### **Instructions for Use**







PB-510, PB-520, PB-530

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WARNING! (risk of injury)



ATTENTION! (to prevent damage occurring)



General explanations, without risk to persons or objects

### Symbols on the control unit



Manufacturer

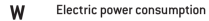


CE mark with identification number of the Notified Body



Foot control











Coolant volume



Upper limit of temperature



Date of manufacture

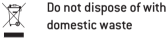
Follow Instructions for Use



Non-ionizing electromagnetic radiation



Catalogue number





Serial number





DC – direct current



MEDICAL — GENERAL MEDICAL EQUIPMENT AS TO ELECTRICAL SHOCK, FIRE AND MECHANICAL HAZARDS ONLY IN ACCORDANCE WITH ANSI/AAMI ES 60601-1:2005, ANSI/AAMI ES60601-1: A1:2012 + C1:2009/[R]2012 + A2:2010/[R]2012, CAN/CSA-C22.2 No. 60601-1:2008, CSA CAN/CSA-C22.2 NO. 60601-1:2014. 25UX — Control No.

### Symbols on the foot control



CE mark with identification number of the Notified Body



Non-ionizing electromagnetic radiation



Catalogue number



Do not dispose of with domestic waste



DC - direct current



Serial number



DataMatrix Code for product information including UDI (Unique Device Identification)



Protection against dripping water



Date of manufacture



UL Component Recognition Mark indicates compliance with Canadian and U.S. requirements



Wireless foot control C-NW



Reset



Manufacturer

### **Symbols**

### radio symbol on the control unit/foot control



GITEKI (MIC) – Japan



RCM - Australian / New Zealand



ANATEL - Brazil



IC - South Korea

MSIP-CRM-BGT-BGM113

Contains FCC ID: QOQBGM113 Contains IC: 5123A-BGM113  $\mathsf{FCC} \, / \, \mathsf{IC} - \mathsf{USA} \, / \, \mathsf{Canada}$ 

## Symbols on the packaging



CE mark with identification number of the Notified Body



This way up



Fragile, handle with care



Keep dry



»Der Grüne Punkt« (The Green Dot) trademark of Duales System Deutschland GmbH



Trademark of RESY OfW GmbH for identification of recyclable transport and outer packaging of paper and cardboard



DataMatrix Code for product information including UDI (Unique Device Identification)



Data structure in accordance with Health Industry Bar Code



Temperature limitation



**Humidity limitation** 



Caution! According to Federal law, this medical device may only be sold by or on the order of a dentist, physician or any other medical practitioner licensed by the law of the State in which he or she practices and intends to use or order the use of this medical device.

#### 1. Introduction

#### For your safety and the safety of your patients

These Instructions for Use explain how to use your medical device. However, we must also warn against possible hazardous situations. Your safety, the safety of your team and, of course, the safety of your patients are of paramount importance to us.



Observe the safety notes.

#### Intended use

PB-510, PB-520, PB-530:

Drive unit with a piezoceramic oscillating system, which moves the tip in a linear oscillation. The drive unit is used for the removal of supragingival calculus and subgingival concretions and for endodontic application and preparation of tooth enamel.

C-NF, C-NW: Foot control for operation of medical electrical equipment.



Misuse may damage the medical device and hence cause risks and hazards for user and third parties.

#### **Qualifications of the user**

We have based our development and design of the medical device on the dentists, dental hygienists, dental employees (prophylaxis) and dental assistants target group.

### Introduction



Production according to EU Directive
The medical device meets the requirements of Directive 93/42/EEC.

0297 The foot control meets the requirements of Directive 93/42/EEC and RED Directive 2014/53/EU.

#### Responsibility of the manufacturer

The manufacturer can only accept responsibility for the safety, reliability and performance of the medical device when it is used in compliance with the following directions:

- The medical device must be used in accordance with these Instructions for Use.
- Only the components approved by the manufacturer may be replaced (0-ring, coolant filter, pump cartridge).
- Modifications or repairs must only be undertaken by an authorised W&H service partner (see page 49).
- The medical device has no components that can be repaired by the user.
- The electrical installation at the premises must comply with the regulations laid out in IEC 60364-7-710 (»Installation of electrical equipment in rooms used for medical purposes«) or with the regulations applicable in your country.
- Unauthorized opening of the medical device invalidates all claims under warranty and any other claims.

Improper use, unauthorized assembly, modification or repair to the medical device, non-compliance with our instructions or the use of accessories and spare parts which are not approved by W&H, invalidates all claims under warranty and any other claims.



Any serious incident that has occurred in relation to the medical device should be reported to the manufacturer and the competent authority!

### 2. Electromagnetic compatibility (EMC)



Medical electrical equipment is subject to particular precautions in regard to EMC and must be installed and put into operation in accordance with the EMC notes included.

W&H guarantees the compliance of the device with the EMC requirements only when used with original W&H accessories and spare parts. The use of accessories and spare parts not approved by W&H can lead to an increased emission of electromagnetic interference or to a reduced resistance against electromagnetic interference.

#### HF communication equipment

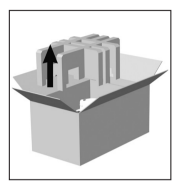
Portable HF communications equipment (including peripherals such as antenna cables and external antennas) should be used no closer than 30 cm (11.8 inch) to the medical device. Otherwise, degradation of the performance of this medical device could result.

The medical device may be interfered by other equipment, even if these other devices comply with CISPR (International special committee on radio interference) emission requirements.

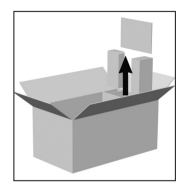
Use of this medical device adjacent to or stacked with other equipment should be avoided because it could result in improper operation. If such use is necessary, this medical device and the other equipment should be observed to verify that they are operating normally.

The medical device is not intended for use in the vicinity of HF surgical devices.

## 3. Unpacking



Remove the insert.



Remove the Instructions for Use and the accessories.



Remove the control unit, the coolant tank and the foot control.

W&H packaging is environmentally friendly and can be disposed of by industrial recycling companies. However, we recommend that you keep the original packaging.

## 4. Scope of delivery

	Control unit (100–240 V)	PB-510 30323000	PB-520 30324000	PB-530 30325000
REF 02675000	Coolant filter	X		
REF 05075600	Coolant hose (0 6 mm, approx. 2 m)	Х		
REF 08016690	Power supply with adaptor	Х	Х	Х
REF 07991190	Coolant tank		Х	Х
REF 08014700	Cable (pairing/charging)			Х

	Optional
REF 30316000	Foot control C-NW
REF 04717300	Foot control C-NF
REF 30326000	Handpiece PB-5 L
REF 30327000	Handpiece PB-5 L Q
REF 30328000	Handpiece PB-5 L S

### 5. Safety notes

#### Control unit/Foot control



- > Before using the medical device for the first time, store it at room temperature for 24 hours.
- > Check the medical device for damage and loose parts each time before using.
- > Do not operate the medical device if it is damaged.
- > Always ensure the correct operating conditions and cooling function.
- > Always ensure that sufficient and adequate cooling is delivered and ensure adequate suction (the exception are endodontics applications).
- > In case of coolant supply failure, the medical device must be stopped immediately. The exception to this is in endodontic applications, where coolant is not used.
  - Maximum operating time without coolant:
- > 2 minutes for the power range 1–30
- > 30 seconds for the power range 31-40
- > Perform a test run each time before using.
- > Never touch the patient and the electrical contacts on the medical device simultaneously.
- > Check the parameter settings every time you restart.
- > Make sure that the supply hose is dry. Moisture in the supply hose can lead to a malfunction (risk of short circuit).
- > Replace damaged or leaking 0-rings immediately.



> Do not twist, kink or squeeze the supply hose (risk of damage).



The medical device is classed as »conventional equipment« (closed equipment without protection against the ingress of water).



The medical device is not approved for operation in potentially explosive explosive atmospheres.



Disconnect the medical device in dangerous situations from the power supply.

> Pull the power supply out of the socket.



> Only use the cable supplied for the foot control (C-NW).

#### System failure

A total system failure does not constitute a critical fault.

Pull the power supply out of the socket and then connect again.

Safety notes Control unit



#### **Control unit PB-510**

- > Disconnect the medical device from the water supply connection after each use (the medical device does not have an automatic aquastop).
- > The medical device is approved for use only with supply units with category 5 backflow prevention devices as defined in EN 1717.
- > Do not connect the medical device to the hot water supply (>30°C/86°F).



#### Control unit PB-520, PB-530

- > Never fill or top up the coolant tank with liquids hotter than 30°C/86°F.
- > Replace a faulty or leaky pump cartridge immediately.



# Control unit PB-510, PB-520, PB-530 Risks due to electromagnetic fields

The functionality of implantable systems, such as cardiac pacemakers and implantable cardioverter defibrillators (ICD) can be affected by electric, magnetic and electromagnetic fields. This medical device complies with the reference values defined in EN 50527-2-1/2016 for unipolar and bipolar pacemakers and is therefore suitable for patients with pacemakers.

- > Find out if patient and user have implanted systems before using the medical device and consider the application.
- > Keep a safe distance of at least 10 cm (3.94 inch) between the medical device and the cardiac pacemakers.
- > Make appropriate emergency precautions and take immediate action on any signs of ill-health.
- > Symptoms such as raised heartbeat, irregular pulse and dizziness can be signs of a problem with a cardiac pacemaker or ICD.



The control unit is designed for use with the W&H handpiece PB-5 L/L S/L Q so only this is to be used with the control unit. The use of other handpieces could lead to a malfunction or destruction of the electronics.

Safety notes Foot control



- > Keep the foot control (C-NW) away from magnetic fields.
- > Replace the foot control as soon as the resistance is noticeably reduced.



> Do not expose the medical device to any violent mechanical impacts.

#### Battery (C-NW)



- > Do not charge the battery unattended.
- > As soon as the charging cycles start to deteriorate send the medical device to an authorized W&H service partner.
- > Defective or worn-out batteries must only be replaced by an authorized W&H service partner.



- > Charge the battery of the foot control as soon as the status LED flashes.
- > Incorrect use of the rechargeable battery can cause fire or corrosion.

Safety notes Foot control C-NW



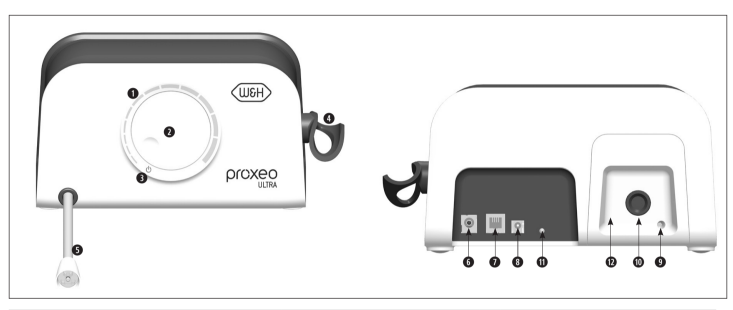
#### Foot control C-NW

#### Risks due to electromagnetic fields

The functionality of implantable systems, such as cardiac pacemakers and implantable cardioverter defibrillators (ICD) can be affected by electric, magnetic and electromagnetic fields. This medical device complies with the reference values defined in EN 50527-2-1/2016 for unipolar and bipolar pacemakers and is therefore suitable for patients with pacemakers.

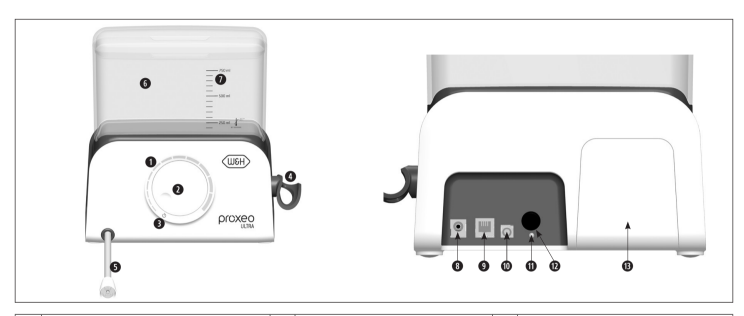
- > Find out if patient and user have implanted systems before using the medical device and consider the application.
- > Keep a safe distance of at least 7 cm (2.76 inch) between the medical device and the cardiac pacemakers.
- > Make appropriate emergency precautions and take immediate action on any signs of ill-health.
- > Symptoms such as raised heartbeat, irregular pulse and dizziness can be signs of a problem with a cardiac pacemaker or ICD.

6. Description Control unit PB-510



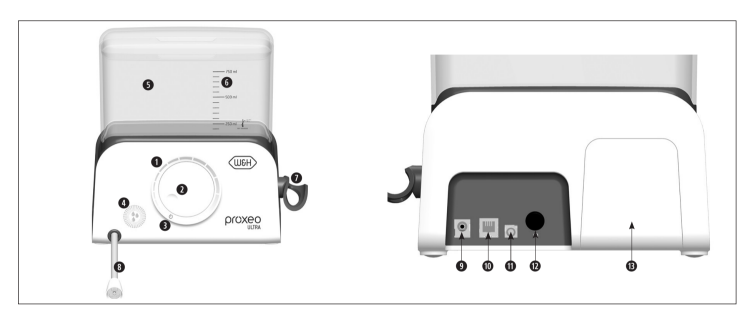
0	Power range		Connections	0	Status LED		
2	Power regulator		Power supply ESI (external service interface)	0	Cover		
3		_	Foot control				
4	Handpiece support (adjustable)		9	9	Coolant hose		
6	Supply hose	0	Coolant regulator				

**Description** Control unit PB-520



0	Power range	6	Coolant tank	0	Status LED
2	Power regulator	0	Filling level indicator	0	Coolant regulator
3	»0FF«		Connections	ß	Cover
4	Handpiece support (adjustable)	8	Power supply ESI (external service interface)		
6	Supply hose	0	Foot control		

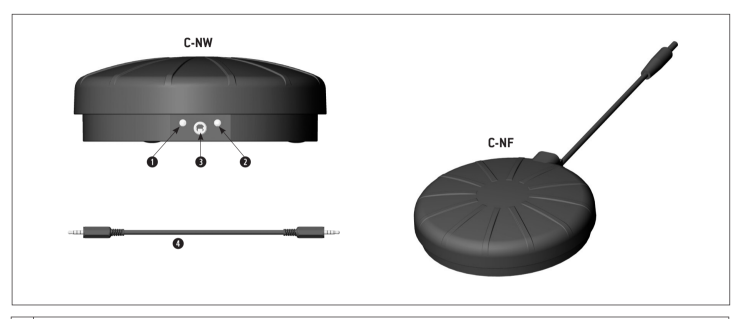
**Description** Control unit PB-530



0	LED display	2	Power regulator	6	Coolant tank		Connections
	> Power range > Battery status foot control	8	»OFF«	6	Filling level indicator	0	Power supply ESI (external service interface)
	> Error message	4	Function button	0	Handpiece support (adjustable)		Cable (pairing/charging)
	> Rinsing function > Cleaning function		> Rinsing function > Cleaning function	8	Supply hose	<b>@</b>	Coolant regulator
	> Pairing		> Pairing			₿	Cover

**Description** 

### Foot control C-NF/C-NW



- Charging LED (orange)
- 2 Status LED (green)
- 3 Connection for cable (pairing/charging)
- 4 Cable (pairing/charging)



#### Standby mode

> The foot control can be activated by pressing.

LED	steady	steady	flashes	flashes intermittently*		
	• @ •	● ⊖ ●	<b>6</b>	• • •		
GREEN		→ Connection to paired medical device established	→ Foot control is attempting to establish a connection to the paired medical device	→ Battery is flat > Charge the battery		
ORANGE	→ Battery is charging					
* LED flashes for	* LED flashes for 40 milliseconds every 4 seconds					

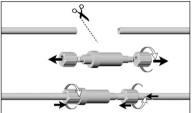
### Control unit general

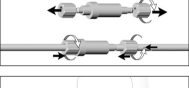


Ensure that the medical device can be disconnected from the power supply at any time.

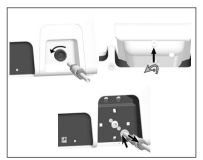


Place the medical device on a flat, level surface.









#### Control unit PB-510 Mount the coolant filter

- Cut through the coolant hose.
- 2 Unscrew the cap nut from the coolant filter.
- Attach the coolant hose through the cap nut onto the coolant filter. Screw the cap nut tight.
- Push the coolant hose until the limit stop.

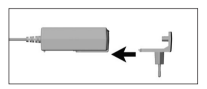
#### Control unit PB-510 Remove the coolant hose

- Screw off the coolant regulator.
- Uncrew the cover and remove it.
- Push the connection ring and simultaneously remove the coolant hose.



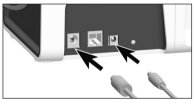
#### Control unit PB-520, PB-530 Coolant tank

• Fill the coolant tank and attach it. The coolant tank snaps audibly into place.



#### Control unit PB-510, PB-520, PB-530

Slide the adapter onto the power supply.

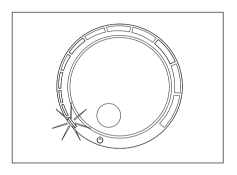


- 2 Connect the power supply.
- 3 Connect the foot control C-NF (control units PB-510, PB-520).



- Plug the power supply into a socket.
- **5** Pull the power supply out of the socket.

Start-up Control unit PB-530



#### **Control unit PB-530**

Power regulator »OFF«

> 1. LED display flashes white

#### Next steps:

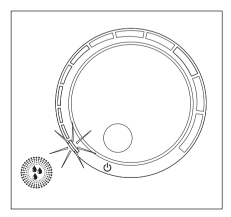
- > Pairing foot control C-NW with control unit PB-530
- > Charging the battery of the foot control C-NW with control unit PB-530



Coolant and handpiece inactive



The foot control C-NW and control unit PB-530 are not paired when delivered!



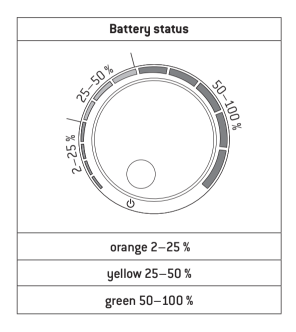
#### Pairing foot control C-NW with control unit PB-530

- Set power regulator to »OFF«
- Connect the cable to the control unit and foot control
- > 1. LED flashs orange/red = not paired
- 3 Press function button for 5 seconds
- > Sequential white LED during pairing
- > 1. LED flashes white = pairing successful

Start-up Control unit PB-530



Charge the foot control fully before you use it for the first time.



#### Charging foot control C-NW with control unit PB-530

- Connect the cable to the control unit and foot control
- > Power regulator »OFF«: The charging level is visible for 5 seconds at the LED display.



Query battery status during charging process with one of these options:

- > Press foot control, battery status visible for 5 seconds
- > Press function button, battery status visible for 5 seconds
- > Disconnect charging cable, battery status visible for 5 seconds
- > Set power
- > Power set: The battery status is visible on the LED display.

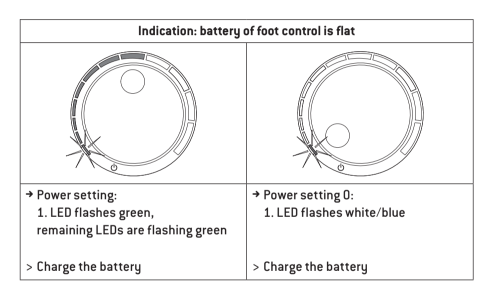


During the charging process the LED display flashes.

The LED display flashes completely when the battery is charged.



Control unit and foot control are not connected with the cable.



Proxeo Ultra	PB-510	PB-520	PB-530
Rinsing function for automatic internal cleaning of the coolant channels	<b>V</b>	<b>V</b>	<b>V</b>
Cleaning function for automatic internal cleaning of the coolant channels	-	-	V



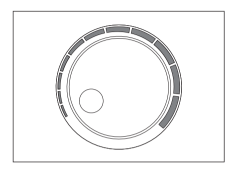
Before every patient: Perform rinsing function for automatic internal cleaning of the coolant channels.

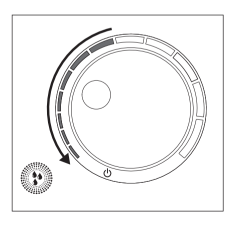
#### Approved coolants and rinsing liquids

- > Physiological saline solution (NaCl, 0.9%)
- > Hydrogen peroxide  $(H_2O_2, 1-3\%)$
- > Liquids with the active substance chlorhexidine (CHX, 0.2%)
- > Tap water



W&H recommends performing a rinsing function with tap water after using one of the approved liquids.





#### Control unit PB-510, PB-520

- Remove the handpiece from the supply
- 2 Set power to 0
- 3 Press foot control 3 times within 3 seconds
- > Rinsing function active for 30 seconds



Cancelling the rinsing function with one of these options:

- > Press foot control
- > Adjust power regulator

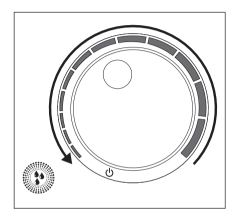
#### Control unit PB-530

- Remove the handpiece from the supply
- Adjust power
- Press funtion button 1 second
- > Rinsing function active for 30 seconds, visible by blue LEDs
- > Rinsing function finished after all blue LEDs go out



Cancelling the rinsing function with one of these options:

- > Press foot control
- > Press funtion button 1 second
- > Set power regulator to »OFF«



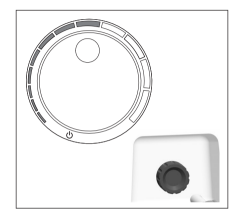
#### Control unit PB-530

- Remove the handpiece from the supply hose
- 2 Set power
- 3 Press funtion button 3 seconds
- > Cleaning function active for 8 minutes, visible by blue LEDs
- > Pump stops several times during the cleaning function
- > Cleaning function finished after all blue LEDs go out



Cancelling the cleaning function with one of these options:

- > Press foot control
- > Press function button 1 second
- > Set power regulator to »OFF«



#### Control unit PB-510, PB-520, PB-530

• Attach the handpiece to the supply hose. Insert tip.

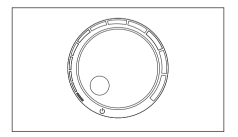


Follow the directions and safety notes in the Instructions for Use of the W&H handpieces.

- 2 Set power and coolant (variable)
- Press foot control
- > Release foot control: Fade-out time of the handpiece LED 30 seconds

#### Control unit PB-530

> Coolant in coolant tank < 50 ml: handpiece LED flashes



## Subgingival flushing Control unit PB-530

- Set power to 0
- > 1. LED flashes blue
- Press foot control

### Test run



Do not hold the handpiece at eye level!

- > Attach the handpiece to the supply hose.
- > Insert the tip.
- > Put the medical device into operation.



In the event of operating malfunctions (e.g., vibrations, unusual noise, overheating, coolant failure or leakage) **stop the medical device immediately** and contact an authorized W&H service partner.

## 9. Error messages



The error messages are indicated at the rear of the control unit by the status LED (flashing green).

Flashing cycle	Description of error	Solution
1x	Overheating	> Switch off conrol unit > Allow the control unit to cool for at least 10 minutes > Observe permissible ambient temperature/operating mode
2x	Foot control	> Release foot control
5x	Time-out (> 15 min)	> Release foot control (must not be active for longer than 15 minutes without interruption)
6x	Handpiece	Check tip     (full engagement, damage, torque)      Dry the handpiece/supply hose      Check plug-in connection of the handpiece/supply hose      If the error message appears again, contact an authorized W&H service partner immediately.
8x	System error	> Start the medical device again > Contact an authorized W&H service partner.

Error messages Control unit PB-530



The error messages are indicated by the LED display (LED steady).

LED display	Colour	Description of error	Solution
1. LED	orange	Overheating	<ul> <li>Switch off control unit</li> <li>Allow the control unit to cool for at least 10 minutes</li> <li>Observe permissible ambient temperature/operating mode</li> </ul>
2. LED	orange	Foot control	> Release foot control
4. LED	orange	Function button	> Release function button
5. LED	orange	Time-out (> 15 min)	> Release foot control (must not be active for longer than 15 minutes without interruption)
6. LED	orange	Handpiece	<ul> <li>Check tip         (full engagement, damage, torque)</li> <li>Dry the handpiece/supply hose</li> <li>Check plug-in connection of the handpiece/supply hose</li> <li>If the error message appears again, contact an authorized W&amp;H service partner immediately.</li> </ul>
12. LED	red	System error	Start the medical device again     If the error message appears again, contact an authorized W&H service partner immediately.

If the error messages described cannot be resolved, a check by an authorized service partner is required.

> In case of a total system failure, switch the medical device off and on again.



Follow your local and national laws, directives, standards and guidelines for cleaning and disinfection.



> Wear protective clothing, safety glasses, face mask and gloves.



> Wipe the entire medical device and the foot control with disinfectant.



> Ensure that no fluids enter the medical device.



> Note that the disinfectant used during pre-treatment is only for personal protection and cannot replace the disinfectant step after cleaning.



W&H recommends performing the rinsing function (PB-520) or the cleaning function (PB-530) using an approved cleaning agent according to the manufacturer's instruction. Fill coolant tank with at least 200 ml of liquid.

#### Approved cleaning agents

- > Citrisil™ (Sterisil, Inc.)
- > Bilpron (ALPRO MEDICAL GMBH)



W&H recommends performing a rinsing function with tap water after using one of the approved liquids.

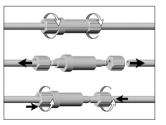
#### 11. Maintenance

Proxeo Ultra	PB-510	PB-520	PB-530
Replacing the 0-ring of the coolant tank	_	V	V
Replacing the coolant filter of the coolant hose	~	_	_
Replacing the pump cartridge	_	~	~



#### Replacing the O-ring of the coolant tank

- Remove the 0-ring with tweezers.
- 2 Slide on the new 0-ring.



#### Replacing the coolant filter of the coolant hose



Replace the coolant filter if it is soiled or after 1 year at the latest.

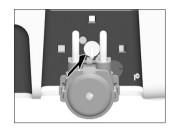
- 1 Unscrew the cap nut from the coolant filter.
- 2 Pull off the coolant hose from the coolant filter.
- 3 Attach the coolant hose through the cap nut onto the new coolant filter. Screw the cap nut tight.

#### Maintenance

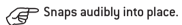
#### Replacing the pump cartridge

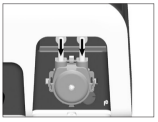


Unscrew cover and remove.

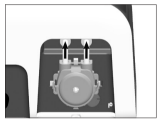


• Attach new pump cartridge.

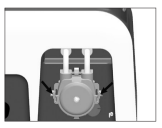




2 Pull off coolant hoses.



• Attach the coolant hoses until the limit stop.



3 Unlock pump cartridge and pull it out.



6 Attach cover and screw tight.

# 12. Servicing



#### Regular checks

Regular servicing of function and safety including the accessories is necessary and should be carried out at least once every three years, unless shorter intervals are prescribed by law.

The inspection must be undertaken by a qualified organisation and must include the following procedures:

#### **Control unit**

- > External visual inspection
- > Measurement of the device leakage current
- > Measurement of patient leakage current
- > Visual inspection of internal components on suspicion of safety interference, e.g., mechanical damage of the enclosure or indicators of overheating

#### Foot control

- > External visual inspection
- > Function test with check to see if the maximum speed can be reached



The regular checking must only be performed by an authorised W&H service partner.

# **Servicing**

#### Repair and returns

In the event of operating malfunctions immediately contact an authorized W&H service partner. Repairs and maintenance work must only be undertaken by an authorized W&H service partner.



> Always return equipment in the original packaging!

# 13. W&H accessories and spare parts



Use only original W&H accessories and spare parts or accessories approved by W&H. **Suppliers:** W&H partners



30326000 Handpiece PB-5 L 30327000 Handpiece PB-5 L Q 30328000 Handpiece PB-5 L S



02675000 Coolant filter 05075600 Coolant hose



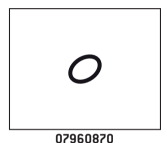
08001660 Pump cartridge



07991190 Coolant tank



08014700 Cable (pairing/charging)



0-ring for coolant tank



08016690 Power supply with adaptor

# W&H accessories and spare parts



30316000 Foot control C-NW with Stick



04717300 Foot control C-NF

#### 14. Technical data

Control unit	PB-510	PB-520	PB-530		
Power supply:		28.5-31.5 V <del></del>			
Mains voltage:		100-240 V			
Nominal current:		max. 830 mA			
Permissible voltage fluctuation:		±10%			
Max. output power to the handpiece under load (ultrasonic):		12 W			
Frequency (ultrasonic):		22-35 kHz			
Operating mode:		S3 (14sec/6sec)			
Max. oscillation amplitude (Tip 1U):	0.2 mm				
Max. water pressure:	1–6 bar				
Max. coolant flow (adjustable):	ca. 50 ml/min				
Dimensions in mm (WxDxH):	120 x 185 x 110	120 x 185 x 205	120 x 185 x 205		
Weight:	807 g	1,064 g	1,106 g		

#### **Ambient conditions**

Temperature during storage and transport:  $-20^{\circ}\text{C to } +60^{\circ}\text{C (-4°F to } +140^{\circ}\text{F)}$  Humidity during storage and transport: 8% to 80% (relative), non-condensing Temperature during operation:  $+10^{\circ}\text{C to } +35^{\circ}\text{C (}+50^{\circ}\text{F to } +95^{\circ}\text{F)}$ 

Humidity during operation: 15% to 80% (relative), non-condensing

#### Technical data

Foot control	C-NW
Battery type:	Li-lon
Runtime:	approx 2 months
Standby:	automatically if not actuated
Charging time:	approx. 3 h
Nominal voltage:	3.7 V
Nominal capacity:	680 mAh
Dimensions (WxDxH):	117 x 117 x 38 mm
Weight:	190 g

#### **Ambient conditions**

Temperature during storage and transport: Humidity during storage and transport: Temperature during operation:

Humidity during operation:

-20°C to +60°C (-4°F to +140°F)

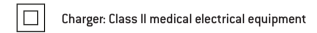
8% to 80% (relative), non-condensing

+10°C to +35°C (+50°F to +95°F)

15% to 80% (relative), non-condensing

#### Technical data

Classification according to Paragraph 6 of the General Specifications for the Safety of Medical Electrical Equipment according to IEC 60601-1/ANSI/AAMI ES 60601-1





The C-NF/C-NW foot control is protected against vertically falling drops of water (IPX1 as per IEC 60529)

Pollution level: 2
Overvoltage category: II

Altitude: up to 3,000 m above sea level

# 15. Disposal



Ensure that the parts are not contaminated on disposal.



Follow your local and national laws, directives, standards and guidelines for disposal.

- > Medical device
- > Waste electrical equipment
- > Packaging

# Explanation of warranty terms

This W&H medical device has been manufactured with great care by highly qualified specialists. A wide variety of tests and controls guarantees faultless operation. Please note that claims under warranty can only be validated when all the directions in the Instructions for Use have been followed.

As the manufacturer, W&H is liable for material or manufacturing defects within a warranty period of 24 months from the date of purchase. Accessories and consumables (pump cartridge, coolant hose, coolant filter, 0-rings) are excluded from the warranty.

We accept no responsibility for damage caused by incorrect handling or by repairs carried out by third parties not authorized to do so by W&H!

Claims under warranty accompanied by proof of purchase must be sent to the vendor or to an authorized W&H service partner. The provision of service under warranty extends neither the warranty period nor any other guarantee period.

# Authorized W&H service partners

Find your nearest authorized W&H service partner at http://wh.com Simply go to the menu option »Service« for full details.

Or simply scan the QR code.



# Manufacturer's declaration

Electromagnetic compatibility (EMC)
WARNING: The use of realise, power supplies, accessories other than those specified by the manufacturer increased entision and/or decreased immufily. Only use original W&H accessories.

reference Manufacturer: W&H GmbH REF: 30316xxx Manufacturer: W&H GmbH REF: 04717300 Manufacturer: GlobTek, Inc. REF: 08016690 1.8 m power supply (GTM96300-3036-6.0-R2) cables and accessories foot controller C-NW foot controller C-NF

Operate the product in a place with a maximum distance to electrical and magnetic interfering transmitters. If operation of the product close to other devices or together in a stack is necessary, observe the correct function of the system.

assure that it is used in an electromagnetic environment as described below.	n electromagnetic en	vironment as descrit	ped below.	process is sustained in an electromagnetic environment as described below.
Immunity Test	IEC 60601-Level	IEC 60601-Level	Compliance	Electromagnetic Environment
	(3rd Ed.)	(4th Ed.)	Level	Guidance
Electrostatic	±6 kV contact	±8kV contact	± 8 kV contact	Floor should be wood, concrete or
discharge (ESD)	±8 kV air	± 15 kV air	± 15 kV air	ceramic tile. If floors are covered
IEC 61000-4-2				with synthetic material, the relative
1				number of the street of the st
Electrical tast	± 2 kV for power	± 2 kV tor power	± 2 kV tor power	Mains power quality should be that
transient/bursts	supply lines	snbbiy lines	supply lines	of a typical commercial and/or
IEC 61000-4-4	±1 kV for	± 1 kV for	± 1 kV for	hospital environment
	input/output lines	input/output lines	input/output lines	
	5kHz repetition	100kHz repetition	Both repetition	
	rate	rate	rates	
Surge	±1 kV	±1kV	±1kV	Mains power quality should be that
IEC 61000-4-5	line(s) to line(s)	line(s) to line(s)	line(s) to line(s)	of a typical commercial and/or
				hospital environment
	±2 kV	±2kV	±2kV	
	line(s) to earth	line(s) to earth	line(s) to earth	
Voltage dips, short	<5% U₁	0% U⊤ 0.5 cycle	Complies to both	Mains power quality should be that
interruptions and		(e)	editions	of a typical commercial and/or
voltage variations on	for 0.5 cycle	0°,45°,90°,135°,1	requirements	hospital environment. If the user of
power supply input		80°,225°,270° &		the product requires continued
lines IEC61000-4-11	40% U <sub>T</sub>	315°		operation during power mains
	(60% dip in U⊤)			interruptions, it is recommended that
	for 5 cycles	0% U₁ 1 cycle		the product be powered from an
		and		uninterruptible power supply or a
	70% U <sub>T</sub>	70% U <sub>T</sub> 25/30*		battery.
	(30% dip in U <sub>T</sub> )	cycles @ 0°		
	for 25 cycles			
		0% U <sub>T</sub> 250/300*		
	<5% U₁	cycle		
	(>95% dip in U⊤)			
	for 5 sec			
Power	3A/m	30A/m	30A/m	Power frequency magnetic fields
frequency(50/60 Hz)				should be at levels characteristic of
IIIagileiic lioid				a typical location in a sypical

Condicioned RF   13 Vm   15
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Test frequency	Band <sup>a)</sup>	Service <sup>a)</sup>	Modulation <sup>b)</sup>	Maximum power	Distance	IMMUNITY TEST LEVEL
(MHz)	(MHz)			(W)	(m)	(M/M)
385	380 -390	TETRA 400	Pulse modulation <sup>b)</sup> 18 Hz	1.8	0.3	27
450	430 – 470	GMRS 460, FRS 460	FM <sup>c)</sup> ± 5 kH deviation 1 kHz sine	2	0.3	28
710		CF 7 G LE	Pulse			
745	704 – 787	LIE Dang 13,	modulation <sup>b)</sup>	0.2	0.3	6
780			217 Hz			
810		GSM 800/900,				
870	800 – 960	TETRA 800, IDEN 820, CDMA 850.	Pulse modulation <sup>b)</sup> 18 Hz	2	0.3	28
930		LTE Band 5				
1720		GSM 1800;				
1845	1700 – 1990	GSM 1900; DECT;	Pulse modulation <sup>b)</sup> 217 Hz	2	0.3	28
1970		LTE Band 1, 3, 4, 25; UMTS				
2450	2400 – 2570	Bluetooth, WLAN, 802.11 b/g/n, RFID 2450, LTE Band 7	Pulse modulation <sup>b)</sup> 217 Hz	2	0.3	28
5240			Pulse			
5500	5100 - 5800	WLAN 802.11 a/n	modulation <sup>b)</sup>	0.2	0.3	6
5785			Z17 HZ			

<sup>&</sup>lt;sup>a)</sup> For some services, only the uplink frequencies are included.
<sup>b)</sup> The carrier shall be modulated using a 50 % duty cycle square wave signal.
<sup>c)</sup> As an alternative to EM modulation, 50 % pulse modulation at 18 Hz may be used because a continuous of the continu

Recommended Separation Distances between portable and mobile HF-communications equipment and the product (Table 6, IEE 60601-12:2007) product (Table 6, IEE 60601-12:2007) product (Table 6, IEE 60601-12:2007) crossioner of the product is hierarcely of use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer of the user of the product can help pevent electromagnetic interference by maintaining a minimum distance between potable and mobile RF promunications equipment (Idansmittels) and the product – according on output power and

smitter in meter (m) 800 MHz to 2.5 GHz	d = 2.3√P	0,23	0,73	2,3	7,3
y of transmitter ii 800 MHz	Ψp		1		
Rated maximum output power of Separation distance according to the frequency of transmitter in meter (m 150 kHz to 80 MHz to 800 MHz to 800 MHz to 2.5 GHz to 800 MHz	$d = 1.2\sqrt{P}$	0,12	0,38	1,2	3,8
Separation distance 150 kHz to 80 MHz	d = 1.24P	0,12	0,38	1,2	3,8
Rated maximum output power of transmitter in watts (W)		0,01	0,1	1	10

For transmitters rated at a maximum output power not listed above, the recommended separation distance d in meters (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 maximum output power

Note 1: At 80 MHz and 800MHz, 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, people and animals.

Electromagnetic Emission (Table 1, IEC 60601-1-2:2007)
The product is suitable for use in a specific electromagnetic.

THE PROGRESS SURBING TO USE III &	specific electroniagnetic en vironiment.	The product is suitable for use in a specific electroning letic environment. The customer and of the product should
assure that it is used in an electrom	assure that it is used in an electromagnetic environment as described below.	
Emission Test	Compliance	Electromagnetic Environment Guidance
RF-emission	Group 1	The product use RF energy only for its internal
CISPR 11		function. Therefore, its RF emissions are very
		low and not likely to cause any interference in
		nearby electronic equipment.
		However, a separation distance of 30 cm shall
		be maintained.
RF-emission	Class B	The product is suitable for use in all
CISPR 11		establishments, including domestic
Harmonic emissions	N/A (P<75W)	establishments and those directly connected to
IEC 61000-3-2 (1)		the public low-voltage power supply network
Voltage fluctuations/	N/A (P<75W)	that supplies buildings used for domestic
flicker emissions		purpose.
IEC 61000-3-3 (1)		
A Property Constitution of the constitution of	W CO CO - 1111 - 11 - 11 - 11 - 11 - 11 -	

#### Manufacturer

**W&H** Dentalwerk Bürmoos GmbH Ignaz-Glaser-Straße 53, 5111 Bürmoos, **Austria** 

t +43 6274 6236-0, f +43 6274 6236-55 office@wh.com wh.com Form-Nr. 50968 AEN Rev. 002 / 01.10.2020

Subject to alterations