



# **OPERATION INSTRUCTION**

I 01 / I 02

**BUILT-IN ULTRASONIC SCALER** 

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# 1. SYMBOLS













Caution! Read the operation instruction

Class II equipment

Applied part, type B

Disposal Manufacturer' s logo

Manufacturer











Used indoor only

Serial number

Autoclavable CE marking:refers to directive 93/42EEC, including EN60601-1 booklet and EN60601-1-2

Refer to instruction manual /



Storage Humidity







Fragile

Authorised representative in the European community

2. DIRECTIONS AND INSTALLATION

## 2.1 INTRODUCTION

I 01, I 02 built-in ultrasonic scaler produced by Guilin Veirun Medical Technology Co., Ltd., is mainly used for removing the calculus and stains on teeth surface along with dental units.

I 01 is attached with a sealed handpiece, I 02 is attached with a detachable handpiece.

# 2.2 COMPONENTS

2.2.1 The Packing List shows exactly what is included in your scaler.

# 2.2.2 Mainly Components

The device is composed of control circuit, a handpiece and scaling tips.

#### 2.2.3 Applicable Scope

Installed in the dental unit, used for removing the calculus and stains on teeth surface.

#### 2.3 TECHNICAL SPECIFICATIONS

- Power Supply: AC 24V (1±10%), 50~60Hz or DC 30V (1±5%)
- Input Power: ≤40VA
- Fuse Protector: 250V, 2A
- Tip Principal Oscillation Offset: 1µm (min), -50% deviation

100µm (max), +50% deviation

- Half-excursion Force: 0.1N (min), -50% deviation
   2N (max), +50% deviation
- Tip Vibration Frequency: 28KHz±3KHz
- Ultrasonic Output Power: 3W~20W
- Weight of Main Unit: 0.3kg
- Weight of Adapter: 0.2kg (optional)
- Operating Mode: Continuous operating
- Electric Shock Protection Type: Class II equipment
- Electric Shock Protection Degree: Type B Applied Part
- Liquids Protection Degree: IPX 0
- Safety Degree of Usage in the presence of a Flammable Anaesthetic Mixture
  with air or with Oxygen or Nitrous Oxide: not suitable for being used in the
  presence of a flammable anaesthetic mixture with air or with oxygen or nitrous
  oxide
- Working Condition: Environment Temp 5°C~40°C, Relative Humidity ≤90°

#### 2.4 CONNECTION AND INSTALLATION

#### 2.4.1 Connection and Installation

(The connection instruction is shown by a picture in the attachment.)

- Port 1 connects with DC 30V adaptor with P1J standard interface mode. The adaptor provided by Guilin Veirun Medical Technology Co., Ltd. is recommended;
- Port 2 and Port 3 connect with electromagnetic valve;
- Port 4 and Port 5 connect with AC 24V from the inside of dental unit;
- Port 6 and Port 7 connect with gas control switch;
- Port 8 to Port 15 are free;
- Port 16 to Port 18 connect with potentiometer;

• Port 19 and Port 20 connect with the handpiece.

#### 2.4.2 Notice

- The gas control switch, pneumatic water value and foot switch are equipped by the dental unit manufacturer or the user. The holes for installing the potentiometer and cable are drilled by the dental unit manufacturer, device supplier or the user.
- Keep enough space for dispersing heat of ultrasonic generator.
- Before turning on the device, turn the potentiometer knob to the minimum and the water control knob the maximum.
- The frequency of ultrasonic scaler is extremely high. Under normal working state of scaling tips, a light touch and a certain to-an-fro motion will eliminate the tartar without obvious heat. Overexertion and longtime lingering are forbidden.

# 3. FUNCTION AND OPERATION

#### 3.1 WORKING PRINCIPLES

#### 3.1.1 Instruction

Built-in Ultrasonic Scaler mainly consists of control circuit, handpiece and scaling tips. Its power and water are controlled by the foot switch and the handpiece holder of dental unit.

# 3.1.2 Working Principles

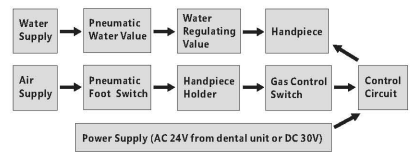


Figure 1

Pull the handpiece out of the Handpiece Holder, step on the Foot Switch, then the air is on. The air gets though the Gas Control Switch and turns on the scaler, the Water Regulating Value is opened at the same time.

#### 3.2 PARTS INSTRUCTIONS

## 3.2.1 Usage Instruction of Wrench

When installing a scaling tip, insert the scaling tip into the hole of the wrench, hold the handpiece and turn the wrench clockwise till the tip stop. The scaling tips must be tightened up to get the best using effects.

When uninstalling a scaling tip, insert the scaling tip, hold the handpice and turn the wrench anti-clockwise.



Figure 2

## 3.2.2 Usage Instruction of Main Parts of Sealed Handpiece



Figure 3

- Cap: Removable, can be screwed out to make the pole cleaned by disinfectant.
- Ruble Ring: Can be removed regularly and cleaned by disinfectant.
- Handpiece: Can not be autoclaved, can only be cleaned by disinfectant.
- Cable: connects the handpiece and water supply and power supply.

# 3.2.2 Usage Instruction of Main Parts of Detachable Handpiece



Figure 4

- Cap: Removable, can be screwed out to make the pole cleaned by disinfectant.
- Ruble Ring: Can be removed regularly and cleaned by disinfectant.
- Handpiece: Can be autoclaved.
- Logo and Symbols: Manufacturer's logo. Can be sterilized by high temperature (135°C) and high pressure (0.22 MPa).
- Cable: connects the handpiece and water supply and power supply.

# 4. MAINTENANCE AND DISINFECTION

## 4.1 STERILIZATION OF SCALING TIPS

Scaling tips can be cleaned by the ultrasound cleaner and sterilized by alcohol cotton or disinfected cloth, and also can be sterilized by high temperature and high pressure.

#### 4.2 STERILIZATION OF WRENCH

The wrench can be cleaned by the ultrasound cleaner and sterilized by neutral non-corrosive disinfectant, and also can be sterilized by high temperature and high pressure.

#### 4.3 STERILIZATION OF HANDPIECE

#### 4.3.1 Sterilization of Sealed Handpiece (I 01)

• The sealed handpiece can be sterilized by wiping with some disinfectant such as iodine, alcohol and glutaraldehyde

- During sterilizing the sealed handpiece, please notice whether the surface of the handpiece is damaged. Do not smear any protective oil on the surface of the handpiece.
- Do not soak the handpiece in any disinfectant.

#### 4.3.1 Sterilization of Detachable Handpiece (I 01)

- Autoclaved by the high temperature and pressure:
  - 121 °C, 1 bar (0.1 Mpa), 20 minutes;
  - 135 °C, 2.2 bar (0.22 Mpa), 15 minutes.
- Pull out the handpiece and uninstall the scaling tips after each use.
- Pack the handpiece in sterile gauze or sterile bag before sterilization.
- Reuse the handpiece after it cool down naturally to avoid burning hands.
- Notice:
- a) Before sterilization, empty the cleaning liquid in the handpiece with compressed air.
- b) Be sure that the scaling tip has been uninstalled from the handpiece. It cannot be sterilized along with other spare parts.
- c) Notice whether the surface of the handpiece is damaged during the treatment and sterilization. Do not smear any protective oil on the surface of the handpiece.
- d) There are two waterproof O-rings at the end of handpiece. Please lubricate them with dental lube frequently, as sterilization and repeated pulling and inserting will reduce their life-span. Change a new one once it is damaged or worn excessively.
- e) Sterilizable parts can be sterilized at least 250 times.
- f) The following sterilizing methods are prohibited:
  - Put the handpiece into any liquid for boiling;
- Soak the handpiece in any disinfectant.
- Put the handpiece into oven or microwave oven for baking.

#### 4.4 TROUBLE SHOOTING

FAULT	POSSIBLE CAUSES	SOLUTION	
	The plug is loose or wrong contact.	Connect as the shown picture.	
The scaling tip doesn't vibrate when stepping on the foot switch.	Handpice connect with the cable irrelevantly. (I 02)	Connect it again.	
	Scaling tip is loose.	Fasten it by the wrench.	
when stepping on the footswitch.	The handpiece socket is too wet. (I 02)	Dry it with hot wind.	
	The handpiece is faulty.	Send it back to the manufacturer or local agent dealer for repair.	
	Water supply of dental unit is off.	Examine the water supply of dental unit.	
The scaling tip vibrates, but no water flowing out.	The liquid pipe in the cable is blocked	Clean the liquid pipe with the multi-function syringe.	
	The liquid way in the handpiece is blocked.	Clean the handpiece with the multi-function syringe.	
The handpiece overheat.	Water shortage.	Turn up the water.	
	The liquid pipe of dental unit is blocked.	Clean the water pipe.	
The water shortage.	The liquid pipe in the cable is blocked.	Clean the liquid pipe with the multi-function syringe.	
	The liquid way in the handpiece is blocked.	Clean the handpiece with the multi-function syringe.	
	Insufficient water pressure.	Enhance the water pressure.	
The vibration becomes weak.	The tip loose.	Fasten the tip.	
	The tip is faulty or worn down.	Change a new tip.	
The potentiometer is out of order.	The potentiometer is faulty.	Change a new potentiometer.	
Water leaks from the coupling between the handpiece and cable.	The water proof O-ring is damaged.	Change two new water proof O-ring.	

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# 5. ACCESSORIES

ITEMS	NAME	SPECIFICATIONS	
1	Control Cuicirt	64mm×75mm×35mm	
2	Detachable handpiece (I 02)		
3	Sealed handpiece (I 01)		
4	Cable (I 02)	BL - 1	
5	Potentiometer and flat cable.	2 ΚΩ	
6	Scaling tips	5 pcs packed in 1 package	
7	Handpiece ruble ring	17mm×15.4mm×2mm	
8	Water proof O-rings	3.5mm×1.5mm	
9	3-way / straight value		
10	Tip wrench	31mm×30mm	
11	Liquid pipe		
12	Power adaptor (optional)	30 V	
13	Electromagnetic valve (optional)		
14	Gas control switch (optional)		
15	Water regulating value (optional)		
16	6-way value (optional)		
17	Normally close value		

**NOTE:** The exactly specifications of accessories are not been shown detailedly in this operation instrument. See the attached page and packing list for more information.

# 6. NOTES

#### **6.1 NOTES OF USAGE**

- Keep the device clean before and after operation.
- The handpiece, scaling tips and wrench must be sterilized before each use.
- Do not install or uninstall a scaling tip when stepping on the foot switch.
- •The scaling tip must be fastened and there must be spray or drip coming out from the tip when operating.
- Change a new tip when it is damaged or worn down. Do not twist or rub the tip.
- Do not use impure water source and be sure not use normal brine instead of pure water source
- Be sure the tail of handpiece and the socket are dry before intalling the handpiece (I 02).
- Do not pull the cable forcibly while operating in order to avoid the handpiece fall off from the socket (I 02).
- •The threads of scaling tips which are produced by some other manufacturer might be coarse, rusty, collapsed or with different specification. They do not match the thread of our handpiece and will damage its thread beyond repair if force screwing down. Please always use the recommending scaling tips produced by Guilin Veirun Medical Technology Co., Ltd..
- In order to avoid affecting the function of scaler, it is not allowed to dismantle the scaler. If you have any special requirement, please contact with us.

#### **6.2 CONTRAINDICATION**

- Hemophiliacs disable.
- Patients or dentists with cardiac pacemaker disable.
- Heart disease patients, pregnant women and infants should be cautious.

#### **6.3 STORAGE AND MAINTENANCE**

- •This device should be handle with care, kept away from vibration and kept or installed in a cool, dry and ventilated place.
- Do not store the device along with any poisonous, mordant, flammable and explosive article.
- Storage humidity: ≤ 90%, storage temperature: -20°C~+55°C.

• Please turn off the power if not use it. If not use for a long time, please turn the power and water on for 5 minutes once per month.

#### 6.4 TRANSPORTATION

- Shock and vibration should be avoid in transport. Handle it with care and avoid to place upside down.
- Do not pack it with any hazardous articles in transport.
- Avoid the sun, rain or snow in transport.

# 7. AFTER SERVICE

- Guilin Veirun Medical Technology Co., Ltd. is responsible for the safrty of the products only in the following two kinds of situations:
- ① The maintaining, repairing and alterations were treated by our company or approved dealers.
- ② The replaced spare parts are original from Guilin Veirun Medical Technology Co., Ltd. and following the instructions to operate.
- Irreparable damage to the equipment caused by non-professional person dose not belong to free warranty.

# 8. DISPOSAL

This equipment dose not contain any poisonous and harmful substance. It can be disposed and destroyed according to the provisions of local authorities.

# 9 WARNING

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.

# 10. EMC Declaration

Table 1

Guidance	and manufacturer's d	eclaration - electromagnetic emissions	
The [K08] is intended for use in the electromagnetic environment specified below. The customer or the user of the [K08] 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

			onment specified below. The customer used in such an environment
Immunity Test	IEC 60601 Test level	Compliance level	Electromagnetic environment - guidance
Electrostatic discharge (ESD) IEC 61000-4-2	±6 kV contact ±8 kV air	±6 kV contact ±8 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 transient/burst IEC 61000-4-4	±2 kV for power supply lines ±1 kV for input/output lines	±2 kV for power supply lines ±1 kV for input/output lines	Mains power quality should be that of a typical commercial or hospital environment.
Surge IEC 61000-4-5	±1 kV line(s) to line(s) ±2 kV line(s) to earth	±1 kV line(s) to line(s) ±2 kV line(s) to earth	Mains power quality should be that of a typical commercial or hospital environment.

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#### Voltage dips, short <5% U<sub>T</sub> <5% U<sub>T</sub> Mains power quality should be that of a typical commercial or hospital interruptions and (>95% dip in U<sub>T</sub>) (>95% dip in U<sub>T</sub>) voltage variations environment. If the user of the [K08] for 0.5 cycle for 0.5 cycle requires continued operation during on power supply input lines 40% U<sub>T</sub> 40% U<sub>T</sub> power mains interruptions, it is IEC 61000-4-11 (60% dip in U<sub>T</sub>) (60% dip in U<sub>T</sub>) recommended the [K08] be for 5 cycle for 5 cycle powered from an uninterruptible power supply or a battery. 70% U<sub>T</sub> 70% U<sub>T</sub> (30% dip in U<sub>T</sub>) $(30\% \text{ dip in } U_T)$ for 25 cycle for 25 cycle <5% U<sub>T</sub> <5% U<sub>T</sub> (>95% dip in U<sub>T</sub>) (>95% dip in U<sub>T</sub>) for 5s for 5s 3 A/m Power frequency 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<sub>T</sub> is the a.c. mians voltage prior to application of the test level.

#### Table3

The [K08] is intended for use in the electromagnetic environment specified below. The customer or the user of the [K08] should assure that it is used in such an environment					
Immunity IEC 60601 Test Test level		Compliance level	Electromagnetic environment - guidance		
			Portable and mobile RF communications equipment should be used no closer to any part of the [K08], including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter.  Recommended separation distance		
Conducted RF IEC 61000-4-6	3Vrms 150KHz to 80MHz	3V	$d = 12\sqrt{P}$		
Radiated RF IEC 61000-4-3	3 V/m 80MHz to 2.5GHz	3 V/m	$d = 1.2\sqrt{P}$ 80 MHz to 800 MHz $d = 2.3\sqrt{P}$ 600 MHz to 2.5 GHz		
			where <i>p</i> is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and <i>d</i> is the recommended separation distance in metres (m)  Field strengths from fixed RF transmitters, as determined by		

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



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.

<sup>a</sup> 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].

<sup>b</sup> 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.

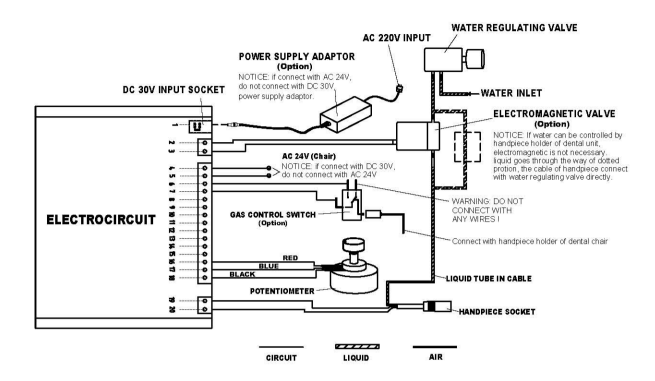
Rated maximum output power of transmitter	Separation distance according to frequency of transmitter m			
W	150 kHz to 80 MHz $d = 12\sqrt{P}$	80MHz to 800MHz $d = 1,2\sqrt{P}$	800MHz to 2.5GHz $d = 2.3\sqrt{P}$	
0,01	0.12	0.12	0.23	
0,1	0.38	0.38	0.73	
1	1.2	1.2	2.3	
10	3.8	3.8	7.3	
100	12	12	23	

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.

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# VRN BUILT-IN UNLTRASONIC SCALER CONNECTION INSTRUCTION



# VRN BUILT-IN UNLTRASONIC SCALER CONNECTION INSTRUCTION

