文件名/NAME	根管长度测量仪英文说明书Woodpex X CE无 Logo 1.3 版 2406111 14.02	2.04.033	代码/code	14.02.0	04.033
尺寸/SIZE	150×140mm, 出血 6mm		版本/REV.	V1.3	
材质/MATERIAL	120g 铜版纸				
工艺/PROCESS	/				
装订&注释/ bind (books etc) &NOTES	骑马订; 保修卡正反面印刷需对应。	印刷 ■C	l颜色/COL MYK	ORS	4X4
修订日期/DATE	2024.06.11				

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

Apex	Locator Warranty Card		
g the dath	(Re	eturn to Manufacturer)	Warranty Instruction
Tel	(,	1. Period validity:
Address Details		IIIn Woodpecker Medical Instrument Co., Ltd. crimities Extratrul Park, Oxfer National High-Tech- re, Oxfer, Comput. 541804 P. R. China In: Data: 1-69-723-5823154/2005099	We offer two years free repair on main unit and six months on spare parts from the date of
Postal Code	Abo 5-m Web	ze-szles Senice Dagz.: +84-1779-5827890 na Lwoodpecken/d0glwoodpeckercom Isster http://www.glwoodpeckercom	purchase, warranty card must be attached.
Model	Distrib	butor:	2. Range of warranty:
Product No.			Within the warranty period of validity, we are responsible for any troubles caused by quality
Purchase Date			problems or products technique and structure.
Contact Person		Seal	3. The following are beyond our warranty:
Date	Maintenance Record/Repairer	r	a. The damage caused by disobeying the operation instruction or lack of the needed condition.
			 c. The damage caused by unstituate operation or usassembly without automization.
			d. There isn't the seal of distributor or the warranty card isn't filled in completed.
			e. Device failures caused by the use of non-original accessories are not under warranty.

* Cut alo		Apex Locator Warranty Ca	ırd	
ng the dashed	Name of Customer		(For Distributor)	Warranty Instruction
Ine	Tel			1. Period validity:
	Address Details		Adult Weschecker Hedical Instrument Co., Ltd. Hormston Houzini Prot., Oal In Netional High-Tech. Zone, Coder, Danapat, Sk1004 P. R. China Sales Dept. 198-773 SISTING235999 Alter Service Dept. 198-0703 SISTING Encode methodisation of the constants arease	We offer two years free repair on main unit and six months on spare parts from the date of
	Postal Code		Website: http://www.gl.woodpediectors	purchase, warranty care most be attached.
	Model	D	Distributor:	2. Range of warranty:
	Product No.			Within the warranty period of validity, we are responsible for any troubles caused by quality problems or products technique and structure.
	Purchase Date			
	Contact Person		Seal	3. The following are beyond our warranty:
	Date	Maintenance Record/Repa	airer	 a. The damage caused by disobeying the operation instruction of tack of the needed conduction. b. The damage caused by unsuitable operation or disassembly without authorization.
				c. The damage caused by unadvisable transportation or preservation.
				d. There isn't the seal of distributor or the warranty card isn't filled in completed.
				e. Device failures caused by the use of non-original accessories are not under warranty.
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Apex locator Instruction Manual

Please read this manual before operating ZMN-SM-828 V1.3-20240611



Guilin Woodpecker Medical Instrument Co., Ltd.

Catalog

1 Introduction	1
2 Notice of installing and using the device	4
3 Installation of the device	6
4 Product function and operation	13
5 Trouble shooting	18
6 Cleaning, Disinfection and Sterilization	20
7 Storage, maintenance and transportation	23
8 Environmental protection	
9 European authorized representative	
10 After service	
11 Symbol instruction	
12 Statement	
13 EMC - Declaration of conformity	

1 Introduction

1.1 Foreword

Guilin Woodpecker Medical Instrument Co., Ltd. is a professional manufacturer in researching, developing and producing dental equipment which has a wholesome quality assurance system.

Products include ultrasonic scaler, curing light, apex locator and ultrasurgery, etc.

1.2 Description of the device

Apex locator is a supporting equipment of endodontic treatment, through the measurement of the length of apical teeth, helping dentists to finish the endodontic treatment.

Features of the device:

a) Equipped with clear bright LCD, clear image and different colors indicate the trajectory of the file clearly.

b) Based on advanced multiple frequency network impedance measurement technology and automatic calibrating ensures the measurements are accurate.

c) The File clip, Lip hook, Touch probe and Endodontic explorer(Optional) can be autoclaved under high temperature and high pressure. Avoiding cross infection effectively.

d) Battery is rechargeable, unnecessary to replace batteries repeatedly.

e) Contact materials with the human body;

File clip: plastic material;

Lip hook: stainless steel;

Touch probe: rubber, stainless steel;

Endodontic explorer(Optional):stainless steel;

1.3 Model and dimensions

1.3.1 Dimensions: 74mm×67mm×123mm

1.3.2 Weight: 240g±10g

1.3.3 Model: Woodpex X

1.4 Components

1.4.1 Picture of the main unit. (Picture 1)







1.5 Structure

Is composed of main unit, measuring wire, lip hooks, file clip, touch probe, Adapter(Optional), Charging cable(Optional) and Endodontic explorer(Optional), etc.

1.6 Intended use

This equipment applies to the measurements below:

1.6.1 Used to help determine the working length of various types of dental root canals during root canal treatment.

1.6.2 Used to test pulp vitality.

1.7 Contraindication

We do not advise the use of the model on patients fitted with pacemakers (or other electrical equipment) or on those patients who are advised not to use the electric equipment (like electric shaver, electric blower) for safety reasons.

1.8 The classification of the device

1.8.1 Type of protection against electric shock: Class II equipment

1.8.2 Degree of protection against electric shock: Type BF applied part

1.8.3 Degree of protection against water shock: Ordinary equipment (IPX0)

1.8.4 Device not suitable for being used in the presence of a flammable anesthetic mixture with air or with oxygen or nitrous oxide.

1.8.5 Operation mode: Continous operation

1.8.6 Applied part: Touch probe, Lip hook, File clip and Endodontic explorer(Optional)

1.9 The main technical specifications

1.9.1 Battery: 3.7V/2000mAh (Model: 18500)

1.9.2 Adapter (ADS-6AM-06N 05050)(Optional):

Input: ~100V-240V 50Hz/60Hz 0.4A MAX

Output: DC5V/1A

1.9.3 Consumption power: ≤0.5W

1.9.4 Screen: 3.8" LCD

1.9.5 Buzzer alert: The buzzer will alert when the endo file is close to the apex.

1.9.6 Release software version: V1

1.9.7 Operation condition

a) Environment temperature: +5°C~+40°C

b) Relative humidity: 30% ~75%.

c) Atmosphere pressure: 70kPa~106kPa

2 Notice of installing and using the device

2.1 Please read the instruction manual carefully before the operation.

2.2 When the indicating bar reaches the position of the dial 0.0 and there is "APEX" on screen, the endo file has reached the anatomical apical foramen. To guarantee the safety, the work length is clinically obtained by subtracting 0.5-1mm from the length measured by the Apex locator.

2.3 The scales 0.5 and 1.0 on the screen dial do not indicate that the distance to the apex is 0.5 mm or 1.0 mm. It just reminds the operator that the file is getting close to or away from the apical foramen.

2.4 If the screen bar graph suddenly makes a large movement or immediate display 'OVER' in the upper part of the canal, continue slightly towards the apex so the signal returns to normal.

2.5 In order to prevent leakage or interference between the root canal and resulting in inaccurate measurements, dry the access cavity with a cotton pellet or air-blower before each use.

2.6 Use a file size adapted to the root canal diameter. The selected file is too small for a large root canal might cause the screen digital display is not steady during the procedure.

2.7 In order to confirm the file clip and measuring wire makes good contact, test the wire connecting before each use(See 3.1.2).

2.8 The file clip, lip hook and touch probe and Endodontic explorer(Optional) are reusable. Please make sure they are autoclaved under high pressure and high temperature before each operation. The endo files should not be used more than 3 times.

2.9 The batteries must be taken out for storage when the device is not used for a long time. 2.10 Please recharge the battery when low battery indicator flashes.

2.11 Please use original components, the components made by other companies may cause inaccurate measurement or un-measurable.

2.12 Avoid the connection between the outside and inside liquid of endodontic during measuring in order to avoid the measuring difference.

2.13 Keep endo file and file clip away from any other metal or instruments.

2.14 To ensure that short circuits do not impair the measurements, be particularly careful with patients fitted with metal crowns or bridges. Please confirm the wetness of the endo to ensure the reliability of the measuring. If it is confirmed that the endo file hasn't reached the apex yet the data showed on the apex locator is too low, please check whether the endo is too dry and confirm it with X-ray.

2.15 This device have electromagnetic interference, the patient or doctor who with a heart pace maker are forbidden to use this device and the device is susceptible to other device which produces electromagnetic interference. Dentists should be cautious about operation under such environment.

2.16 The guarantee is valid for normal usage conditions. Any disassembly will render the guarantee void, the professionals of Woodpecker company will offer the repair service during guarantee period.

2.17 Any modification will render the guarantee void and may cause harm to the patient.

2.18 Only the original adapter or the original charging cable with the charging plug that meets the sparameter requirements(see 2.23) and the original lithium battery could be used to this machine.

2.19 Not to position equipment to make it diffcult to operate the disconnection device.

2.20 The adapter must be connected to an appropriate power source in the instructions.

2.21 Error in replacing lithium batteries can lead to unacceptable risks, so use the original lithium battery and replace the lithium battery according to the correct steps in the instructions.

2.22 Please to remove the battery if the me equipment is not likely to be used for some time.

2.23 If the charging method of charging cable with self-purchased charging plug is adopted, the charging plug needs to comply with CE certification, and the parameters meet: Input: 100-240VAC 50/60Hz 0.4A Max

Output: 5V 1A-2A

2.24 The device must be operated in hospital and clinic by the qualified dentists.

2.25 It is recommended that users employ a DG16 instrument with a narrow, sharp, and stiff tip for optimal performance. To minimize errors, it is suggested to use the manufacturer-provided DG16.

3 Installation of the device

3.1 Apex Locator Mode

3.1.1 Preparation

Insert the plug of the measuring wire into the right side socket of the unit.

Attention:

a) Please be careful to use the device, keep it stable and avoid hit. Incautious use will lead to the damage or the failure of the machine.

b) Measurement can not be proceeded without the complete insertion of the plug.

c) Be sure not to hit the plug. Keep the device away.

Insert the file clip and lip hook respectively into the two sockets of the measuring wire. When the Apex Locator is used alone, there is no difference between the gray end and the white end of measuring wire [Picture 3]. But if the Apex Locator is connected to the Endo Motor, please connect the white end with the lip hook and the gray end is suspended.



Attention:

Be sure not to pull the wire when inserting or pulling out the the measuring wire and the file clip. [Picture 4 (a)]

Correct operation showed as in picture 4 (b).

3.1.2 Test the wire connecting (Test before each use)

a) Press the power switch. Make sure the scene of measuring the length of the root canal displayed on the LCD screen.

b) Make sure if the plug of the measuring wire is inserted into the socket correctly.



Picture 5



Picture 6

c) Make sure if the file clip and lip hook are connected well to the measuring wire.

d) Make the lip hook touch the bent wire of the file clip [as showed in Picture 5] make

sure the connection icon on the LCD screen shows steadily [as showed in Picture 6], otherwise, it means that the file clip or the measuring wire is damaged, should be replaced.

3.1.3 Determine the working length

a) When the indicating bar reaches the position of the dial 0.0[Picture 7(a)], [Picture 7(b)] and there is "APEX" on screen, the endo file has reached the anatomical apical foramen. On the basis of measured length, subtract 0.5-1.0 mm to get the working length.

b) When the indicating bar reaches the red area "OVER" [Picture 7(c)], it indicates that the endo file has exceeded the apical foramen.



Picture 7

* The working length will differ somewhat depending on each individual tooth. This discrepancy must be judged by the dentist as he/she works on the tooth.

* Make sure to take an X-ray to check the results.

3.1.4 Apical Stop Setting

Set the Apical Stop between 0.0 and 0.5 by pressing the middle button, and the set parameter will be automatically saved. When the file reaches the Apical Stop, the device will beep continually.





3.1.5 Testing the device by tester (Test every two weeks)

Users can use the tester to check if the device work properly, specific operation is as follows:

a) Pulling out the the measuring wire and turn off the device.



Picture 9

b) Insert the tester.

c) After powered on, If the indicating bar indicates within ± 1 bars away from the dial 0.0 the device functions normally [Picture 9]. If the indicating bar is outside the range, the device cannot measure accurately. On this occasion, please contact authorized distributor or manufacturer for help.

3.1.6 Connect to compatible Endo Motor.

Plug one end of the USB line into the USB socket on the right side of the device, and connect the other end with compatible Endo Motor as shown in Picture 10 (a). There is no difference between those two ends. As shown in Picture 10 (b), when the Contra-angle icon is lit, the Apex locator and Endo Motor can communicate normally, so that the 2-in-1 function can be realized in Endo Motor.



[Cautions] :

① Please use the Apex locator carefully and do not drop it or hit it. Careless use may bring risk of damage to the machine or malfunction.

2 If the USB wire was not completely plugged into the USB socket, the Apex locator

cannot communicate with the Endo Motor.

③ After plugging the USB wire into the USB socket, please do not drop anything on it, and do not hit the USB socket.

3.2 The Micro Hole Negotiator (MHN) Mode (Match only device with endodontic explorer) The MHN mode is designed to aid clinicians in three key areas during root canal treatment:

A) Locating the pulp chamber when preparing an access cavity with the aim of reducing the removal of excessive amounts of dentin and preventing damage to the floor of the pulp chamber.

B) Locating highly calcified root canal orifices that are difficult to visualize and challenging to detect using conventional methods.

C) Evaluating the leakage of non-electro conductive root canal fillings (e.g. gutta-percha) to assist in the decision to remove the root filling and retreat the canal system.

Connect the measuring line, file/probe clip, and lip hook to the interfaces on the device. Turn on the device and select the MHN mode by holding the mode button for more than 2 seconds, as shown in Picture 11.



Picture 11

3.2.1 Locating the Pulp Chamber

Method of operation:

1. Start to prepare an access cavity through enamel and dentin using an appropriate bur.

2. Isolate the tooth and dry the access cavity to ensure accurate readings.

3. Attach the lip hook to the patient and the probe clip to a endodontic explorer DG 16, etc.).

4. Place the tip of the explorer on the dentin at the estimated center of the pulp chamber and monitor the readings on the screen of the device.

a) Reading 0.00 - 1.00: No connection; consider more dentin removal with a bur and/or reevaluate depth/alignment of the access cavity using radiographs/CBCT images.

b) Reading 2.00 - 3.00: Connection established; remove dentin using an ultrasonically activated tip without damaging the pulp chamber floor.

c) Reading above 3.00: Pulp chamber is detected. The access cavity at the point of pulp chamber exposure can now be enlarged/widened carefully with ultrasonic tips to remove the dentine of the pulp chamber roof in order to allow the floor of the pulp chamber to be explored.

3.2.2 Locating Root Canal Orifices

Method of operation:

1. Prepare an access cavity using good illumination and magnification.

2. Isolate the teeth, then debride and clean the pulp chamber with NaOCl and an ultrasonic device.

3. Attach the lip hook to the patient and the probe clip to the endodontic explorer or an endodontic file or spreader.

4. Dry the pulp chamber floor to ensure accurate readings.

5. Place the tip of the explorer (or file/spreader) on the dentin at the estimated position of the orifice and monitor the readings of the screen of the device whilst moving the explorer around the region.

a) Reading 0.00 to 1.00: indicates no connections. The operator should consider removing

dentin with an ultrasonic device or carefully examine other areas where the orifice may lie. b) Reading 2.00 to 3.00: indicates a connection and the potential location of the orifice. The dentin at the location should be removed ultrasonically to reveal the canal orifice.

c) Reading above 3.00: indicates the location of the orifice. A suitable hand file can now be used to enter and negotiate the root canal.

Note 1: The values on the device set out above are for reference only; they are not absolute values. The readings will be affected by moisture within the pulp chamber, the shape of the pulp chamber and the diameter of the root canal.

Note 2: The MHN mode is an auxiliary function and should only be used as an adjunct to the clinician's clinical judgment.

4 Product function and operation

4.1 Usage requirements

Apex locator should be precise, repeatable, and easy to operate. The following requirements are necessary besides the proper operation method.

4.1.1 The operation should be according to the manual.

4.1.2 The dentists should have the knowledge of teeth position and average length and the skill to operate the device.

4.1.3 An fully exposed access cavity to show the pulpal cabin.

4.1.4 A X-ray photo to show the whole length and root canal of the teeth.

4.1.5 The endo file should not be too big nor too small to avoid cutting through the apical foramen.

4.1.6 Mark an anatomized symbol on the diseased tooth and memorize it on the case history. This symbol should be marked on the health bridge or on the tooth filled integrated. The position of the mark should be on the incisal edge of the anterior tooth or on the spire of the molars. For those bridge that's broken obviously, this symbol should be on the tooth surface supported by the dentin instead of on the suspended enamel.

4.1.7 The acute inflammation surrounding the apex has been gone and the infected materi-

al has been cleaned. It is also necessary to get rid of the pulp and necrosis tissue.

4.1.8 The following cases are not suited for a normal measurement:

a) The size of the root similar to the size of apical foramen.

In this case, the measurement result of the length of the root canal will be shorter than its real because of the hypoplasia of the root [Picture 12].

b) Bleeding or the blood overflow from the apical foramen.

In this case, the blood will overflow from the root canal and reaches gingival that the blood and the gingival will be on a conducting state which will cause an inaccurate result while measuring. The measurement can be continued when the bleeding is stopped [Picture 13].

c) The tooth crown is broken.

The tissue of the gingival may reach the cavity of the endo hole at the broken point which will cause inaccuracy because of the electronic conduction. The measurement can be continued when the crown is fixed by gypsum or other insulators [Picture 14]







Picture 13



Picture 14

d) There is a crack on the tooth root.

In this case, the crack may cause the electric leakage which will affect the accuracy of measurement [Picture 15].

e) A retreatment to an endo which was filled with gutta-percha.

Clean the remaining material in the root canal and fill it with little normal saline before a measurement [Picture 16].

f) There is a metal crown which has connected to the gingival.

It will cause an inaccuracy when the endo file touches metal crown [Picture 17].



Sometimes, the results of the Apex Locator and X-rays do not meet each other, which is neither bdcause the machine is not normal, nor the photo is incorrect taken. The actual position of the apical foramen is different from the anatomical one, it is very common that the apical foramen slightly to the side of the root canal crowns. In this case, according to the shooting angle as the belowing picture show, it will cause illusion that the front tip of the root canal haven't reached the canal tip. [Picture 18]

(Because of the angle of X-rays, sometimes it can't take photo of the apical foramen properly, so it can't show the accurate position of the apical foramen.)

4.2 Instruction

4.2.1 Insert the plug of measuring wire into the socket in the side of main unit. Turn it on. The battery is on the left of screen.

4.2.2 The volume is adjustable. Please press the volume bottom for a setting.

4.2.3 Hang the lip hook on the lip, make sure it contact the oral mucosa as a reference electrode [Picture 19].

4.2.4 Clip the file with file clip, approach to the apex, then there will be continuous alarm when the distance is less than 2mm [Picture 20].



Attention:

a) When gripping the root canal with a canal file, please grip the upper of the metal part (near the root canal at the needle handle). If you grip the lower part (blade or moving part), it will wear the file clip. [Picture 21]

b) When measuring the length of root canal, please use the canal file with the resinous handle.

If you operate the device without the dentistry glove, it will cause leakage and the result of measurement will be inaccurate. Therefore, please use the resin needle file and remember don't touch the metal part with finger.

c) Please don't use the worn file clip, and it will make the result of measurement inaccurate.

d) Please reference the [Picture 22 (a)] to grip the needle file. If as [Picture 22 (b)], it can't properly measure the length of the root canal due to the improper force, and the front of the root canal pin is easy to wear.



4.2.5 When the file reaches the apex, adjust the rubber piece set on the endo file to the reference point (incisal edge or fossa edge), then pull out the endo file, measure the length between the top of the file and the rubber piece, and this is the working length of the tooth. It also can be use with the touch probe instead of file clip, when it is inconvenient to measure the back teeth [Picture 23].





4.2.6 Please remove the lip hook, file clip or touch probe after shutting down.

4.2.7 The components that touch body must be autoclaved under high temperature and high pressure. The shell and measuring wire should be cleaned by 75% alcohol.

Attention: Avoid the silk-screen when cleaning.

4.2.8 Adjust the Sleep Duration.

a. The device with apex locator only.without performing

root canal measurements, adjust the volume to the maximum. Long-press the mode button

until you hear a "beep".

b. The device with Micro Hole Negotiator (MHN) mode, long-press the adjustment button until you hear a "beep".

It indicates a successful adjustment of the sleep duration. There are three sleep durations adjustable: 10 min, 20 min, and 30 min. The default setting sleep times is 20 min

4.2.9 Adjust the sound level. In the Apex Locator Mode, adjust the volume to the maximum, long-press the adjustment button until you hear a "beep", indicating a successful adjustment of the sound level. There are 2 sound levels adjustable.

4.2.10 The device will sleep automatically after 3 minutes without operation. After the sleep times, the device will shutdown automatically.

Problems	Possible cause	Solutions
No power and no signal on the screen after the power on.	 If the battery is placed correctly? If the battery with no power? 	 Re-install the battery. Recharge the battery.
The length of the root canal cannot be measured.	 If the measuring wire is connect- ed correctly? If the measuring wire is broken? 	Confirm the measuring wire is plugged firmly, link the lip hook with the file clip to check if the measuring wire is broken.
No sound of alarm.	If the volume is set at "mute"?	Adjust the sound level.
The charging LED indicator goes out.	 The adapter is not connected well. Have used faulty adapter with excessive output. The battery is not installed well. The battery has been damaged. 	 Reconnect the adapter. Change the adapter, must use the original adapter or the charging plug that meets the sparameter requirements(see 3.3.5) Reinsert the battery and then reconnect the adapter. Change the battery and then reconnect the adapter.

5 Trouble shooting

Problems	Possible cause	Solutions
	If the connection between the lip	Make sure the lip hook has contacted the oral
	hook and the oral mucosa is ok?	mucosa at a good position.
	Is there a blood/saliva overflowing, glued to the crown?	Blood, liquid overflow from the root canal, glued to the crown or the tooth neck, will cause short-circuit then cause the in-normal phenomena. Clean the blood and the liquid.
Display not steady while measuring: the measurement result is rather longer or shorter; numerical display irregular.	If the root canal is filled with blood, liquid?	Once the endo needle contact the surface of the root canal which is filled with blood, liquid, it will display "OVER" immediately. In this case, push the needle to the apical root canal, then the display will be normal, you can measure the length of the root canal correctly.
	If there is liquid, scrap on the tooth surface?	Clean the tooth surface.
	If the endo needle contact the gums?	The LCD will display "OVER" if the endo needle contact the gums.
	If there is still pulp in the root canal?	If there is much pulp left in the root canal, the root canal length can't be measured correctly.
Display not stor dy while	If the needle touched the metal repaired material?	Once the needle touched the metal repaired material, current measurement from the gums to the periodontal tissue loss, the screen will display "OVER".
measuring: the measurement result is rather longer or shorter: numerical diplay	If the adjacent surface has caries?	Current measurement flow from caries of the adjacent surface to gums, then the root canal length can't be measured correctly.
irregular.	Whether there is collateral or the tooth root is broken?	Once the needle reached the collateral or the broken part of the tooth root, current measure- ment will overflow from periodontal ligament, it displays "OVER".
	Is it because in addition to the top pulp chamber, low tooth crown? Or there are residues left?	Use rubber dam to prevent the current flow to gums.

Problems	Possible cause	Solutions
Display not steady while	Are there cysts apical?	If there has cysts, the length of root canal can't be measured accurately.
result is rather longer or	Whether the file clip is not clean or broken?	Clean the file clip by alcohol, or replace it.
irregular.	Whether the measuring wire is broken or poor contact?	Contact the both end of the measuring wire directly, it displays "-3".
	Whether the root canal is occlu- sive?	The display will be normal after penetrating the narrow part of apical.
The length measurement indi- cator only full display near	If the root canal is too dry?	Wet the root canal with normal saline solution or sodium hypochlorite solution.
narrow part of the apical.	If the endo file is too small for a large root canal?	Replace the current endo file with a larger one.

* If all above measures do not work, please contact us.

6 Cleaning, Disinfection and Sterilization

The procedure for cleaning, disinfection and sterilization applies only to the accessories file clip, touch probe and lip hook.

Reprocessing procedures have only limited implications to this file clip, touch probe and lip hook. The limitation of the numbers of reprocessing procedures is therefore determined by the function/ wear of the device. However with every renewed preparation for use, thermal and chemical stresses will result in ageing of the products. The specified maximum times of sterilization for File clip is 500 times. The specified maximum times of sterilization for Touch probe and Lip hook is 1000 times.

In case of damage the device should be reprocessed before sending back to the manufacturer for repair.

6.1 Preparation at the Point of Use:

Disconnect the file clip, touch probe and lip hook from the measuring wire. Remove gross soiling of he instrument with cold water ($<40^{\circ}$ C) immediately after use. Don't use a fixating detergent or hot water ($>40^{\circ}$ C) as this can cause the fixation of residuals which

may influence the result of the reprocessing process. Store the instruments in a humid surrounding.

6.2 Transportation:

Safe storage and transportation to reprocessing area to avoid any damage and contamination to environment.

6.3 Preparation for Decontamination:

The devices must be reprocessed in a disassembled state.

6.4 Pre-Cleaning:

Do a manual pre-cleaning, until the instruments are visually clean. Submerge the instruments in a cleaning solution and flush the lumens with a water jet pistol with cold tap water for at least 10 seconds. Clean the surfaces with a soft bristol brush.

6.5 Cleaning:

Regarding cleaning/ disinfection, rinsing and drying, it is to distinguish between manual and automated reprocessing methods. Preference is to be given to automated reprocessing methods, especially due to the better standardizing potential and industrial safety. Automated Cleaning:

Use a washer-disinfector meeting the requirements of the ISO 15883 series. Put the instrument into the machine on a tray. Connect the instrument with the WD by using suitable adapter and start the program:

• 4 min pre-washing with cold water (<40°C);

• emptying

• 5 min washing with a mild alkaline cleaner at 55°C

• emptying

• 3 min neutralising with warm water (>40°C);

• emptying

• 3 min neutralising with warm water (>40°C);

• emptying

• 5 min intermediate rinsing with warm water (>40°C)

• Emptying

The automated cleaning processes have been validated by using 0.5% neodisher Medi-Clean forte (Dr. Weigert). Note Acc. to EN ISO 17664 no manual reprocessing methods are required for these devices. If a manual reprocessing method has to be used, please validate it prior to use.

6.6 Disinfection:

Automated Thermal Disinfection in washer/disinfector under consideration of national requirements in regards to A0 value (see EN 15883). A disinfection cycle of 5 min disinfection a 93°C has been validated for the device to achieve an A0 value of 3000.

6.7 Automated Drying:

Drying of outside of instrument through drying cycle of washer/ disinfector. If needed, additional manual drying can be performed through lint free towel. Insufflate cavities of instruments by using sterile compressed air.

6.8 Functional Testing, Maintenance:

Visual inspection for cleanliness of the instruments and reassembling. Functional testing according to instructions of use. If necessary, perform reprocessing process again until instruments is visibly clean. Before packaging and autoclaving, make sure that the file clip, touch probe and lip hook has been maintained acc. to manufacturer's instruction.

6.9 Packaging:

Pack the instruments in an appropriate packaging material for sterilization. The packaging material and system refer to EN ISO 1167.

6.10 Sterilization:

Sterilization of instruments by applying a fractionated pre-vacuum steam sterilization process (according to EN 285/EN 13060/EN ISO17665) under consideration of the respective country requirements. Minimum requirements: 3 min at 134°C (in EU: 5 min at 134°C) Maximum sterilization temperature: 137°C.

Note: Flash sterilization is not allowed on lumen instruments!

6.11 Storage:

Storage of sterilized instruments in a dry, clean and dust free environment at modest temperatures, refer to label and instructions for use.

7 Storage, maintenance and transportation

7.1 Storage

7.1.1 This equipment should be stored in a room where the relative humidity is $10\% \sim 93\%$, atmospheric pressure is 70kPa to106kPa, and the temperature is $-20^{\circ}C \sim +55^{\circ}C$.

7.1.2 Avoid the storage in a too hot condition. High temperature will shorten the life of electronic components, damage battery, reshape or melt some plastic.

7.1.3 Avoid the storage in a too cold condition. Otherwise, when the temperature of the equipment increases to a normal level, there will be dew that will possibly damage PCB board.

7.2 Maintenance

7.2.1 This device do not include accessories for repair usage, the repair should be carried out by authorized person or authorized after service center.

7.2.2 Keep the equipment in a dry storage condition.

7.2.3 Do not throw, beat or shock the equipment.

7.2.4 Do not smear the equipment with pigments.

7.2.5 Replace the battery if it seems to be running out of power sooner than it should. Please use the original lithium battery. The procedure for battery replacement is as follows.

a) Turn the power off.

b) Remove the battery cover.

c) Remove the old battery and disconnect the connector.

d) Connect the new battery and put it in the Battery compartment.

e) Replace the battery cover.

It is recommended to contact local distributors or manufacturer to replace the battery.

7.3 Transportation

7.3.1 Excessive impact and shake should be prevented in transportation. Lay it carefully and lightly and don't invert it.

7.3.2 Don't put it together with dangerous goods during transportation.

7.3.3 Avoid solarization and getting wet in rain and snow during transportation.

8 Environmental protection

Please dispose according to the local laws.

9 European authorized representative

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

10 After service

From the date this equipment has been sold, based on the warranty card, we will repair this equipment free of charge if there are quality problems. Please refer to the warranty card for the warranty period.

11 Symbol instruction



Class II equipment Date of manufacture



Type BF applied part



Ordinary equipment



Used indoor only



Appliance compliance WEEE directive





Handle with care



12 Statement

All rights of modifying the product are reserved to the manufacturer without further notice. The pictures are only for reference. The final interpretation rights belong to GUILIN WOODPECKER MEDICAL INSTRUMENT CO., LTD. The industrial design, inner structure, etc, have claimed for several patents by WOODPECKER, any copy or fake product must undertake legal responsibilities.

13 EMC - Declaration of conformity

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.

Guidance and manufacturer's declaration - electromagnetic emissions			
The model Woodpex X is intended for use in the electromagnetic environment specified below. The			
customer or the user of the model should assure that it is used in such an environment.			
Emissions test Compliance Electromagnetic environment - guidance			

RF emissions CISPR 11	Group 1	The model Woodpex X use RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in near- by electronic equipment.
RF emissions CISPR11	Class B	
Harmonic emissions IEC 61000-3-2	Complies	The model Woodpex X is suitable for used in domestic establishment and in establishment directly connected
Voltage fluctuations / flicker emissions IEC 61000-3-3	Complies	to a low voltage power supply network which supplies buildings used for domestic purposes.

Guidance	&	Declaration -	electromagnetic	immunity
Guiuanee	~	Deciaration	ciecti omagnetic	munuty

The model Woodpex X is intended for use in the electromagnetic environment specified below. The customer or the user of the model Woodpex X should assure that they are used in such an environment.

Immunity test	IEC 60601 test level	Compliance level	Electromagnetic environment - guidance
Electrostatic dis- charge (ESD) IEC 61000-4-2	±8 kV cantact ±2 kV, ±4 kV, ±8 kV, ±15 kV air	±8 kV contact ±2 kV, ±4 kV, ±8 kV, ±15 kV air	Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30 %.
Electrical fast tansient/burst IEC 61000-4-4	±2kV for power supply lines ±1 kV for lnput/out- put lines	±2kV for power supply lines	Mains power quality should be that of a typical commercial or hospital environment.

Surge IEC 61000-4-5	$\pm 0.5 \text{ kV}, \pm 1 \text{ kV}$ line to line $\pm 0.5 \text{ kV}, \pm 1 \text{ kV}, \pm 2$ kV line to ground	±0.5 kV, ±1 kV line to line	Mains power quality should be that of a typical commercial or hospital environment.		
Voltage dips, short interruptions and voltage variations on power supply input lines IEC 61000-4-11			Mains power quality should be that of a typical commercial or hospi- tal environment. If the user of the model Woodpex X require continued operation during power mains inter- ruptions, it is recommended that the model Woodpex X be powered f rom an uninterruptible power supply or a battery.		
Power f requency (50/60 Hz) magnetic field IEC 61000-4-8	30 A/m	30 A/m	Power f requency magnetic f ields should be at levels characteristic of a typical location in a typical com- mercial or hospital environment.		
NOTE U_T is the a.c. mains voltage prior to application of the test level.					

Guidance & Declaration - Electromagnetic immunity			
The model Woodpex X is intended for use in the electromagnetic environment specified below. The customer or the user of that the model Woodpex X should assure that it is used in such an environment.			
Immunity test	IEC 60601 test level	Compliance level	Electromagnetic environment - guidance

Conducted RF IEC 61000-4-6	3 Vrms 150 kHz to 80 MHz 6 Vrms in ISM and amateur radio bands	3 Vrms 150 kHz to 80 MHz 6 Vrms in ISM bands	Portable and mobile RF communications equipment should be used no closer to any part of the model Woodpex X, including cables, than the recommended separation dis-
Radiated RF	3 V/m	3 V/m	tance calculated from the equation applicable
IEC 61000-4-	80 MHz to 2.7	80 MHz to 2.7	to the f requency of the transmitter.
3	GHz	GHz	
	385MHz-	385MHz-	If higher IMMUNITY TEST LEVELS than
	5785MHz Test	5785MHz Test	those specified in Table 9 are used, the mini-
	specifications	specifications	mum separation distance may be lowered.
	for	for	Minimum separation distances for higher
	ENCLOSURE	ENCLOSURE	IMMUNITY TEST LEVELS shall be calcu-
	PORT	PORT	lated using the following equation:
	IMMUNITY to	IMMUNITY to	$E = [6/d] \times P^{1/2}$
	RF wireless	RF wireless	Where P is the maximum power in W, d is the
	communication	communication	minimum separation distance in m, and E is
	equipment	equipment	the IMMUNITY TEST LEVEL in V/m.
	(Refer to table 9	(Refer to table 9	
	of IEC 60601-1-	IEC 60601-1-	
	2:2014+A1:2020)	2:2014+A1:2020)	
NOTE 1 At 80 MHz and 800 MHz. the higher f requency range applies.			
NOTE 2 These	guidelines may not	apply in all situation	ons. Electromagnetic propagation is af fected

by absorption and ref lection f rom 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 is used exceeds the applicable RF compliance level above, the model should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as reorienting or relocating the model.

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

Recommended separation distances between portable and mobile RF communications equipment and the model Woodpex X

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

Rated maximum out-	Separation distance according to frequency of transmitter m		
put power of trans-	150kHz to 80MHz	80MHz to 800MHz	800MHz to 2,5GHz
mitter W	$d=1.2 \times P^{1/2}$	$d=1.2 \times P^{1/2}$	$d=2.3 \times P^{1/2}$
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 distanced 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 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.

Apex locator in the above specified electromagnetic environment, it will be safe, and it can provide the basic properties such as article 1.6.1-1.6.3;

1. Measurement of pulpitis, pulp necrosis, periapical periodontitis and tooth length.

2. Measurement of the tooth length before restoration of post crown.

3. Measurement of the tooth length of transplantation and retransplantation.

Cautions:

1. Cautions: User must regard EMC, please install and put in service the model according to the EMC information provided in the accompanying documents

2. Cautions: Portable and mobile RF communications equipment can affect medical electrical equipment.

3. Use is not specified for the Apex locator the model of the adapter(Optional), charging cable(Optional), measuring wire, file clip may increase the radiation quantity or reduce the interference ability of the Apex locator system. A list of all cables and maximum lengths of cables is as follows, transducers and other accessories with Guilin Woodpecker Medical Instrument Co., Ltd. claims compliance with the requirements of Emission and Immunity. Please use original accessories.

Serial Number	Accessories name	Cable length	Whether shielding
1	adapter(Optional)	2	No
2	measuring wire	1.7	No

3	file clip	0.2	No

4. Cautions: The use of accessories, transducers and cables other than those specified, with the exception of transducers and cables sold by Guilin Woodpecker Medical Instrument Co., Ltd. as replacement parts for internal components, may result in increased Emissions or decreased Immunity of the model.

5. The model should not be used adjacent to or stacked with other equipment and that if adjacent or stacked use is necessary, the model should be observed to verify normal operation in the configuration in which it will be used.

6. The accessories adapter(Optional), charging cable(Optional), battery, measuring wire, file clip of Apex locator the model may affect the radiation quantity. The original accessories are in compliance with the requiments of the IEC 60601-1-2. Please use original accessories.

Scan and Login website for more information





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Zone, Guilin, Guangxi, 541004 P. R. China Sales Dept.: +86-773-5873196/2350559 After-sales Service Dept.: +86-0773-5827898 E-mail: woodpecker4@glwoodpecker.com Website: http://www.glwoodpecker.com Website: http://www.glwoodpecker.com ECREP Borkstrasse 10 - 48163 Muenster · Germany

Apex Locator Warranty Card

Name of Customer	(For Distributor)	
Tel		
Address Details	Gulin Woodpecker Medical Instrument Co.,Lt Information Industrial Park, Gulin National High-Te Zone, Gulin, Guangus, 541004 P. R. China Sales Dept. + 86-773-6973196/2305699 After-sales Service Dept. + 86-0773-6827898	
Postal Code	E-mail: woodpecker4@gjwoodpecker.com Website: http://www.glwoodpecker.com	
Model	Distributor:	
Product No.		
Purchase Date		
Contact Person	Seal	
Date	Maintenance Record/Repairer	

Warranty Instruction

1. Period validity:

We offer two years free repair on main unit and six months on spare parts from the date of purchase, warranty card must be attached.

2. Range of warranty:

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

3. The following are beyond our warranty:

a. The damage caused by disobeying the operation instruction or lack of the needed condition.

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

c. The damage caused by unadvisable transportation or preservation.

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

e. Device failures caused by the use of non-original accessories are not under warranty.

Apex Locator Warranty Card

The dashe of Customer		(Return to Manufacturer)	
Tel			
Address Details		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	
Postal Code		After-sales Service Uept: +86-07/3-982/998 E-mail: woodpecker/@glwoodpecker.com Website: http://www.glwoodpecker.com	
Model		Distributor:	
Product No.			
Purchase Date			
Contact Person		Seal	
Date	Maintenance Record/F	Maintenance Record/Repairer	

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

e. Device failures caused by the use of non-original accessories are not under warranty.

Harm of fake products

p---- and *pre* are two brands of Guilin woodpecker medical instrument company. Recently, growing fake ultrasonic scaler handpieces, tips curing lights are produced and sold on the market, which do harm to users' interest. On this issue, We Woodpecker will crack down fake products and provide safe and secure medical instrument products.

1. Harm of fake ultrasonic scaler handpieces.

1.1 Fake handpieces with poor-designed inner structure can lead to frequent power leakage, which may cause medical accidents.

1.2 Material used on fake handpieces don't pass biocompatible test, which can easily lead to irritability and poisoning.

1.3 Fake handpieces have quality problems of overheating, non-vibration and cracking, which cause ultrasonic scalers out of order.

1.4 Fake handpieces can' t be compatible with ultrasonic scalers, thus leading to circuit burn out.

2. Harm of fake scaler tips.

2.1 Fake tips are low in toughness, poor in resistance and easy to crack, thus easily cause medical accident.

2.2 Fake tips' screw threads are roughly processed, which can cause handpiece's screw loosing and cracking.

2.3 Material used on fake tips is inferior and easily rusting, which can cause infection of patient.

2.4 Fake tips have used problem of poor water-spraying, bad screw-thread fit and water leaking, which leads ultrasonic scalers work wrongly.

3. Harm of fake curing light.

3.1 Fake curing light's batteries can cause self-ignite, even explosion with poor-quality material and no complete charging management.

3.2 Light intensity of fake curing light is not constant, when battery level goes down under 60%, it would lead to incomplete solidification of resin, causing secondary dental caries.

