooling».



Symbol for «peristaltic pump».



Recyclable materials.



Recyclable electrical and electronic material.

Sterilisable up to the specified temperature.



Operating mode intermittent.



Manufacturer.

STERILE EO

Sterilise with Ethylene Oxyde

2

2 Description

Identification	motor with v A peristaltic p contaminated The device's	lly controlled tabletop device for dentistry allowing operation of an MX-LED CHIROPRO a variable speed control by a pedal. c pump conveys the physiological liquid via a disposable irrigation line without being ted. 's LCD display indicates the stage of implant fitting, the instrument's ratio, the bur speed, ue and irrigation flow setting.			
Intended use	igned to con to cut hard a The system is Any use othe	em is to be used by dentists and surgeons in dental offices and hospitals. The system is de control a dental micromotor which can drive a dental hand-piece fitted with appropriate t rd and soft tissues in the mouth and to screw dental implants. em is intended for use in dentistry for implantology, dental surgery and endodontic work. other than that for which this product is intended is unauthorised and may be dangerous. ical device meets all the current legal requirements.			
Environment	\wedge				
Environment	The device is	s not designed for use in ar	n explosive atmosphere (anaesthetic gas).		
	Working	Temperature: Relative humidity: Atmospheric pressure:	+10°C (50°F) to +25°C (77°F) 30% to 80%, including condensation 700 hPa to 1060 hPa		
	Transport and storage	Environmental conditions Temperature: Relative humidity: Atmospheric pressure:	-25°C (-13°F) to +70°C (158°F) 10% to 100%, including condensation 500 hPa to 1060 hPa		

Environmental protection and information for disposal of the instrument



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



This device and its accessories must be recycled.

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 2002/96/EC).

3 Set supplied

1x	CHIROPRO L control	REF	1600679-001
1x	Micromotor MX-LED CHIROPRO	REF	1600605-001
1x	Cable for MX-LED micromotor	REF	1600606-001
1x	Pack of 10 disposable sterile lines	REF	1500984-010
1x	10 attachments collars for fastening the sterile irrigation line to a cable	REF	1303711-010
1x	Bracket for fluid bottle	REF	1303393-001
1x	Pedal 3 buttons	REF	1600631-001
1x	Cable system 3P, US / Asia, length 2,00 m	REF	1300067-001
1x	Instruction	REF	2100189

4 **Options**

Contra-angle handpiece CA 20:1 L (light)	REF 1600598-001
Contra-angle handpiece CA 20:1 (without light)	REF 1600632-001
Contra-angle handpiece CA 20:1 L Micro-Series (light)	REF 1600692-001
Micromotor MX-LED CHIROPRO	REF 1600605-001
Cable for MX-LED CHIROPRO micromotor	REF 1600606-001
Pack of 10 disposable sterile lines	REF 1500984-010
10 attachments collars for fastening the sterile irrigation line to a cable	REF 1303711-010
Pedal 3 buttons	REF 1600631-001
Cable system 3P, Switzerland, length 2,00 m	REF 1300065-001
Cable system 3P, Europe, length 2,50 m	REF 1300066-001
Cable system 3P, US / Asia, length 2,00 m	REF 1300067-001
10 fuse T4.0A L 250 VAC	REF 1301560-010

5 Technical Description: Technical data

Voltage

100 – 240 VAC 50 – 60 Hz

Fuses

2 fuses T4.0A L 250 VAC, breaking capacity 40A

Power demand

300 VA

Classification

Class IIa in accordance with European Directive 93/42/EEC concerning medical devices.

Electric insulation class

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

Degree of protection

IP 40 (protection against insertion of objects larger than 1 mm).

Dimensions L x W x H

 $309 \ x \ 220 \ x \ 123 \ mm.$ Height with bracket $506 \ mm$

Weight

Housing	2.7 kg	Pedal	830 g
Cable	105 g	Bracket	115 g

Memory

Implantology mode:	Storage in memory of 8 implant fitting
	sequences of 10 steps each.
Endodontics mode:	Storage in memory of an endodontics
	sequence of 10 steps.
Surgery mode:	Storage in memory of 4 separate
	programs.

Interface Languages

French, German, English, Italian, Spanish, Portuguese, Japanese and Russian.

List of errors & Toubleshooting

Page 15

Bracket

Stainless steel

Intended for use with:

Micromotor MX-LED CHIROPRO	REF 2100161
Cable for MX-LED CHIROPRO micromotor	REF 2100163
Contra-angle CA 20:1, without light	REF 2100209
Contra-angle CA 20:1 L, with light	REF 2100209
Contra-angle CA 20:1 L Micro-Series, with light	REF 2100209

see instruction

The use of the system with other handpieces, motors or cables has not been validated/certified.

Peristaltic pump

```
Pump delivery:
Hose for pump:
```

From 30 to 150 ml/min. (5 levels). External \emptyset 5.60 mm, internal \emptyset 2.40 mm Wall thickness 1.60 mm.

Pedal

REF 1600631-001Dimensions (LxWxH) 250 x 205 x 54 mmwith handle:250 x 205 x 144 mmThe pedal is waterproof (IP X8 in accordance with CEI 529).

Cables

Length of cables: Pedal cable 2,90 m Motor cable 2,00 m

WARNING

To prevent any risk of electric shock, this device must be connected only to a power supply network provided with protective earth. Modification of the device forbidden. The system is not adapted to be used in the presence of inflammable gases (e.g. anaesthetic gas). Do not attempt to open the apparatus when it is connected to the electric mains. Beware of electric shocks.

Applied parts (per IEC 60601-1)

 Micromotor MX-LED CHIROPRO
 REF 1600605-001

 CA 20:1 L
 REF 1600598-001

 CA 20:1
 REF 1600632-001

 CA 20:1 L Micro-Series
 REF 1600692-001

 Irrigation lines
 REF 1500984-010

Operating mode:

Intermittent ON: 5 min. OFF: 40 min.

5 Technical Description: Electromagnetic compatibility

Precautions regar- ding Electromagnetic Compatibility (EMC)	Electro-medical equipment needs special precautions regarding EMC and needs to be installed and pu into service according to the EMC information provided in the present document.			
computering (arro)	ment, co ence the such as	ellular phones, etc. shall not e performance of the device. High Frequency surgical and	C requirements according to IEC 60601-1-2. Radio transmitting equip- be used in the close proximity of the device since they could influ- Particular precaution is required when using strong emission sources d similar equipment so that the HF cables are not routed on or near ct a qualified technician or Bien-Air Dental.	
		sary, CHIROPRO L should be	djacent or stacked with other equipment. If adjacent or stacked use e monitored to verify normal operation in the configuration in which	
WARNING!	The use of accessories, transducers and ca-bles other than those specified, with the exception of transducers and cables sold by Bien-Air Dental as replacements parts for internal components, may result in increased emissions or decreased immunity of CHIROPRO L. 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.			
Guidance and manu-			the electromagnetic environment specified below.	
facturer's declaration - electromagnetic emissions			PRO L should ensure that it is used in such an environment.	
Emissions test		Compliance	Electromagnetic environment - guidance	

Emissions test	Compliance	Electromagnetic environment - guidance
RF emissions CISPR 11	Group 1	CHIROPRO L uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.
RF emissions CISPR 11	Class B	CHIROPRO L is suitable for use in all Establishments, including domestic establishments and those directly connected
Harmonic emissions IEC 61000-3-2	Not applicable	to the public low-voltage power supply network that supplies buildings used for domestic purposes.
Voltage fluctuations/flicker emissions IEC 61000-3-3	Conform	

Guidance and manufacturer's declaration - **electromagnetic immunity** CHIROPRO L is intended for use in the electromagnetic environment specified below. The customer or the user of CHIROPRO L should ensure that it is used in such an envir

Immunity test	IEC 60601 test level	Compliance level	Electromagnetic environment - guidance
Electrostatic	±6 kV contact	±6 kV contact	Floors should be wood, concrete or ceramic tile. If floors
discharge (ESD)			are covered with synthetic material, the relative humidity
TTC (1000 / 0	±8 kV air	±8 kV air	should be at least 30%.
IEC 61000-4-2	.2.1.1.6	-21X6	
Electrical fast transient/burst	±2 kV for power	±2 kV for power	Mains power quality should be that of a typical commer-
IEC 61000-4-4	supply lines No input/output lines	supply lines	cial or hospital environment.
IEC 01000-4-4	No input/output intes	No input/output lines	
Surge	±0.5 kV line to line	±0.5 kV line to line	
IEC 61000-4-5	±1 kV line to line	± 1 kV line to line	Mains power quality should be that of a typical commer-
			cial or hospital environment.
	±0.5 kV line to earth	±0.5 kV line to earth	cial of hospital environment.
	±1 kV line to earth	±1 kV line to earth	
	±2 kV line to earth	± 2 kV line to earth	
Voltage dips, short	<5% UT	<5% UT	
interruptions and	$(>95\% \text{ dip in } U_{T})$	(>95% dip in $U_{\rm T}$)	Mains power quality should be that of a typical commer-
variations de tension	for 0,5 cycle	for 0,5 cycle	cial or hospital environment. If the user of CHIROPRO L
voltage variations	(00) II	(00) 1	requires continued operation during power mains inter-
on power supply	$40\% U_{\rm T}$	$40\% U_{\rm T}$	ruptions, it is recommended that CHIROPRO L be pow-
input lines 1	$(60\% \text{ dip in } U_{\mathrm{T}})$	(60% dip in $U_{\rm T}$)	ered from an uninterruptible power supply or a battery.
	for 5 cycles 70% $U_{\rm T}$	for 5 cycles 70% $U_{\rm T}$	
IEC 61000-4-11	$(30\% \text{ dip in } U_{\rm T})$	$(30\% \text{ dip in } U_{\rm T})$	
IEC 01000-4-11	for 25 cycles	for 25 cycles	
	ior 29 cycles	ior 29 cycles	
	<5% UT	<5% UT	
	$(>95\% \text{ dip in } U_{\rm T})$	$(>95\% \text{ dip in } U_{T})$	
	for 5 sec	for 5 sec	
Power frequency			Power frequency magnetic fields should be at levels
(50/60 Hz)	3 A/m	3 A/m	characteristic of a typical location in a typical commer-
magnetic field			cialor hospital environment.
IEC 61000-4-8			1

NOTE $U_{\rm T}$ is the a.c. mains voltage prior to application of the test level.

5 Technical Description: Electromagnetic compatibility

Guidance and manufacturer's declaration - electromagnetic immunity

CHIROPRO L is intended for use in the electromagnetic environment specified below. The customer or the user of CHIROPRO L should ensure that it is used in such an environment.

Immunity test	IEC 60601 test level	Compliance level	Electromagnetic environment - guidance
			Portable and mobile RF communications equipment should be used no closer to any part of CHIROPRO L, including cables, than the recommended separa- tion distance calculated from the equation applicable to the frequency of the transmitter.
			Recommended separation distance
Conducted RF IEC 61000-4-6	3 Vrms 150 kHz to 80 MHz	3 V	$d = 1.2\sqrt{P}$ $d = 1.2\sqrt{P}$ $d = 2.3\sqrt{P}$ 80 MHz to 800 MHz 800 MHz to 2,5 GHz
Radiated RF IEC 61000-4-3	3 V/m 80 MHz to 2,5 GHz	3 V/m	where <i>P</i> is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and <i>d</i> is the recommended separation distance in metres (m). 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. ^b Interference may occur in the vicinity of equipment marked with the following symbol: $((\bullet))$

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

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

^a Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the CHIROPRO L is used exceeds the applicable RF compliance level above, the CHIROPRO L should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as reorienting or relocating the CHIROPRO L.

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

Recommended separation distances between portable and mobile RF communications equipment and the CHIROPRO L

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

Rated maximum output	Separation distance according to frequency of transmitter					
power of transmitter	m					
\mathbf{W}	150 kHz to 80 MHz	150 kHz to 80 MHz 80 MHz to 800 MHz 800 MHz to 2,5 GHz				
	$d = 1.2\sqrt{P}$	$d = 1.2\sqrt{P}$	$d = 2.3\sqrt{P}$			
0.01	0.12	0.12	0.23			
0.1	0.38	0.38	0.73			
1	1.2	1.2	2.3			
10	3.8	3.8	7.3			
100	12	12	23			

For transmitters rated at a maximum output power not listed above, the recommended separation distance d in metres (m) can be estimated using the equation applicable to the frequency of the transmitter, where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer.

NOTE 1 At 80 MHz and 800 MHz, the separation distance for the higher frequency range applies.

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

6 Installation

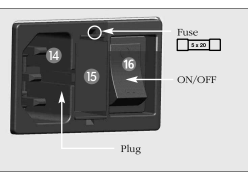
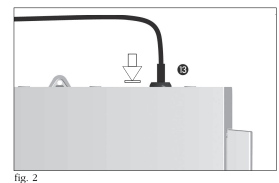


fig. 1



Installation

A. CHIROPRO L may be positioned on a table, on a trolley or another surface, but in no case on the floor.Power plug (2) is the device for disconnection in case of problems,

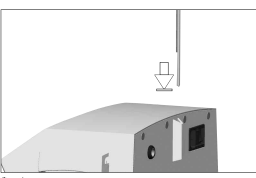
and it must be easily accessible at all times.

- B. The fuse box may be opened with a screwdriver . 100 - 240 Vac = fuse T-4.0 A L 250 VAC REF 1301560-010
- C. The apparatus is powered by your line voltage (100/115/230 Vac). Connect the power cable to the plug **fig. 1**.
- D. Connect the pedal cable to the output provided on the rear panel, guiding the connector and plug by means of the index pin on the connector **fig. 2**.

 \triangle Do not raise the pedal using the connection cable.

- E. Connect the MX-LED CHIROPRO micromotor cable to the motor output, guiding the connector and plug by means of the index pin and red dot on the connector **fig. 3**.

fig. 3



F. Align and attach the bracket to the housing provided on the console's rear and suspend the flask or bottle **fig. 4**.

fig. 4



G. Check the packaging integrity, as well as the expiration date of the irrigation line.

Only lines supplied by Bien-Air Dental ensure trouble-free operation. These lines are sterile and for single use. Re-use may result in microbiological contamination of the patient.

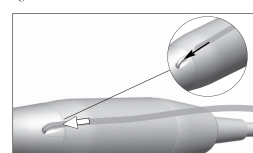




6 Installation



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



Fitting on the spray tube

I. Connect the flexible hose of the irrigation line to the spray tube of the handpiece or contra-angle **fig. 7**.



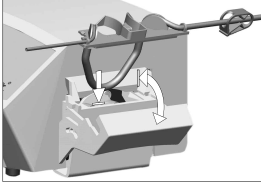


fig. 8

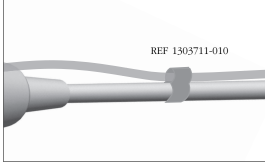


fig. 9

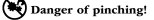
Stopping procedure

The device can be safely stopped using the main switch (6).

Installation on the peristaltic pump

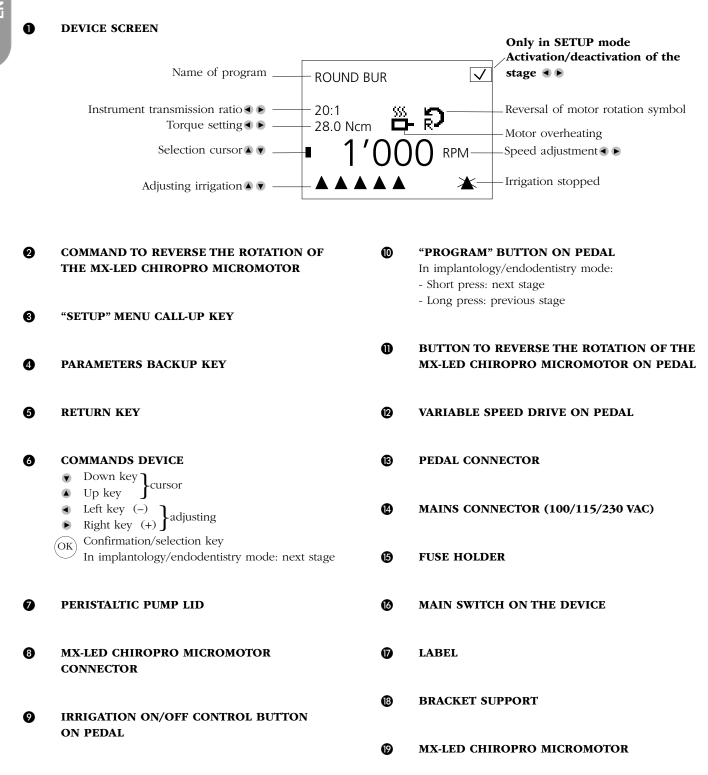
J. Install the plastic cassette in the peristaltic pump.
Check that the cassette is clipped correctly.
Close the pump lid, **fig. 8**.
If there is resistance to closing, open the lid again and check the correct positioning of the cassette.

Warning! Do not run the pump while the lid is open.



- K. Perforate the cap of the physiological liquid flask with the pointed end of the irrigation line after removing the protective cap.
- L. Attach the irrigation line on the motor cable using the attachment collars REF 1303711-010 **fig. 9**.

7 Description of keys and elements



9 List of errors & Toubleshooting

Message		Cause of error	Action
\wedge	The pedal is pressed when starting the device.	Safety	Release the pedal and press
Release the		Salety	again.
pedal	The motor is blocked for more than 2 sec.		0
	The motor control card limits the power supplied to the motor to	Safety	Avoid extended use.
	prevent motor overheating.	Salety	Avoid extended use.
Equipment ini	tialisation error		
	rror may occur at start-up of CHIROP	RO L	
1. Check on th	e integrity of the CHIROPRO L m	nemory	
INIT ERROR 1	The memory is corrupt! Please contact Bien-Air Dental SA. ESC: restore	The memory data check failed.	Press the ESC key to try to restore the memory. Contact Bien-Air Dental SA.
Device operati The following e	i ng error rrors may occur during operation of t	he device	
L. Loss of ped	al connection		
ERROR 1	The pedal is not connected! Please check the connection. ESC: exit	The pedal is not connected correctly.	Check pedal connection. Contact Bien-Air Dental SA.
2. Peristaltic p	oump overheating		
ERROR 2	Irrigation pump overheating! Please wait for it to cool.	Peristaltic pump motor overheating	Wait until the system cools. Contact Bien-Air Dental SA.
	ESC: exit		
3. Peristaltic p	ump general error	1	
ERROR 3	Irrigation pump fault! Please contact Bien-Air Dental SA. ESC: exit	Peristaltic pump electrical fault.	Contact Bien-Air Dental SA.
4. Loss of mot	or connection		
ERROR 4	The motor is not connected! Please check the connection. ESC: exit	Loss of motor phase fault. The motor is not connected correctly.	Check motor connection. Contact Bien-Air Dental SA.
5. Motor cable	fault		
ERROR 5	Motor cable fault! Please change cable. ESC: exit	Motor power fault. The motor cable may be defective.	Check motor cable. Contact Bien-Air Dental SA.
6. Motor contr	ol overheating	1	
ERROR 6	System overheating! Please wait for it to cool. ESC: exit	Overheating of motor control card (electrical control of motor).	Wait until the system cools. Contact Bien-Air Dental SA.
7. System elec	trical fault		
GEN ERROR	System electrical fault!	Communication fault with motor	Contact Bien-Air Dental SA.
Error code]	Please contact Bien-Air Dental SA.	control card: [EC100]	
	ESC: exit	Motor control card power supply undervoltage: [EC101]	
		Motor control card power supply overvoltage: [EC102]	
		Other motor control card faults: [EC120]	

10 Default values

Implantology: Default values page 33

The table shows the default operating values for 8 different implantology sequences.

Endodontics:

Default values page 34

The table shows the default operating values for the endodontics sequence.

11 Maintenance

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.

Servicing

Never disassemble the device. For any modification and repair, we recommend that you contact your regular supplier or Bien-Air Dental directly. Bien-Air Dental asks the user to have its dynamic instruments checked or inspected at least once a year.

Information

The technical specifications, illustrations and dimensions contained in these instructions are given only as a guide. They may not be the subject of any claim. The manufacturer reserves the right to make technical improvements to its equipment, without amending these instructions. For all additional information, please contact Bien-Air Dental SA at the address indicated on the back cover.

12 Generalities and guarantee

General information

The device must be used by a qualified professional in compliance with the current legal provisions concerning workplace safety, health and accident prevention measures, and these working instructions. In accordance with such requirements, the operators:

• must only use devices that are in perfect working order; in the event of irregular functioning, excessive vibration, abnormal heating or other signs that may indicate malfunction of the device, the work must be stopped immediately; in this case, contact a repair centre that is approved by Bien-Air Dental;

• 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, and must avoid contamination through the use of the product.

Terms of guarantee

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

The device is covered by this guarantee for 24 months from the date of invoicing.

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

Surgery:

Default values page 34

The table shows the default operating values for 4 different surgical sequences.

Cleaning-disinfection

Disinfect the surfaces of the console and pedal with a clean cloth soaked in a suitable product.
Do not exert any pressure on the screen.
Do not immerse in disinfectant solution
Not designed for an ultrasonic bath.
Use a new sterile irrigation line for each patient.
AAMI TIR 12:2004 - Disinfection level: intermediate.

Important

-	
For maintenance:	see instruction
- MX-LED CHIROPRO micromotor	REF 2100161
- Cable for micromotor MX-LED CHI	ROPRO REF 2100163
- Contra-angle CA 20:1 L	REF 2100209
- Contra-angle CA 20:1	REF 2100209
	BEE 2100200

- Contra-angle CA 20:1 L Micro-Series REF 2100209

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

Bien-Air Dental 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 "fibre optic" 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 authorised by Bien-Air Dental.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.

Implantology: default values

The table shows the default operating values for 8 different implantology sequences. These default values represent common settings used during implant procedures.

Implant PRG1 / PRG8	Implant PRG2	Implant PRG3	Implant PRG4	Implant PRG5	Implant PRG6	Implant PRG7
ROUND BUR 1	ROUND BUR	ROUND BUR	ROUND BUR	ROUND BUR	ROUND BUR	PILOT DRILL 1
20:1	20:1	20:1	20:1	20:1	20:1	20:1
28.1 Ncm	28.1 Ncm	28.1 Ncm	28.1 Ncm	28.1 Ncm	35.3 Ncm	28.1 Ncm
1'000 RPM	2'000 RPM	1'000 RPM	1'200 RPM	1'500 RPM	1'500 RPM	800 RPM
ROUND BUR 2	PILOT DRILL	DRILL 1	DRILL 1	DRILL 1	DRILL 1	PILOT DRILL 2
20:1	20:1	20:1	20:1	20:1	20:1	20:1
28.1 Ncm	35.3 Ncm	35.3 Ncm	35.3 Ncm	35.3 Ncm	35.3 Ncm	28.1 Ncm
1'000 RPM	800 RPM	800 RPM	800 RPM	500 RPM	1'500 RPM	800 RPM
DRILL 1	DRILL 1	DRILL 2	DRILL 2	DRILL 2	DRILL 2	DRILL 1
20:1	20:1	20:1	20:1	20:1	20:1	20:1
35.3 Ncm	35.3 Ncm	35.3 Ncm	35.3 Ncm	35.3 Ncm	35.3 Ncm	35.3 Ncm
800 RPM	800 RPM	800 RPM	800 RPM	500 RPM	1'500 RPM	600 RPM
DRILL 2	DRILL 2	DRILL 3	DRILL 3	DRILL 3	DRILL 3	DRILL 2
20:1	20:1	20:1	20:1	20:1	20:1	20:1
35.3 Ncm	35.3 Ncm	35.3 Ncm	35.3 Ncm	35.3 Ncm	35.3 Ncm	35.3 Ncm
600 RPM	800 RPM	800 RPM	800 RPM	500 RPM	1'500 RPM	500 RPM
DRILL 3	DRILL 3	DRILL 4	DRILL 4	DRILL 4	DRILL 4	DRILL 3
20:1	20:1	20:1	20:1	20:1	20:1	20:1
35.3 Ncm	35.3 Ncm	35.3 Ncm	35.3 Ncm	35.3 Ncm	35.3 Ncm	35.3 Ncm
500 RPM	800 RPM	800 RPM	800 RPM	500 RPM	1'500 RPM	400 RPM
DRILL 4	DRILL 4	DRILL 5	DRILL 5	DRILL 5	DRILL 5	SHAPING DRILL
20:1	20:1	20:1	20:1	20:1	20:1	20:1
35.3 Ncm	35.3 Ncm	35.3 Ncm	35.3 Ncm	35.3 Ncm	35.3 Ncm	35.3 Ncm
400 RPM	800 RPM	800 RPM	800 RPM	500 RPM	1'500 RPM	250 RPM
TAPPING	TAPPING	TAPPING	TAPPING	TAPPING	TAPPING	TAPPING
20:1	20:1	20:1	20:1	20:1	20:1	20:1
35.3 Ncm	35.3 Ncm	35.3 Ncm	35.3 Ncm	35.3 Ncm	35.3 Ncm	35.3 Ncm
15 RPM	15 RPM	15 RPM	15 RPM	15 RPM	20 RPM	20 RPM
TAP UNSCREWING	TAP UNSCREWING	TAP UNSCREWING	TAP UNSCREWING	TAP UNSCREWING		TAP UNSCREWING
20:1	20:1	20:1	20:1	20:1	20:1	20:1
42.5 Ncm	42.5 Ncm	42.5 Ncm	42.5 Ncm	42.5 Ncm	42.5 Ncm	42.5 Ncm
15 RPM REV	15 RPM REV	15 RPM REV	15 RPM REV	15 RPM REV	15 RPM REV	
15 RPM REV	15 RPM REV	15 RPM REV	15 RPM REV	15 RPM REV	15 RPM REV	20 RPM REV
AAAA IMPLANT SCREWING 20:1	AAAAA IMPLANT SCREWING 20:1	AAAA IMPLANT SCREWING 20:1		AAAA IMPLANT SCREWING 20:1		AAAAA IMPLANT SCREWING 20:1
A A A A IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM	AAAAA IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM	AAAAA IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM	AAAAA IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM	AAAA IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM	△△△△△ IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM	AAAA IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM
AAAA IMPLANT SCREWING 20:1 35.3 Ncm			△ △ △ △ IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM △ △ △ △ △	△△△△ IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM △△△△△		
A A A A IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM A A A A UNSCREWING	AAAA IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM AAAAA UNSCREWING	AAAA IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM AAAAA UNSCREWING	A A A A IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM A A A A UNSCREWING	AAAA IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM AAAAA UNSCREWING	AAAA IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM AAAAA UNSCREWING	AAAA IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM AAAAA UNSCREWING
AAAA IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM AAAAA UNSCREWING 20:1	△△△△ IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM △△△△△ UNSCREWING 20:1		A A A A IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM A A A A UNSCREWING 20:1	△△△△ IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM △△△△△ UNSCREWING 20:1	AAAA IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM AAAA UNSCREWING 20:1	AAAA IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM AAAAA UNSCREWING 20:1
A A A A IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM A A A A UNSCREWING	AAAA IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM AAAAA UNSCREWING	AAAA IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM AAAAA UNSCREWING	A A A A IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM A A A A UNSCREWING	AAAA IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM AAAAA UNSCREWING	AAAA IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM AAAAA UNSCREWING	AAAA IMPLANT SCREWING 20:1 35.3 Ncm 15 RPM AAAAA UNSCREWING

Default values

Endo : default values

The table shows the default operating values for the endodontics sequence.

ENDODONTICS
OPEN PULP CHAMBER 1:5
7.20 mNm
100'000 RPM
ENDO FILE 1
1:1
30.2 mNm
250 RPM
ENDO FILE 2
1:1
10.1 mNm
250 RPM
ENDO FILE 3
1:1
14.9 mNm
250 RPM
ENDO FILE 4
1:1
20.2 mNm
250 RPM
ENDO FILE 5
1:1
30.2 mNm
250 RPM
ENDO FILE 6
1:1
20.2 mNm
250 RPM
1:1
14.9 mNm
250 RPM
ENDO FILE 8
1:1
14.9 mNm
250 RPM
ENDO FILE 9
1:1
10.1 mNm
250 RPM

Surgery : default values

The default values of the 4 available procedures are representative of settings commonly used by clinicians for surgical procedures.

SURGERY					
PROCEDURE 1	PROCEDURE 2	PROCEDURE 3	PROCEDURE 4		
1:5	1:2	1:5	1:5		
0.72 Ncm	2.40 Ncm	0.72 Ncm	0.72 Ncm		
100'000 RPM	80'000 RPM	50'000 RPM	100'000 RPM		

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