Pulse Oximeter JPD-500G(Bluetooth)

FCC statement

This device complies with Part 15 of the FCC Rules. Operation is subject to the following two conditions :

(1) This device may not cause harmful interference.

(2) This device must accept any interference received, including interference that may cause undesired operation.

- NOTE: This equipment has been tested and found to comply with the limits for a Class B digital device, pursuant to Part 15 of the FCC Rules. These limits are designed to provide reasonable protection against harmful interference in a residential installation. This equipment generates, uses and can radiate radio frequency energy and, if not installed and used in accordance with the instructions, may cause harmful interference to radio communications. However, there is no guarantee that interference will not occur in a particular installation. If this equipment does cause harmful interference to radio or television reception, which can be determined by turning the equipment off and on, the ser is encouraged to try to correct the interference by one or more of the following measures:
 - Reorient or relocate the receiving antenna.
 - Increase the separation between the equipment and receiver.
 - •Connect the equipment into an outlet on a circuit different from that to which the receiver is connected.

•Consult the dealer or an experienced radio/TV technician for help. **Warning:** Changes or modifications to this unit not expressly approved by the party responsible for compliance could void the user's authority to operate the equipment.

Precautions

• Do not attempt to repair he Oximeter unless you are professional engineers. Only professionals with maintenance qualification are allowed to perform interior maintenance as necessary.

• Change the contact position between the Oximeter probe and the finger periodically if you are monitoring your SpO2 levels and pulse rate for more than 2 hours.

• Stop immediately if you have broken skin or the blood circulation of your finger is affected during prolong use.

• This product is not designed to be used by newborn babies.

Seek for medical care if the measured value goes beyond the normal range and you are sure that the instrument is not malfunctioning.

• The pulse oximeter uses infrared light (invisible to your eyes) to measure your SpO2 levels. Hence, please do not stare at the light-emitting components of the Oximeter, as that could cause harm and/or potentially blind your eyes.

 This pulse oximeter is not a medical device and is not intended to diagnose or treat any medical condition or disease. It is intended for non-medical use in healthy people to monitor their pulse and blood oxygen levels during sports and/or aviation only.

People who need SpO2 and pulse rate measurements because of a medical condition should not use the oximeter and should consult with their physician. • For details about clinical limitations and contraindications, please carefully consult relevant medical literatures.

The following factors may affect the accuracy of the measurement:

• The Oximeter is used in an environment involving high-frequency devices, such as high-frequency electric knives and CT apparatuses.

• Ambient light intensity is too bright. Hence, please avoid direct exposure to strong light (such as beams from operating lamps or sunlight) during measurement.

- The probe of the Oximeter is placed on the same arm that a blood pressure cuff arterial duct or intravenous injection.
- The user suffers from hypotension, severe vascular atrophy, severe anemia, or low oxygen.
- The user is in sudden cardiac arrest or shock state.
- The user is wearing nail polish or artificial nails.

Warnings

Warning: Do not use the Oximeter in an environment with any flammable gases, flammable anesthetic, or other flammable substances.

Warning: Keep unit and lanyard away from children as the included lanyard may present an entanglement or choking hazard to small children. Adult supervision required; never leave children unattended with unit or lanyard

Warning: Do not throw the batteries into fire, as that could cause an explosion. Warning: Do not attempt to charge the included batteries, as that could cause leakage, fire disaster, or even explosion. Dispose the used batteries in accordance to the local laws and regulations.

Warning: Do not use the Oximeter in an MRI or CT environment.

Warning: Caution: Do not operate the Oximeter if it is wet. Avoid moving the oximeter from a cold to a hot and humid environment.

Warning: Install the batteries properly before powering on the Oximeter for normal use. Please remove the batteries if you are not planning to use the Oximeter for a long time.

Warning: Close the battery cover when the instrument is in use.

Symbols

Meaning
Type BF applied part
Caution: Please see this manual.
Symbol of oxygen saturation
Symbol of pulse rate
No SpO2 alarms.
Bluetooth
When end users abandon this product, they must send the product to the collection place for recycling.

Overview

Oxygen saturation is the percentage of oxyhemoglobin (HbO2) that is combined

with oxygen against all combinable hemoglobin (Hb). It is an important physiological

parameter involved in respiration and circulation. The oxygen saturation of arterial

blood in a normal human body is 98%. Oxygen saturation is an important indicator of the oxygen condition in the human body. In general, the normal values of oxygen saturation shall not be lower than 94%. If the measured value of oxygen saturation is lower than 94%, an insufficient supply of oxygen is considered.

The pulse rate is the number of pulse beats per minute. Normally, the pulse rate is consistent with the heart rate. In general, the pulse rate of every people is 60 to 90 beats per minute.

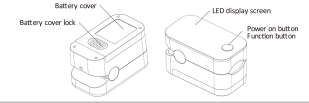
The Perfusion Index (PI) usually reflects the limb perfusion status of an examined patient, and shows the detection precision of the instrument as well; that is, examination can still be performed even in the low or weak perfusion condition.The PI of a normal human body is 3% or greater.

Working Principles and Usage

Based on full digital technology, the Finger Pulse Oximeter non-invasively measures the actual content (oxygen saturation) of oxyhemoglobin (HbO2) in arterial blood using the optical transmittance method.

The Finger Pulse Oximeter measures the blood oxygen saturation and pulse rate of a human body via finger artery. It is applicable to a wide range of fields, such as families, clinics, oxygen bars, social medical care institutions, and sports & health. Use this instrument for measurement before or after sports. You are not advised to use this instrument during sports activities. Do not use it for continuous care for patients.

Schematic Diagram of Display



Schematic Diagram of Display

The following figure shows the information display on the LED screen of the Oximeter in normal detection state:



Hold: Press the button for more than 0.5 seconds.

Brightness Setting

Hold the power-on button while the oximeter is in powered-on state, then the oximeter shows a brightness setting interface(as "Interface 1" below shows, "br" represents brightness). Hold the button to adjust brightness. There 3 brightness settings(1,2,3). 3 is the brightest.



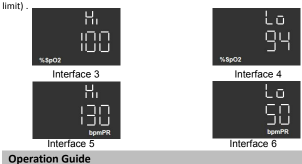
Alarm Setting

After setting the brightness, press the power-on button to enter the alarm setting interface(as "interface 2" below shows, "AL" represents alarm). Then hold the button to set "AL" to on or off. When "AL" is set to on and the measured values of the blood oxygen saturation and pulse rate go beyond the upper limit or lower limit, the oximeter will beep to alarm.



Alert Range Setting

When "AL" is set to on, you can set the upper limit and lower limit of SpO2 Alert and PR Alert. Press it to switch an option(SpO2 upper limit, SpO2 lower limit, PR upper limit and PR lower limit). Hold the power-on button to adjust the limits. (as "Interface 3,4,5,6" below show, "Hi" represents upper limit, "Lo" represents lower



Stick one finger completely into the finger chamber

of the oximeter. The fingernail should be facing upward.

Release the clip and press the power-on button to power

on the pulse oximeter.

 \angle If you do not insert your finger completely into the

chamber, measurement will be inaccurate.

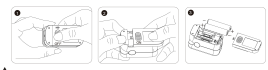
To keep your finger still during measurement. It is also not advisable to use this instrument during sports activities as movement may lead to inaccuracies. Once the reading stabilizes, read the measured values of oxygen saturation and pulse rate on the screen.

NOTE: The oximeter will automatically shut down 10 seconds after you remove your

finger.

Connecting the Instrument to a smartphone via Bluetooth

Note: For details on specific operations, see the JUMPER Health User Manual.



 Δ Replace the batteries when the batteries run out of power and the symbol

 (\Box) flickers on the screen.

Install the two AAA dry batteries into the battery slot according to polarity indication, and mount the battery cover.

Cleaning

Power off the instrument and remove the batteries before cleaning. Ensure that the appearance of the instrument is neat, dust-free, and dirt-free. Clean the outer surface of the instrument (including the LED screen) using a piece of dry soft cloth dipped with 75% medical alcohol

Caution: Avoid liquid flowing into the instrument during cleaning. Caution: Do not immerse any part of the instrument into any liquid.

Disinfection

Before measurement with the instrument, wipe the rubber finger pad using a piece of dry soft cloth dipped with 75% medical alcohol. Clean the finger to be measured using the medical alcohol for disinfection purposes before and after use.

Do not disinfect the instrument by means of high-temperature/high-

pressure or gas disinfection.

Maintenance

- Remove the batteries from the battery slot and properly store them if you do not plan to use the Oximeter for a long period of time.
- Avoid using the Oximeter in an environment with inflammable gases or using it in an environment where the temperature or humidity is excessively high or low.
- Check the accuracy of the oxygen saturation and pulse rate readings by using • an appropriate calibration apparatus.

Technical Specifications

- 1. Dimensions: 58.0 mm (Width) × 32.0 mm (Depth) × 32.9 mm (Height) Weight: 50.4 g (including two AAA dry batteries)
- 2. Peak wavelength range of the light emitted from the probe: red light 663 nm ± 3; infrared light 900 nm ± 7.
- 3. Maximum optical output power of the probe: 60 mW for infrared light (905 nm).

Bluetooth module:4.2 4

5. Normal working condition

110								
	Working Temperature	5°C to 40°C (41°F to 104°F)						
	Relative Humidity	15% to 80%, non-condensing						
	Atmospheric Pressure	70 kPa to 106 kPa						
	Rated Voltage	DC 3.0 V						
	Atmospheric Pressure	70 kPa to 106 kPa						

Default values and conditions of alert

	Parameter	Value				
	Our contraction	Upper limit: 99				
	Oxygen saturation	Lower limit: 94				
	Pulse rate	Upper limit: 130				
	Puise rate	Lower limit: 50				
	Alert condition	When the alert switch is on and the				
		actual measured value goes beyond				
		the preset alert parameter range, the				
		Oximeter gives an alert sound.				

7. Technical parameters

eci	echnical parameters							
	Param	eter	Value					
	Display range	Oxygen saturation	35% to 99%					
		Pulse rate	35 bpm to 250 bpm					
	Resolution	Oxygen saturation	1%					
		Pulse rate	1 bpm					
	Measurement precision	Oxygen	±2% (70% to 99%)					
		saturation	No requirement (≤ 69%)					
		Pulse rate	±2 bpm					
		Oxygen	Upper limit: 50% to 100%					
		saturation	Lower limit: 50% to 100%					
	Alert range	Pulse rate	Upper limit: 35 bpm to 250 bpm Lower limit: 35 bpm to 250 bpm					
		Oxygen saturation	± 1% of the preset value					
	Alert error	Pulse rate	The greater of ±10% of the preset value and ±5 bpm					
	PI	Weak PI	Min. 0.3%					

Safety Type

Anti-electric-shock type: internal power supply device Anti-electric-shock degree: Type BF applied part Running mode: continuous working Waterproof grade: IP22

Storage and Transportation

Temperature : $-10^{\circ}C - 50^{\circ}C(14^{\circ}F - 122^{\circ}F)$ Relative humidity : 10%-93% (no condensation) Atmospheric pressure : 50kPa-106 kPa.

ELECTROMAGNETIC COMPATIBILITY (EMC) TABLES

Guidance and manufacturer's declaration - electromagnetic emissions							
The device is intended for use in the electromagnetic environment specified below.							
The customer or the user assure that it is used in such an environment.							
Emissions test	Compliance	Electromagnetic environment - guidance					
RF emissions CISPR11	Group 1	The devuce device use RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.					
RF emissions CISPR11	Class B	The device is suitable for use in all establishments other than domestic and					



Harmonic emissions	Not applica		ectly connected to the public ge power supply network that		in U for 5	T) 5 sec				•	nobile radios, amateu t be predicted theore		
IEC61000-3-2 Voltage fluctuations/ Ficker emissionsIEC61000 3-3	Not applica		supplies buildings used for domestic		Power frequency (50Hz/60Hz) 3A/m 3A/m Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.				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 Blood Pressure Monitor is used exceeds the applicable RF compliance level above, the Blood Pressure Monitor should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as reorienting or relocating the Blood Pressure Monitor.				
	ufo oturo r'o do olo			NOTE UT is	the a.c. mai	ins voltage	prior to ap	plication of the test level.	41	-	80 MHz, field strengt		
Guidance and man			environment specified below.	Guidance and	l manufactu	ire's declara	ation – eleo	ctromagnetic immunity]	paration distances bet	ween portable and m	obile RF	
The customer or the user of the device should assure that it is used in such an environment.			The device is intended for use in the electromagnetic environment specified below. The customer or the user of device should assure that it is used in such an				•	communications equipment and the device. The device is intended for use in an electromagnetic environment in which radiated					
Immunity test	IEC 60601 Test level	Compliance level	Electromagnetic environment-guidance	environment.	IEC				41		omer or the user of the maintaining a minim	e device can help um distance between	
Electrostatic discharge (ESD)	±6kV contact	±6kV contact ±8kV air	Floors should be wood , concrete or ceramic tile. If	Immunity test	60601 test	Compliar level		Electromagnetic environment - guidance	11 .	ters) and the device			
IEC61000-4-2	±8kV air		floors are covered with synthetic material, the relative humidity should be	Radiated RF IEC	level 3 V/m 80 MHz	3 V/m		Portable and mobile RF communications equipment should		Separation distance according to frequency of transmitte (m)			
Electrical fast transient/burst IEC 61000-4-4	±2 kV for power supply lines	not applicable	at least 30% not applicable (For INTERNALLY POWERED ME EQUIPMENT)	61000-4-3	to 2.5 GHz			be used no closer to any part of the Blood Pressure Monitor, including cables, than the recommended separation distance calculated from	Maximum output power of transmitter	80 MHz to 800	800 MHz to 2.5	800 MHz to 2.5	
120 01000-4-4	± 1 kV Input/ output line						1	the equation applicable to the frequency of the transmitter. Recommended separation distance	(W)	$d = 1.2\sqrt{P}$	GHz $d = 1.2\sqrt{P}$	GHz $d = 1.2\sqrt{P}$	
Surge IEC 61000-4-5	± 1 kV Differential mode	not applicable	not applicable (For INTERNALLY POWERED ME EQUIPMENT)					$d = 1.2\sqrt{P}$ $d = 1.2\sqrt{P}$ 80 MHz to 800 MHz	0.01	/	0.12	0.23	
	voltage ± 2 kV							$d = 1.2\sqrt{P}$ 80 MHz to 800 MHz $d = 1.2\sqrt{P}$ 800 MHz to 2.5 GHz	1 10	/	1.2 3.8	2.3 7.3	
	Common mode							Where P is the maximum output power rating of the transmitter in watts (W) according to the	100 For transmitters ra		12 tput power not listed	23 above, There	
Voltage dips, short interruptions and voltage variations on	voltage <5% UT (>95% dip in UT) for 0.5 cycle	not applicable	not applicable (For INTERNALLY POWERED ME EQUIPMENT	watts (w) according to the transmitter manufacturer and d is the recommended separation distance in metres (m). Field strengths from fixed RF transmitters, as determined by an				 commended separation distance d in meters(m) can be estimated using the equation applicable to the frequency of the transmitter, where P is the maximum output power rating of the transmitter in watts(W) accordable to the transmitter manufacturer. NOTEI At 80 MHz and 800 MHz. the separation distance for the higher frequency range applies. 					
power supply input lines IEC 61000-4-11	40% UT (60% dip in UT) for 5 cycles					be less each fre Interfer of equi	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:	-	uidelines may not apply in all situations. Electromagnetic affected by absorption and refection from structures, objects and				
	70% UT (30% dip in							((•))	EC REP Wellkang Ltd Suite B, 29Harley Street, LONDON, W1G9QR,U.K.			J.K.	
	UT) for 25 cycles <5% UT			NOTE 2 T propagation i people.	hese guidel s affected b	ines may no y absorptio	ot apply in on and refle	· •	Shenzhen Jumper Medical Equipment Co., Ltd Address: D Building, No. 71, Xintian Road, Fuyong Street, Baoan,Shenzhen, Guangdong,China E-mail: info@jumper-medical.com Tel: +86-755-26692192,26696279 Web:www.jumper-medical.com				
	<pre>>>>>>>>>>>>>>>>>>>>>>>>>>>>>>>>>>>></pre>					from fixed transmitters, such as base stations for radio			Tel: +86-755-26692192 26696279				