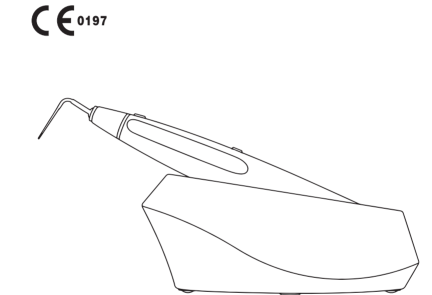
文件名/NAME	热熔牙胶充填机英文说明书 Fi-P CE	代码/code 14.02.11.037
尺寸/SIZE	130×190mm,出血 6mm	版本/REV. V1.6
材质/MATERIAL	120g 铜版纸	
工艺/PROCESS	/	
装订&注释/ bind (books etc) &NOTES	骑马订 保修卡正反面内容需对应,印刷于同一页纸上。	印刷颜色/COLORS 4X4 ■CMYK
修订日期/DATE	2024年07月24日	

请勿打印此页,仅供参考。/DO NOT PRINT THIS PAGE, REFERENCE ONLY.

Gutta Percha Obturation Device Instruction Manual



Fi-P

ZMN-SM-039 V1.6-20240724

Guilin Woodpecker Medical Instrument Co., Ltd.

Contents

Introduction1
1 Product introduction1
2 European authorized representative6
3 Standard icons6
4 Contraindications7
5 Installation and disassembly method of accessories7
6 Operation method9
7 Charging instruction9
8 Safety precautions9
9 Cleaning, Disinfection, Sterilization and Maintenance 10
10 Troubleshooting 16
11 After-sales service 17
12 Environment protection 17
13 EMC-Declaration of comformity 17
14 Statement 21

Introduction

Thank you for purchase Fi-P Heating and Packing Instrument developed by Guilin Woodpecker Medical Instrument Co., Ltd, a Hi-tech enterprise developing, manufacturing, and selling dental instruments. Woodpecker has excellent Quality Control System. To guarantee correct and safe operation, please read this Instruction Manual carefully before use.Depending on the level of risk involved, safety requirements are classed under the following indications:

Danger: (always referred to personal injury)

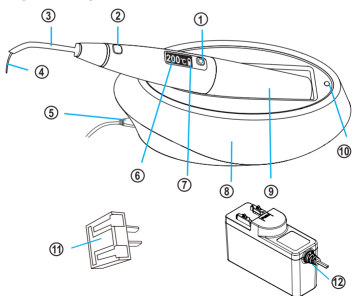
Warning: (referred to possible damage to property)

1 Product introduction

1.1 Intended use

It is used to provide heat to the Work Tip, cut the gutta-percha point, and soften and pressurize the gutta-percha. And the applied part is Work Tip.

1.2 Diagram of components and control buttons



1. "ON/OFF" button

3. Work Tip Protector

5. Connecting hole for power adapter

- 7. Battery level
- 9. Battery cartridge
- 11. Power adapter plug

1) "ON/OFF" button:

- 2. Heating button
- 4. Work Tip
- 6. Temperature Level
- 8. Charging base
- 10. Charging indicator
- 12. Power adapter unit

Under shutdown state, shortly press "ON/OFF" button to start the device.

Under shutdown state, long press "ON/OFF" button to start the device and change the direction of screen display, that is to say, the direction of display can be change to adapt to the operation in left hand or right hand.

Under ON state, long press "ON/OFF" button to shut down the device. (Time for long press is about 1s.)

Note: If there is no operation for 10 minutes, the Heating and Packing Instrument will automatically shut down.

Under ON state, shortly press "ON/OFF" button to change the preset temperature of Work Tip. The preset temperature will change to the next with the sequence $150^{\circ}C \rightarrow 180^{\circ}C \rightarrow 200^{\circ}C \rightarrow 230^{\circ}C$ after each press. And then go back to $150^{\circ}C$ after short press at temperature of $230^{\circ}C$.

150℃월 180℃월 200℃월 230℃월

Figure 1 Preset temperature

2) Heating button:

Under the ON state, connect the Work Tip, and press Heating button to start heating. Release the Heating button to stop heating, followed by the fall of Work Tip temperature.

Note: If press and hold the Heating button for more than 10 seconds, the device will stop heating. If need to continue heating, please release the Heating button and press again.

3) Volume set

After power-on ,press "ON/OFF" button and "Heating" button simultaneously to enter the voice volume setting mode, then short press the "ON/OFF" button to select suitable voice volume, the last, short press the "Heating" button to exit the voice volume setting mode as shown in Figure2.





Figure2a Low volume Figure2b Medium volume Figure2c High volume

4) Battery level:

The actual power of the battery is displayed in real time on the screen. When the battery is fully charged, the power of the OLED display is displayed as five grids. When the battery level is one grid, it indicates that the battery is low and needs to be charged in time. When the battery level is displayed as a space, it indicates that the battery is very low and needs to be charged immediately.

Note: During normal use, try not to let the battery level reduced to space status (completely no power) before charge, which will shorten the service life of battery.

Warning:

If the device has not been used for more than one month, the battery needs to be recharged. If the device is not in use for a long time, please be sure to charge it at least once a month to protect the battery. The service life of battery of Heating and Packing Instrument will be shortened when it is in a low battery state for a long time or when it leaves the charging base for a long time.

5) Temperature Level:

When the temperature is preset, the display screen shows the preset temperature value. About 1s after the temperature preset, the OLED screen will display the real-time temperature of the Work Tip. When the Heating and Packing Instrument is in the heating state, the temperature indicator will simultaneously display the current temperature of the Work Tip.

6) Charging base:

Firstly, connect the power adapter plug to the power adapter as shown in Figure 3. Then connect the power adapter to the power connecting hole on the charging base as shown in Figure 4 and connect the power adapter to a standard socket. Place the Heating and Packing Instrument correctly on the charging base as shown in Figure 5, so that the charging connector under the Heating and Packing Instrument can be reliably connected to the output connector of the charging base. When the Heating and Packing Instrument is properly connected to the charging base, the LED charging indicator on the base will be on constantly. If the LED is flashing or not lit, please check all the cables carefully.

There are charging status indicators on the charging base. When the Heating and Packing Instrument is not placed on the charging base, the indicator will flashes in yellow and green alternately. When the Heating and Packing Instrument is placed on the charging base, if the charging is being charged, the yellow indicator will be on constantly. When the battery is full, the yellow indicator will be off and the green indicator will be on constantly.

Notes: After receiving the device, please charge it immediately. Before use, please be sure that battery is fully charged. When the device is fully charged, the battery level of the Heating and Packing Instrument led display screen is the highest. After the battery runs out, the time of battery charging takes at least 2 hours and 30 minutes.



Figure 3 Installation of power adapter

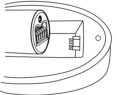


Figure 4 Connection to power supply

Figure 5 Charging

1.3 Device includes

- 1. Heating and Packing Instrument
- 2. Charging base
- 3. Power adapter with cord
- 4. Work Tips (The models are as shown in Table 2)
- 5. Work Tip Protector
- 6. Instruction Manual
- 7. Qualified Certification
- 8. Warranty card
- 9. Packing list

Model	Work Tip	Taper
	Size(mm)	
WP4004	0.40	0.04
WP4504	0.45	0.04
WP5506	0.55	0.06

WP5508	0.55	0.08	
T-1-1-2 M- 4-1 - CW- 1- Tim-			

Table 2 Model of Work Tips

1.4 Introduction and scope of application

1.4.1 Features:

a) The display can be set to both right and left sides, to meet the needs of both left-hander and right-hander.

b) Cordless design for Heating and Packing Instrument effectively broadens the operation space.

c) Sensitive temperature control, simple display, and convenient operation; Press temperature setting button to set suitable working temperature.

d) Four preset temperatures are for option: 150°C, 180°C, 200°C, 230°C.

e) If there is no operation for 10 minutes, the Heating and Packing Instrument will automatically shut down.

1.4.2 Scope of application:

Used in the root canal obturation stage in endodontic treatment.

1.5 Product specifications

Size	Heating and Packing Instrument	23.8mm×158.3mm×23.8mm
Size	Charging base	75.5mm×149.7mm×62.6mm
	Heating and Packing Instrument	80g
Weight	Charging base	195g
	Power adapter	167g

1.6 Technical parameters

Classification	Class II(AC/DC power adapter)		
Optional preset temperatures	150°C→180°C→200°C→230°C		
Time consumption for charging	About 2.5h		
Power supply	Input	AC100V-240V 50/60Hz 800mA	
	Output	DC15V/1.6A	
Battery capacity	Chargeable battery	2000mAh	
Heater Rating	10W		

1.7 Environmental parameters

Temperature: $+5^{\circ}C \sim +40^{\circ}C$ Humidity: 30% ~ 75% Air pressure: 70kPa ~ 106kPa

1.8 Storage and transport

1. The device should be handled carefully and lightly. Be sure that it is far from the vibration, and is installed or kept in a cool, dry, and ventilated place.

2. Do not store the device together with the articles that are combustible poisonous, caustic, or explosive.

3. The device should be stored in a room where the relative humidity is $10\% \sim 93\%$, the air pressure is 70kPa ~ 106 kPa, and the temperature is -20°C ~ +55°C.

4. Please avoid the device from strong shock or vibration during transport. And please handle it carefully.

5. Please do not mix the device with hazardous articles during transport.

6. Please avoid the device from sun, rain, and snow during transport.

2 European authorized representative

ECREP

MedNet EC-REP C IIb GmbH Borkstrasse 10 · 48163 Muenster · Germany

3 Standard icons

SN	Product serial number		Follow Instructions for Use
	Manufacturer	\sim	Date of manufacture
T	Type B applied part		Class II device
Ċ	Power switch	IPX0	Ordinary equipment
	Used indoor only		Caution,hot surface
134℃ ∫	Can be autoclaved	DC 15V	DC 15V

(f 0197	CE marked product		
	- 1		
	Device complies with WE	LEE direct	lve
	Attention! Please refer to t	the accom	panying documents.
93%	Humidity limit for storage: 10% ~ 93%		
70kPa	Atmospheric pressure for storage: 70kPa ~ 106kPa		
-20°C	Temperature limit for storage: $-20^{\circ}C \sim +55^{\circ}C$		
ECREP	Authorised Representative COMMUNITY	e in the EU	JROPEAN

4 Contraindications

1. People who are allergic to known natural latex and metals such as stainless steel, silver, copper, etc. are forbidden to use this device.

- 2. The patient with hemophilia is forbidden to use this device.
- 3. The patients with heart pacemaker are forbidden to use this device.
- 4. The dentists with heart pacemaker are forbidden to use this device.

5. Heart disease patients, pregnant women and children should be cautious to use the equipment.

5 Installation and disassembly method of accessories

5.1 Connection of power adapter

Connect the output point of power adapter to the charging base, and connect the input point to the socket that meets the standard of this power adapter. Please install in accordance with the procedures in Figure 3, Figure 4, and Figure 5. (Note: The installation in Figure 3 had been finished before delivery.)

5.2 Installation and removal of Work Tip

1. After turning off the power switch, you can directly pull the Work Tip off the Heating and Packing Instrument.

- 2. Place the used Work Tip in a certain container and disinfect it.
- 3. Select the desired work Work Tip and the hexagonal plug on the

Work Tip (as indicated by the red arrow in Figure 6). When installing the work Work Tip as shown in Figure 7, you can select the appropriate direction according to the usage to insert the Work Tip into the Heating and Packing Instrument.

4. Install the Work Work Tip Protector to the Work Tip as shown in Figure 8, to prevent scalding patient's mouth during operation.

5. Under ON state, if the Work Tip hasn't been installed or is in poor connection, there would be an error code on display screen as shown in Figure 9.

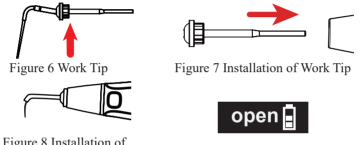


Figure 8 Installation of Work Tip Protector Figure 9 Error code 5.3 Installation and replacement of battery

When replacing the battery, as shown in Figure 10, first rotate the battery barrel counterclockwise to remove the battery tube, then take the old battery out of the battery tube, replace it with a new one, and finally tighten the battery tube clockwise according to the corresponding thread.

Warning: When removing the battery, the screw under the battery barrel (pointed by the arrow in Figure 10) does not need to be unscrewed, just push the connector slightly inward to remove the battery. Improper replacement of lithium batteries may result in unacceptable risks, so replacement of lithium batteries requires trained personnel.

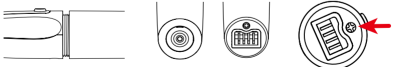


Figure 10 Replacement of battery

6 Operation method

1. According to the situation of patient, select suitable Work Tip and install it. When installing the Work Tip, chose a suitable angle to install the Work Tip.

/ Danger:

Don't turn on the device when installing the Work Tip, to prevent scalding the user by mistakenly pressing the heating button.

2. After pressing the "ON/OFF" button, the display screen of Heating and Packing Instrument lights up and display the preheating temperature and power status.

3. According to the actual situation, lightly press the temperature setting button, and select suitable preheating temperature as per the instruction on display screen.

4. During operation, lightly press the heating button so as to heat up to the preset temperature, soften and pressurizing the gutta-percha with careful, continuous and stable motion with the help of vertical pressurizer.

Note: The continuous heating time on gutta-percha cannot exceed 4s, or there would be risk of scalding.

5. After operation, please clean, disinfect, and sterilize the Work Tip. The specific method is shown in Chapter 9.

7 Charging instruction

7.1 Use the corresponding charging base for this device. Connect the power adapter with the charging base, connect the power supply, and then correctly place the Heating and Packing Instrument into the charging base.

7.2 The battery used in this product has no memory and can be used at any time or charged at any time.

7.3 Before first use of this device, please charge it at least for 3 hours.

Warning: Only unplug the adapter to disconnect from the network power.

8 Safety precautions

1. Do not polish the Work Tip.

- 2. Do not knock or scratch the Heating and Packing Instrument.
- 3. Keep the heating pressurizer, Work Tip, etc. under heating state

away from inflammable and explosive materials.

4. Please keep the device clean before and after operation. Before each use, please disinfect Work Tip and its accessories.

5. The product should be in strict accordance with relevant operation specifications of medical authority and relative regulations. The product can only be operated by trained doctors or technicians.

6. Do not install, remove, or replace the Work Tip under heating state. Please power off before replace the Work Tip.

7. The Work Tip must be correctly installed to prevent it from falling off.

8. When the Work Tip is bent or worn, it will cause uneven heating. The operator should replace the Work Tip in time according to the clinical conditions;

9. After operation, please turn off the power immediately.

Woodpecker is specialized in producing medical instrument. We are only responsible for the safety on the following conditions:

a) The maintenance, repair, and modification are made by the manufacturer or the authorized dealers.

b) The charged components are original of "Woodpecker" and operated according to instruction manual.

9 Cleaning, Disinfection, Sterilization and Maintenance

The cleaning, disinfection and sterilization of Work Tip. Unless otherwise stated, it will be hereinafter referred to as "product".

/ Warnings

The use of strong detergent and disinfectant (alkaline pH>9 or acid pH <5) will reduce the life span of product. And in such cases, the manufacturer takes no responsibility. This product shall not be exposed to high temperature above 138° C.

9.1 Processing limit

The product have been designed for a large number of sterilization cycles. The materials used in manufacture were selected accordingly. However with every renewed preparation for use, thermal and chemical stresses will result in ageing of the product. The maximum number of sterilizations for Work Tip is 100 times. And each time you carry out Cleaning; disinfection and sterilization, you must make corresponding

records. And each time you carry out cleaning and disinfection, you must make corresponding records

9.2 Initial processing

9.2.1 Processing principles

It is only possible to carry out effective sterilization after the completion of effective cleaning and disinfection. Please ensure that, as part of your responsibility for the sterility of products during use, only sufficiently validated equipment and product-specific procedures are used for cleaning/disinfection and sterilization, and that the validated parameters are adhered to during every cycle. Please also observe the applicable legal requirements in your country as well as the hygiene regulations of the hospital or clinic, especially with regard to the additional requirements for the inactivation of prions.

9.2.2 Post-operative treatment

The post-operative treatment must be carried out immediately, no later than 30 minutes after the completion of the operation. The steps are as follows:

1. Turn off the Heating Handpiece and allow it to stand on the base for 1 minutes to cool down to room temperature at the Work Tip;

2. Use a cotton swab or a clean soft cloth to remove the remaining Gutta-percha material from the Work Tip,

3. Dry the product with a clean, soft cloth and place it in a clean tray.

9.2.3 Preparation before cleaning

Steps

Tools: tray, clean and dry soft cloth.

1. Remove the Work Tip Protector from the handle and put it into a clean tray.

2. Remove the Work Tip from the handle and place it in a clean tray.

3. Wipe the Work Tip with a soft cloth until no dirt can be seen on the surface. Then dry it with a clean soft cloth and put them into a clean tray. Cleaning agent can be pure water.

Notes:

The pure water temperature should not exceed 45 °C, otherwise the protein will solidify and it is difficult to remove.

9.3 Cleaning

The cleaning should be performed no later than 24 hours after the operation. The cleaning adopt automated cleaning.

The cleaning procedure are as follows.

1) Pre-wash with pure water at 25 ° C for 3 minutes.

2) Clean with the condition recommended by the cleaning agent manufacturer for 5 minutes. For example the detergent use RUHOF ENDOZIME AW PLUS WITH APA, Dilution Ratio1: 270, temperature 25°C. Clean for 5minutes.

3) Rinse twice with pure water at 25 $^{\circ}$ C for 1 minute each.

Notes:

a)The solution used the pure water and only freshly prepared solutions can be used.

b)During the use of cleaner, the concentration and time provided by manufacturer shall be obeyed.

c)The cleaner is proved to be valid by CE certificationin accordance with EN ISO 15883.

d)The cleaning procedure is suitable for the product, and the irrigating period is sufficient.

9.4 Disinfection

Disinfection must be performed no later than 2 hours after the cleaning phase. Automated disinfection is preferred if conditions permit.

For the thermal disinfection here, the temperature is 93 $^{\circ}$ C, the time is 5 min, and A0>3000.

Cleaning and disinfecting steps by using Washer-disinfector

1. Carefully place the product into the disinfection basket. Fixation of product is needed only when the product is removable in the device. The product is not allowed to contact each other.

2. Start the program.

3. After the program is finished, remove the product from the washerdisinfector, inspect (refer to section "Inspection and Maintenance") and packaging (refer to chapter "Packaging"). Dry the product repeatedly if necessary (refer to section"Drying").

The intrinsic suitability of the product for effective cleaning and disinfection using the above automated cleaning and disinfection procedures was verified by a certified facility.

Notes:

a) Before use the washer-disinfector, you must carefully read the operating instructions provided by the equipment manufacturer to familiarize yourself with the disinfection process and precautions.

b) With this equipment, cleaning, disinfection and drying will be carried out together.

c) Only pure water with a small amount of microorganisms (<10 cfu/ ml) can be used for all rinsing steps. (For example, pure water that is in accordance with the European Pharmacopoeia or the United States Pharmacopoeia).

d) The air used for drying must be filtered by HEPA.

e) Regularly repair and inspect the disinfector.

9.5 Drying

If your cleaning and disinfection process does not have an automatic drying function, dry it after cleaning and disinfection.

Methods

1) Spread a clean white paper (white cloth) on the flat table, point the product against the white paper (white cloth), and then dry the product with filtered dry compressed air (maximum pressure 3 bar). Until no liquid is sprayed onto the white paper (white cloth), the product drying is completed.

2) It can also be dried directly in a medical drying cabinet (or oven). The recommended drying temperature is $80^{\circ}C\sim120^{\circ}C$ and the time should be $15\sim40$ minutes.

Notes:

a) The drying of product must be performed in a clean place.

b) The drying temperature should not exceed 138 °C;

c) The equipment used should be inspected and maintained regularly.

9.6 Inspection and maintenance

In this chapter, we only check the appearance of the product. After inspection, ensure that there is no problem.

9.6.1 Check the product. If there is still visible stain on the product after cleaning/ disinfection, the entire cleaning/disinfection process must be repeated.

9.6.2 Check the product. If it is obviously damaged, smashed, detached, corroded, it must be scrapped and not allowed to continue to be used.

9.6.3 Check the product. If the accessory is found to be damaged, please replace it before use. And the new accessory for replacement must be cleaned, disinfected and dried.

9.6.4 If the number of times of the product reaches the specified number of times, please replace it in time.

9.7 Packaging

Install the disinfected and dried product and quickly package it in a medical sterilization bag (or special holder, sterile box).

Notes:

a) The package used conforms to ISO 11607;

b) It can withstand high temperature of 138 °C and has sufficient steam permeability;

c) The packaging environment and related tools must be cleaned regularly to ensure cleanliness and prevent the introduction of contaminants;

d) Avoid contact with parts of different metals when packaging. <u>9.8 Sterilization</u>

Use only the following steam sterilization procedures (fractional prevacuum procedure*) for sterilization, and other sterilization procedures are not recommended:

1. The steam sterilizer complies with EN13060 or is certified according to EN 285 to comply with EN ISO 17665;

2. The sterilization time is 5 minutes at a temperature of 134° C and a pressure of 2.0 bar ~ 2.3 bars.

Verification of the fundamental suitability of the products for effective steam sterilization was provided by a verified testing laboratory.

Notes:

a) Only the product that have been effectively cleaned and disinfected are allowed to be sterilized;

b) Before using the sterilizer for sterilization, read the Instruction Manual provided by the equipment manufacturer and follow the instructions.

c) Do not use hot air sterilization and radiation sterilization as this may result in damage to the product;

d) Please use the recommended sterilization procedures for sterilization. It is not recommended to sterilize with other sterilization procedures such as ethylene oxide, formaldehyde and low temperature plasma sterilization. The manufacturer assumes no responsibility for the procedures that have not been recommended. If you use the sterilization procedures that have not been recommended, please adhere to related effective standards and verify the suitability and effectiveness.

* Fractional pre-vacuum procedure = steam sterilization with repetitive pre--vacuum. The procedure used here is to perform steam sterilization

through three pre-vacuums.

9.9 Storage

9.9.1 Store in a clean, dry, ventilated, non-corrosive atmosphere with a relative humidity of 10% to 93%, an atmospheric pressure of 70KPa to 106KPa, and a temperature of $-20 \degree$ C to $+55 \degree$ C;

9.9.2 After sterilization, the product should be packaged in a medical sterilization bag or a clean sealing container, and stored in a special storage cabinet. The storage time should not exceed 7 days. If it is exceeded, it should be reprocessed before use.

Notes:

a) The storage environment should be clean and must be disinfected regularly;

b) Product storage must be batched and marked and recorded.

9.10 Transportation

1. Prevent excessive shock and vibration during transportation, and handle with care;

2. It should not be mixed with dangerous goods during transportation.

3. Avoid exposure to sun or rain or snow during transportation.

<u>9.11 The cleaning and disinfection of main unit and charging base are as</u> follows.

Warnings: Do not clean the main unit and charging base with ultrasound cleaning machine.

• Before each use, wipe the surface of the main unit and charging base with a soft cloth or paper towel soaked in 75% medical alcohol. Repeat the wipe for at least 3 times.

• After each use, wipe the surface of the main unit and charging base with a soft cloth soaked in clean water (pure water) or a clean disposable wipe. Repeat the wipe for at least 3 times.

9.12 Daily maintenance

1. When the device is not used, please turn off the power and unplug the power supply plug.

2. If the Heating and Packing Instrument is in a low battery state for a long time, the service life of battery will be shortened. Please charge it in time if the battery level is low.

3. When the device is not used, please charge it for 1 hour once a month.

9.13 Repair of device

This product does not contain self-repairing spare parts. If there is any abnormality in the equipment, please contact our company for maintenance and do not disassemble without authorization. With our company's consent, we will provide circuit diagrams, component part lists, descriptions, calibration instructions to assist to SERVICE PERSONNEL in parts repair.

Fault	Cause	Solution
No indications,	1. Inadequate battery power	1. Connect to power
no response	2. Battery is damaged.	supply to charge. /
	3. The charging interface is	Replace the battery.
	short-circuited, causing the	2. Replace the battery.
	lithium battery to enter a	3. Remove the substance
	protection state;	that causes the short
	4. Heating and Packing	circuit, put the device
	Instrument is damaged.	into the charging base
		to charge, and then the
		device will return to
		normal;
		4. Contact local distributor
		or manufacturer.
Automatic	If there is no operation for	Reboot
shutdown	10 minutes, the device will	
	automatic powers off.	
Work Tip	1. The Work Tip is damaged.	1. Replace the Work Tip
works	2. Malfunction of main unit	2. Send it to the repair
abnormally		center.
Charging	1. The power supply is not	1. Unplug and reconnect.
failure after	correctly connected;	2. Replace the battery.
connecting to	2. The power supply is	3. Wipe the thimble
power supply	damaged, or the specification	with alcohol, dry it, and
	doesn't match.	reconnect.
	3. There are impurities on the	
	contact thimble of charging	
	base.	

10 Troubleshooting

The service	The battery ages and the	Contact local distributor
time after each	battery capacity become	or manufacturer to
charging is	smaller.	buy new batteries for
shortened		replacement.
OPEN code	1. The Work Tip is damaged.	1. Replace the Work Tip.
appears on	2. The Work Tip is not	2. Install the Work Tip.
display screen	installed.	3. Unplug the Work Tip,
	3. The Work Tip is not well	and reconnect.
	installed.	

If the problem still cannot be solved, please contact your local dealer or our company.

11 After-sales service

Since the date of sales, if the device cannot work normally for quality problem, our company will be responsible for the repair of device during the warranty period. Please refer to the Warranty Card for warranty period and warranty scope.

12 Environment protection

The device does not contain any harmful ingredients. It can be handled or destroyed in accordance with the relevant local regulations.

Note:

1) Without Woodpecker agreement and authorization, private modification of device may result in the electromagnetic compatibility problem of that device or other devices.

2) The design and test of Heating and Packing Instrument complies with the related operation regulations of electromagnetic compatibility.

13 EMC-Declaration of comformity

The device has been tested and homologated in accordance with EN 60601-1-2 for EMC. This does not guarantee in any way that this device will not be effected by electromagnetic interference Avoid using the device in high electromagnetic environment.

Technical Description Concerning Electromagnetic Emission

Table 1: Declaration - electromagnetic emissions

Guidance and manufacturer's declaration - electromagnetic emissions

The model Fi-P is intended for use in the electromagnetic environment specified below. The customer or the user of the model Fi-P should assure that it is used in such an environment.

Emissions test	Compliance	Electromagnetic environment - guidance
RF emissions CISPR 11	Group 1	The model Fi-P 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 CISPR11	Class B	The model Fi-P is suitable for used in all establishments,
Harmonic emissions IEC 61000-3-2	Class A	including domestic establishments and those
Voltage fluctuations / flicker emissions IEC 61000-3-3	Complies	directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.

Technical Description Concerning Electromagnetic Immunity

Table 2: Guidance & Declaration - electromagnetic immunity

Guida	ince & Declarati	on — electromagn	etic immunity
The model Fi-P is intended for use in the electromagnetic environment specified below. The customer or the user of the model Fi-P should assure that It is used in such an environment.			
Immunity test	IEC 60601 test level	Compliance level	Electromagnetic environment - guidance
Electrostatic discharge (ESD) IEC 61000-4-2	±8kV contact ±2, ±4, ±8, ±15kV air	±8kV contact ±2, ±4, ±8, ±15kV 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	±2kV for power supply lines ±1kV for Input/ output lines	±2kV for power supply lines	Mains power quality should be that of a typical commercial or hospital environment.

Surge	$\pm 0.5, \pm 1$ kV line	$\pm 0.5, \pm 1$ kV line to	Mains power quality	
IEC 61000-4-5	to line	line	should be that of a	
	$\pm 0.5, \pm 1, \pm 2kV$	$\pm 0.5, \pm 1, \pm 2 kV$	typical commercial or	
	line to earth	line to earth	hospital environment.	
Voltage	<5 % UT	<5 % UT	Mains power quality	
dips, short	(>95% dip in	(>95% dip in UT.)	should be that of a	
interruptions	UT.)	for 0.5 cycle	typical commercial or	
and voltage	for 0.5 cycle	<5 % UT	hospital environment. If	
variations on	<5 % UT	(>95% dip in UT.)	the user of the models	
power supply	(>95% dip in	for 1 cycle	Fi-P requires continued	
input lines	UT.)	70% UT	operation during power	
IEC 61000-4-11	for 1 cycle	(30% dip in UT)	mains interruptions, it is	
	70% UT	for 25 cycles	recommended that the	
	(30% dip in UT)	<5% UT	models Fi-P be powered	
	for 25 cycles	(>95 % dip in UT)	from an uninterruptible	
	<5% UT	for 250 cycles	power supply or a	
	(>95 % dip in		battery.	
	UT)			
	for 250 cycles			
Power frequency	30A/m	30A/m	Power frequency	
(50/60 Hz)			magnetic fields should	
magnetic field			be at levels characteristic	
IEC 61000-4-8			of a typical location in	
			a typical commercial or	
			hospital environment.	
NOTE UT is the a.c. mains voltage prior to application of the test level.				

Table 3: Guidance & Declaration - electromagnetic immunity concerning
Conducted RF & Radiated RF

Guidance & Declaration - Electromagnetic immunity			
The model Fi-P is intended for use in the electromagnetic environment specified below. The customer or the user of the models Fi-P should assure that it is used			
in such an environment.			
Immunity test		Compliance	Electromagnetic environment -
	test level	level	guidance

Conducted RF	3 Vrms	3V	Portable and mobile RF
IEC 61000-4-6	150 kHz to 80	6V	communications equipment
Conducted RF	MHz	3V/m	should be used no closer to any
IEC 61000-4-6	6 Vrms		part of the models Fi-P, including
Radiated RF	ISM		cables, than the recommended
IEC 61000-4-3	frequency		separation distance calculated
	band		from the equation applicable to the
	3 V/m		frequency of the transmitter.
	80 MHz to 2.7		Recommended separation distance
	GHz		$d=1.2 \times P^{1/2}$
			$d=2 \times P^{1/2}$
			$d=1.2 \times P_{1/2}^{1/2}$ 80 MHz to 800 MHz
			d=2.3×P ^{1/2} 800 MHz to 2.7 GHz
			where P is the maximum output
			power rating of the transmitter
			In watts (W) according to the
			transmitter manufacturer and d
			Is the recommended separation
			distance in meters (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:
NOTE I At 80 N	MHz end 800 M	Hz. the high	er frequency range applies.
NOTE 2 These	midalinas may	not apply in	all situations. Electromagnetic

NOTE 1 At 80 MHZ end 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 model Fi-P is used exceeds the applicable RF compliance level above, the model Fi-P should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as reorienting or relocating the model Fi-P. b Over the frequency range 150 kHz to 80 MHz, field strengths should be less

than 3V/m.

 Table 4: Recommended separation distances between portable and mobile

 RF communications equipment and the model Fi-P

	ecommended separa mobile RF commun Fi		
	ntended for use in eleances is controlled.	-	
	ent electromagnetic ir		
(transmitters) and t	ortable and mobile R he model Fi-P as reco ower of the commun	ommended below, ac	
Rated maximum	Separation distance according to frequency of transmitter		
output power	m		
of transmitter		80MHz to 800MHz	800MHz to
W	$d=1.2 \times P^{1/2}$	$d=1.2 \times P^{1/2}$	2,7GHz
			$d=2.3 \times P^{1/2}$
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 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 in watts (W) accordable to the transmitter manufacturer.

NOTE I 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.

14 Statement

Woodpecker reserves the right to change the design of the equipment, the technique, fittings, instruction manual and the content of the original packing list at any time without further notice. The pictures are only for reference. The final interpretation rights belong to Guilin Woodpecker Medical Instrument Co., Ltd.

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EC.

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Name of	Warranty Card	
Customer		-
Address Details		
Postal Code		
Tel		(1)
Model		For Distribute
Product No.		
Purchase Date		
Contact Person		
Date	Maintenance Record	Repairer
Inform Zone, Sales I After-	Woodpecker Medical Instrumen hation Industrial Park, Guilin National Guilin, Guanysi, 541004 P. R. China Dept.: +86-773-5873196/2350599 sales Service Dept.: +86-0773-5827i I: woodpecker4@glwoodpecker.com te: http://www.glwoodpecker.com	High-Te

Gutta percha Obturation Device Warranty Card

Name of Customer		
Address Details		
Postal Code		
Tel		(II)
Model		Return to Manufacturer
Product No.		
Purchase Date		
Contact Person		
Date	Maintenance Record	Repairer

Guilin Woodpecker Medical Instrument Co.,Ltd. Information Industrial Park, Guilin National High-Tech Zone, Guilin, Guangxi, 541004 P. R. China Sales Dept.: +86-773-5873196/2350599 After-sales Service Dept.: +86-0773-5827898 E-mail: woodpecker4@glwoodpecker.com Website: http://www.glwoodpecker.com

Distributor:

1

1

Seal

Warranty Instruction

I Period validity:

Since the date of sales, our company will be responsible for the repair of main unit, charging stand, and power adapter for one year. Please refer to the Warranty Card for warranty period and warranty scope. The warranty period of lithium battery is one year.

II Range of warranty:

Within the warranty period of validity, we are responsible for any troubles caused by quality problems or products technique and structure.

III The following are beyond our warranty:1. The damage caused by disobeying the operation instruction or lack of the needed condition.

2. The damage caused by unsuitable operation or disassembly without authorization.

3. The damage caused by unadvisable transportation or preservation.

4. There isn't the seal of distributor or the warranty card isn't filled in completed.

5. Except for the main unit, charging stand, power adapter and lithium battery, the other accessories are not covered by the warranty.

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I Period validity:

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 The damage caused by unsuitable operation or disassembly without authorization.
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5. Except for the main unit, charging stand, power adapter and lithium battery, the other accessories are not covered by the warranty.