

OPERATION INSTRUCTION



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for more information

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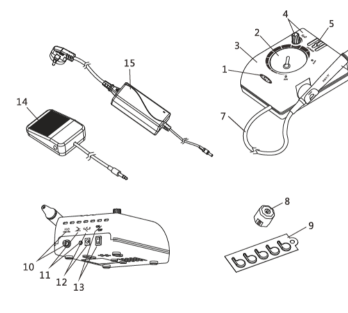
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K08B ULTRASONIC SCALER

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COMPONENTS



- 1 Model and LOGO
- 2 Vibration intensity knob and icon
- 3 Unit
- 4 Water volume knob and icon
- 5 Power LEDs
- 6 Sealed handpiece
- 7 Sealed cable
- 8 Tip wrench
- 9 Tips
- 10 Fluid circuit connection and icon
- 11 Foot switch connection and icon
- 12 External power supply connection and icon
- 13 Power switch and icon
- 14 Foot switch
- 15 External power supply

Copyright and statement

Copyright © Guilin Veirun Medical Technology Co., Ltd.

Congratulations to become valued customers of Guilin Veirun Medical Technology Co., Ltd. Thanks for using K08B ultrasonic scaler. It will bring you the new experiences and conveniences.

This instruction manual is including the latest information when printing. Guilin Veirun Medical Technology Co., Ltd reserves the rights to change the design of the equipment, the technique, fittings the instruction manual and the content of original packing list at any time without notice. If there are some differences between blueprint and real product, take the real product as the norm.

This instruction manual is protected by copyright law. Without any official authorization, any form of duplication, copy or translated into another language is not allowed.

Read the instruction manual originally delivered with each of the components before using this product, and follow the instruction manual to operate. Otherwise, Guilin Veirun Medical Technology Co., Ltd does not take any responsibility for any errors and damage to the product because of violation operation.



Note: Guilin Veirun Medical Technology Co., Ltd does not promise any implied guarantee to the product special use for merchantability and applicability.

If you need after-sale service support, please contact with authorization dealer or manufacturer.

Wainning

The device is not suitable for use in the presence of flammable anesthetic mixtures with air or with oxygen or nitrous oxide.

The device requires no calibration.

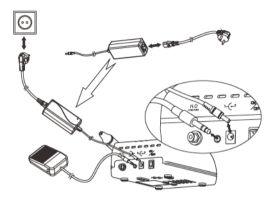
The device is not repairable and contains no user serviceable parts.

No modification of this equipment is allowed. (3rd Edition)

The user must check that the equipment functions safely and see that it is in proper working condition before being used.

The manufacturer does not require such preventive inspections by other persons.

INSTALLATION AND CONNECTION





Always check that the unit is installed in a safe and stable position. An eventual fall may suddenly pull on the cord and the handpiece, hurting the patient and the user, and damaging the unit.

Check that the rated voltage of the power supply corresponds to the local line voltage to prevent damaging the unit.

Use the unit only in a FI protected mains supply (FI = Residual current protection).

For USA or Canada: connect only to an Hospital Grade outlet.

PREPARING FOR TREATMENT



Clean, disinfect and sterilize the different pieces and accessories of this product before each use. Please refer to the information provided in the operating instructions. Non-sterile pieces and accessories may cause bacterial or viral infections.

MANUAL FOR WRENCH



With a special structure design, wrench makes sure that users can handling the tip effectively during Using it, and protects user's hand from hurting in the process of loading and unloading the tip.

Sterile environment: Temperature 135 degrees, Pressure 0.22Mpa, in the steam.

Steps:

- (1)Align the legs of the wrench to the tip.
- (2)Install the tip: grip the handle, rotate the tip in right-handwise till the tip dose not turn round anymore.
- (3)Unloading the tip: grip handle then rotate the tip in counter-clockwise by wrench to remove it.
- (4)Once after using, please put the wrench into disinfection cabinet to disinfect.
- (5)After disinfection, the surface temperature of the wrench is too high to be used again, so users must wait until it is cool down, avoiding hurt.
- (6) When not use the wrench, put it at some place that is dry and ventilated, and keep it clean.

SETTING THE ULTRASOUND POWER



Using the vibration intensity knob increases or decreases the power of the ultrasound. Adjust vibration ntensity on-demand, generally, the medium is right, it also can be adjusted in the clinical course according to the sensitivity of patients and the hardness of the dental stone.



Adjust power to the lowest when just turn on the host.

SETTING THE LIQUID FLOW RATE



Depress the foot switch, and the tip starts vibrating, and then revolve the water volume knob and the water spraying puffs, it will cool down the handle and clean the surface of teeth.

Never work dry to avoid heat damage on the tooth. The tip of the instrument immediately heats up



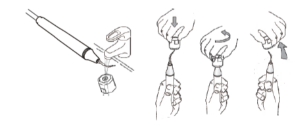
when used dry.

Adjust water volumn to larger when just turn on the host.

FOOT SWITCH FUNCTIONS



END OF TREATMENT





Clean the instrument with running water before disinfection.

Please conform to the recommendations of the "Reprocessing Instructions" manual delivered with your product regarding the procedures of cleaning, disinfecting and sterilizing of the components.

CLEANING, DISINFECTING AND STERILIZING

Handpiece and disinfection

Select disinfectant to wipe and cleaning. Such as iodine or alcohol or glutaraldehyde disinfectant and so on.

Notices

- (1)Remove the tip after working every time.
- (2)Be sure to remove the tip from the handpiece before disinfecting, and avoid mixing it together with other equipment.
- (3)In the process of using and disinfection, please feel free to pay attention that the handpiece is with external damages or not. Prohibit from applying and protected oil on the surface of the handpiece.
- (4)Prohibit to soak the handpiece into disinfectant, for example iodine, alcohol, glutaral dehyde and so on.

Tip Sterilization

The tips can be sterilizated by alcohol or disinfectant wipes, it can also be sterilizated by HTY.

Wrench Sterilization

It can be sterilizated by any neutral disinfectant without any corrosivity and HYT.

Wrench and tips Cleaning

The wranch and tips can be cleaned in the ultrasonic cleaner.

Notices

Prohibit to disinfect in the following ways:

- * Put the wranch into boiling solution
- * Soak the wrench and tips into disinfectant ,for example iodine,alcohol ,glutaraldehyde and so on.
- * Put the wrench and tips into oven or microwave to bake in high-temperature.

Follow present-day regulations enforced in the country about reprocessing.

CLEANING THE UNIT



Clean the unit only with an alcohol-based, commercially-available, colorless disinfectant (ethanol, isopropanol). The use of scouring powder or an abrasive sponge will damage its surface. The unit is not protected against water spray. It cannot be sterilized.

DISINFECTING THE CIRCUIT



It is recommended to disinfect the liquid circuit daily.

Disinfection is necessary to reduce crystalline deposits and the number of bacteria in the

liquid circuit of the unit; The following procedure achieves a 4 log reduction of target pathogens:

- 1. Make sure the water you are using is distilled(or dematerialized) water.
- 2. Remove the handpiece. Revolve the water volume knob to the max point.
- 3. Turn on the scaler and depress the foot switch.
- 4. Allow the disinfecting water to rinse the circuit for 3 to 5 minutes.

MAINTENANCE



Always use VRN original parts. Using non-original instruments may damage the unit, and practitioner or patient may be injured.

SAFETY PRECAUTIONS

VRN and the dealer of this product accept no liability for direct or consequential injury or damage resulting from improper use, arising in particular through non-observance of the operating instructions, or improper preparation and maintenance.



Contraindication: ultrasonic oscillations may prevent cardiac pace makers and defibrillators from functioning properly. Therefore, we recommend that patients with a cardiac pace maker or a defibrillator should not be treated with this product.

Use for the intended purpose only: before using the product, make sure that you have studied the operating instructions. This also applies to any equipment used with this product. Failure to observe the operating instructions may result in the patient or user suffering serious injury or the product.

Risk of explosion: do not use this product in the presence of inflammable anaesthetics or gases.

This product must be used only by trained and qualified personnel.

Clean, disinfect and sterilize the different pieces and accessories of this product before each use.

Please refer to the information provided in the operating instructions. Non-sterile pieces and accessories may cause bacterial or viral infections.

This product has been investigated with regard to safety from electrical shock and fire hazard.

ETL (Electrical Testing Laboratories) has not investigated the physiological effects.

Always examine the product for damage before commencing treatment. Damaged accessories or a damaged unit must not be used and must be replaced. Use original VRN spare parts and accessories only.

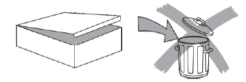


Have this product exclusively repaired by an approved VRN repair center.

Use exclusively the external power supply delivered with the VRN unit.

Certain types of mobile telecommunication equipment could potentially interfere with this oroduct. The separation distances recommended in the "Electromagnetic compatibility" manual must be taken into account.

STORING THE PRODUCT





 $\label{the conditions} \textbf{Keep the original packaging until the product is to be disposed of permanently}.$

It can be used for shipping or storing at any time.

Should you wish to put your product out of use for an extended period of time:

- Proceed as described in the "Cleaning, disinfecting and sterilizing" chapter,
- · Pack the product and all accessories in the original packaging.

Please refer to the storage and transport conditions in the "Technical data" chapter.

DISPOSAL

The product must not be discarded in domestic household waste.

Should you wish to definitively dispose of the product, please comply with the regulations which apply in your country.

Waste Electrical and Electronic Equipment belonging to customers located in the European Union may be shipped to VRN for recycling in accordance to the WEEE regulations. The costs of recycling, exclusive of shipping fees, are covered by VRN.

WARRANTY

Damages due to non-adherence to the operating instructions or wear out of parts are excluded from warranty.



The warranty of your product will be cancelled if you try to open it.

ACCESSORIES

Accessories are available from VRN or any authorized dealers. Please contact your customer service directly.

VRN SERVICE

Should your product need additional servicing or repairs, please send it to your dealer or to an authorized VRN Repair Center.

In the case of non-authorized repairs or damages due to non-adherence to the operating instructions, VRN accepts no liability whatsoever. This will also void the warranty.

It is best to ship your product in the original packaging. It protects your product against damage during shipment.



Risk of transmitting micro-organisms! To protect the personnel of the repair center and for safety reasons during transport and shipment, the product and the accessories returned to the factory for repair or revision must be cleaned, disinfected and sterilized in accordance with the operating manual. Repair can be refused for

When sending your product directly to the approved VRN repair center, please include the name of your dealer. This simplifies processing.

SYMBOLS

¥vrniii°	Manufacturer's logo	\triangle	Caution! Read the operation instruction		Class II equipment	†	Applied part, type BF
Z	Disposal	•••	Manufacturer		Used indoor only	SN	Serial number
135°C	Autoclavable		Storage Humidity	-20 °	Storage Temp	T	Keep dry
	Fragile		Refer to instruction manual / booklet	EC REP	Authorised representative in the European community	C € 0123	CE marking:refers to directive 93/42 EEC,including EN60601-1 and EN60601-1-2
	Electrical Outlet	<u>></u>	Foot switch connection	H₂O 0.01Mpa-0.5Mpa	Water supply	ON OFF	Power Switch

TECHNICAL DATA

Description Models

Classification EN 60601-1

Classification 93 / 42 EEC

Operating mode Power Supply

Adaptor Input Adaptor Output

Power consumption

External power supply Ultrasonic specifications:

Maximum output Frequency range

Weight

Dimensions in mm (HxWxD)

Operating conditions

Storage and transport conditions

Manufacturer Guilin Veirun Medical Technology Co., Ltd. K08B

- Class II
- Applied part, Type BF
- IP 20, unit
- IP X1, foot pedal

Not category AP/APG equipment

Class IIa

Continuous operation

100-240 VAC / 50-60 Hz. 1.3A

30VDC 30VA~48 VA

45W 30V 2m R5.5- 2.5+

8 Watts 25-31kHz Approx. 1.45Kg 75 x 150 x 160 +10 °C to +40 °C

30% to 75% relative humidity

-10 °C to +40 °C

10% to 95% relative humidity 500 hPa to 1 060 hPa air pressure

TROUBLESHOOTING

Fault	Possible cause	Solutions
The scaling tip doesn't vibrate	The power plug is in loose contact	Make the plug insert to the socket well
and there is no water flowing out when stepping on the foot Switch	The foot pedal is in loose contact (wire foot Switch)	Insert the foot switch to its socket tightly
	The fuse in the main unit is broken	Contact the dealers or us
	The battery for wireless control foot Switch died	Change the new batteries
The scaling tip does not vibrate	The tip is in loose contact	Screw the tip on the handpiece tightly
but there is water flowing out when stepping on the foot Switch	The connect plug between the handpiece and the circuit board is in loose contact	Contact the dealer or us
	Handpiece problem	Send the handpiece to dealer or us to repair
	Cable problem	Contact the dealer or us
The scaling tip vibrates but there	The water control knob is not on	Turn on the water control knob [note 1]
is no spray when stepping on the foot Switch	There is impurity in the electric-magnetic valve	Take the electric-magnetic valve apart or contact the dealer or us
	The water system is blocked	Clean the water pipe by multi-function syringe [note 2]
There is still water flowing out	The electric-magnetic valve	Contact the dealer or us
after the power is off	problem	
The handpiece generates heat	The water control knob is in a low setting	Turn the water control knob to a higher grade [note 1]

Fault	Possible cause	Solutions
The amount of spouting water is	The water control knob is in a low	Turn the water control knob to a higher grade
too little	setting	[note 1]
	The water pressure is not high enough	Make the water pressure higher
	The water system is blocked	Clean the water pipe by multi-function syringe [note 2]
The vibration of the tip becomes weak	The tip has not been screwed on to the handpiece tightly	Screw the tip tightly
	The coupling between the handpiece and the cable is not dry	Dry it by the hot air
	The tip is damaged [note 3]	Change the new scaling tip
There is water leaking from the coupling between the handpiece and the cable	The waterproof O-ring is damaged	Change the new waterproof O-ring
The U-file does not vibrate	The screw is loose	Tighten it
	Endochuck is damaged	Change the new endochuck
There is noise coming from the endochuck	The screw is loose	Tighten it



Notice:

If the problem still cannot be solved, please contact with local dealer or manufacturer.



Notes

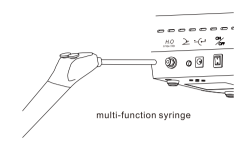
【note 1】

Turn the water control knob as symbol showed. It comes to the min, on the contrary direction, it comes to the max.

[note 2]

- 1)Clean the water pipe with the multi-function syringe of the dental unit;
- 2)Disconnect the water pipe from main unit;
- 3)Get through to the power and turn on the power switch;
- 4)Connect the multi-function syringe of dental unit to the water pipe;
- 5)Disassemble the tip or handpiece;
- 6)Step on the foot switch;
- 7)Turn on the switch of the multi-function syringe, press the water into the machine and the impurity blocked in the water pipe can be eliminated.

[note 3] If the scaling tip has been screwed on tightly and there is fine spray too, the following phenomena show that the scaling tip is damaged:



TROUBLESHOOTING FOR THE ELECTRIC-MAGNETIC VALVE

If the water is not clean or the electric-magnetic valve works too long, there will be the scale and impurity generated on the inside wall and valve plug. It will cause the water way to be blocked, and could repair as following steps:

- 1) Follow the [note 2]
- 2) Uninstall the electric-magnetic valve by following steps:
- a)Turn off the power, and unplug the power cable;

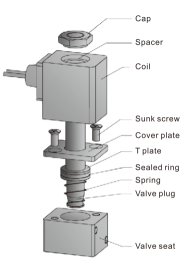
b)Unscrew the cap, and move out the hood of main unit; c)Unscrew the cap on the electric-magnetic valve, and Uninstall the electric-magnetic valve as shown on

Figure;

d)Take out the valve plug, and clean out the impurity on the inside wall;

e)Install the parts as shown on the figure.

f)Ensure the water way is working by repeat [note 2]



EMC DECLARATION

Table 1

Guidance and manufacturer's declaration - electromagnetic emissions				
The [KO8] is intended for use in the electromagnetic environment specified below. The customer or the user of the [KO8] should assure that it is used in such an environment				
Emissions test	Compliance	Electromagnetic environment - guidance		
RF emissions CISPR 11	Group 1	The [K08] uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.		
RF emissions CISPR 11	Class [B]	The [K08] is suitable for use in all establishments other than domestic, and may		
Harmonic emissions IEC 61000-3-2	Class A	be used in domestic establishments and those directly connected to the public low-voltage power supply network that supplies buildings		
Voltage fluctuations/ flicker emissions IEC 61000-3-3	Complies	used for domestic purposes, provided the following warning is heeded: Warning: This equipment/system is intended for use by healthcare professionals only. This equipment/ system may cause radio interference or may disrupt the operation of nearby equipment. It may be necessary to take mitigation measures, such as re-orienting or relocating the [KO8] or shielding the location.		

Table 2

Guidance and manufacturer's declaration - electromagnetic emissions					
The [K08] is intended for use in the electromagnetic environment specified below. The customer					
or the us	er of the [K08] sho	uld assure that it is u	sed in such an environment		
Immunity Test	Immunity Test IEC 60601 Compliance level Electromagnetic environment -				
	Test level		guidance		
Electrostatic	±6 kV contact	±6 kV contact	Floors should be wood, concrete or		
discharge (ESD)	±8 kV air	±8 kV air	ceramic tile. If floors are covered		
IEC 61000-4-2			with synthetic material, the relative		
			humidity should be at least 30 %		
Electrical fast	±2 kV for power	±2 kV for power	Mains power quality should be that		
transient/burst	supply lines	supply lines	of a typical commercial or hospital		
IEC 61000-4-4	±1 kV for	±1 kV for	environment.		
	input/output	input/output			
	lines	lines			
Surge	±1 kV line(s) to	±1 kV line(s) to	Mains power quality should be that		
IEC 61000-4-5	line(s)	line(s)	of a typical commercial or hospital		
	±2 kV line(s) to	±2 kV line(s) to	environment.		
	earth	earth			

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Valence dine decut	4F0/ 11	4E0/ 11	Bitating a service and the about all his short	
Voltage dips, short	<5% U _⊤	<5% U _T	Mains power quality should be that	
interruptions and	(>95% dip in U _T)	(>95% dip in U _T)	of a typical commercial or hospital	
voltage variations	for 0.5 cycle	for 0.5 cycle	environment. If the user of the [K08]	
on power supply			requires continued operation during	
input lines	40% U _T	40% U _T	power mains interruptions, it is	
IEC 61000-4-11	$(60\% \text{ dip in U}_T)$	(60% dip in U _T)	recommended the [K08] be	
	for 5 cycle	for 5 cycle	powered from an uninterruptible	
			power supply or a battery.	
	70% U _T	70% U _T		
	(30% dip in U _T)	(30% dip in U _⊤)		
	for 25 cycle	for 25 cycle		
	,	·		
	<5% U _T	<5% U _T		
	(>95% dip in U _⊤)	(>95% dip in U _T)		
	for 5s	for 5s		
Power frequency	3 A/m	3 A/m	Power frequency magnetic fields	
(50/60Hz)			should be at levels characteristic of	
magnetic field			a typical location in a typical	
IEC 61000-4-8			commercial or hospital	
			environment.	
NOTE U_T is the a.c. mians voltage prior to application of the test level.				

Table3

les .				
Guidance and manufacturer's declaration - electromagnetic emissions				
The [KO8] is intended for use in the electromagnetic environment specified below. The customer or the				
	user of the	e [K08] should	assure that it is used in such an environment	
Immunity	IEC 60601	Compliance	Electromagnetic environment - guidance	
Test	Test level	level		
			Portable and mobile RF communications equipment should	
			be used no closer to any part of the [KO8], including cables,	
			than the recommended separation distance calculated from	
			the equation applicable to the frequency of the transmitter.	
			Recommended separation distance	
Conducted RF	3Vrms	3V	$d = 1,2\sqrt{P}$	
IEC 61000-4-6	150KHz to			
	80MHz			
Radiated RF	3 V/m	2 1//22	$d = 1.2\sqrt{P}$ 80 MHz to 800 MHz	
IEC 61000-4-3	80MHz to	3 V/m		
IEC 61000-4-3	2.5GHz		$d = 2.3\sqrt{P}$ 800 MHz to 2,5 GHz	
	2.5GH2			
			where p is the maximum output power rating of the	
			transmitter in watts (W) according to the transmitter	
			manufacturer and <i>d</i> is the recommended separation distance	
			in metres (m)	
			Field strengths from fixed RF transmitters, as determined by	

an electromagnetic site survey, a should be less than the compliance level in each frequency range. Interference may occur in the vicinity of equipment marked with the following symbol:



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

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

^a Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the [K08] is used exceeds the applicable RF compliance level above, the [K08] should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as re-orienting or relocating the [K08].

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

Table 4

Recommended separation distances between portable and mobile RF communications equipment and the [K08]

The [K08] is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the [K08] can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the [K08] as recommended below, according to the maximum output power of the communications equipment.

Rated maximum output power of transmitter	Separation distance according to frequency of transmitter					
W	150 blis as 90 balls 900balls as 900balls 900balls as 3 50lb					
VV	150 kHz to 80 MHz 80MHz to 800MHz 800MHz to 2.5GHz					
	$d = 1.2\sqrt{P}$	$d = 1,2\sqrt{P}$	$d = 2,3\sqrt{P}$			
0,01	0.12	0.12	0.23			
0,1	0.38	0.38	0.73			
1	1.2	1.2	2.3			
10	3.8	3.8	7.3			
100	12	12	23			

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

NOTE 1 At 80 MHz and 800 MHz, the separation distance for the higher frequency range applies. NOTE 2 These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

