Pulse Oximeter

User Manual



Instructions to User

Dear users, thank you very much for purchasing the Pulse Oximeter

This Manual is written and compiled in accordance with the council directive MDD93/42/EEC for medical devices and harmonized standards. In case of modifications and software upgrades, the information contained in this document is subject to change without notice.

The Manual describes, in accordance with the Pulse Oximeter's features and requirements, main structure, functions, specifications, correct methods for transportation, installation, usage, operation, repair, maintenance and storage, etc. as well as the safety procedures to protect both the user and equipment. Refer to the respective chapters for details

Please read the User Manual carefully before using this product. The User Manual which describes the operating procedures should be followed strictly.Failure to follow the User Manual may cause measuring abnormality, equipment damage and human injury. The manufacturer is NOT responsible

for the safety, reliability and performance issues and any monitoring abnormality, human injury and equipment damage due to users' negligence of the operation instructions. The manufacturer's warranty service does not cover such faults.

Owing to the forthcoming renovation, the specific products you received may not be totally in accordance with the description of this User Manual. We would sincerely regret for that.

This product is medical device, which can be used repeatedly.

WARNING:

- Uncomfortable or painful feeling may appear if using the device ceaselessly, especially for the microcirculation barrier patients. It is recommended that the sensor should not be applied to the same finger for over 2 hours.
- For the special patients, there should be a more prudent inspecting in the placing process. The device can not be clipped on the cdema and tender tissue.

- The light (the infrared is invisible) emitted from the device is harmful to the eyes, so the user and the maintenance man should not stare at the light.
- **●** Testee can not use enamel or other makeup.
- Testee's fingernail can not be too long.
- Please refer to the correlative literature about the clinical restrictions and caution.
- This device is not intended for treatment.

The User Manual is published by our company. All rights reserved.

CONTENTS

C - C - 4-

1	Saie	ty	I
	1.1	Instructions for safe operations	1
	1.2	Warning	1
	1.3	Attention	2
2	Over	rview	4
	2.1	Features	5
	2.2	Major applications and scope of	
	appl	lication	5
	2.3		
3	Prin	ciple	
4	Tech	nical specifications	7
	4.1	Main performance	7
	4.2	Main Parameters	7
5	Insta	allation	8
	5.1	View of the front panel	8
	5.2		
	5.3	Accessories	9
6	Oper	rating Guide	9
	6.1	Application method	9
	6.2	Attention for operation	14
	6.3	Clinical restrictions	15
7	Mair	ntain、transportation and storage	16
	7.1	Cleaning and Disinfecting	16
	7.2	Maintain	16
	7.3	Transportation and storage	
8	Trou	bleshooting	17
9	Key	of Symbols	19
10		tion Specification	

1 Safety

1.1 Instructions for safe operations

- Check the main unit and all accessories periodically to make sure that there is no visible damage that may affect patient's safety and monitoring performance about cables and transducers. It is recommended that the device should be inspected once a week at least. When there is obvious damage, stop using the device.
- Necessary maintenance must be performed by qualified service engineers ONLY. Users are not permitted to maintain it by themselves.
- The oximeter cannot be used together with devices not specified in User's Manual.Only the accessory that appointed or recommendatory by manufacture can be used with this device.
- This product is calibrated before leaving factory.

1.2 Warning

- Explosive hazard—DO NOT use the oximeter in environment with inflammable gas such as some ignitable anesthetic agents.
- ♠™ DO NOT use the oximeter while the testee measured by MRI and CT.
- The person who is allergic to rubber can not use this device

- ◆ The disposal of scrap instrument and its accessories and packings(including battery, plastic bags, foams and paper boxes) should follow the local laws and regulations.
- ♠™ Please check the packing before use to make sure the device and accessories are totally in accordance with the packing list, or else the device may have the possibility of working abnormally.
- Please choose the accessories and probe which are approved or manufactured by the manufacturer, or else it may damage the device.
- Please don't measure this device with functional tester for the device's related information

1.3 Attention

- ⊕ Keep the oximeter away from dust, vibration, corrosive substances, explosive materials, high temperature and moisture.
- $\mathrel{\ \mathclap{\oplus}\ }$ If the oximeter gets wet, please stop operating it.
- A When it is carried from cold environment to warm or humid environment, please do not use it immediately.
- ⊕ DO NOT operate keys on front panel with sharp materials
- A High temperature or high pressure steam disinfection of the oximeter is not permitted. Refer to User Manual in the relative chapter (7.1) for

instructions of cleaning and disinfection.

- Do not have the oximeter immerged in liquid. When it needs cleaning, please wipe its surface with medical alcohol by soft material. Do not spray any liquid on the device directly.
- $\ensuremath{\triangle}$ When cleaning the device with water, the temperature should be lower than $60\,^{\circ}\text{C}$.
- $\ensuremath{\mathfrak{S}}$ As to the fingers which are too thin or too cold, it would probably affect the normal measure of the patients' SpO₂ and pulse rate, please clip the thick finger such as thumb and middle finger deeply enought into the probe.
- ⊕ The update period of data is less than 5 seconds, which is changeable according to different individual pulse rate.
- ⊕ Please read the measured value when the waveform on screen is equably and steady-going, This measured value is optimal value. And the waveform at the monment is the standard one.
- A If some abnormal conditions appear on the screen during test process, pull out the finger and reinsert to restore normal use
- The device has normal useful life for three years since the first electrified use
- The device may not work for all patients. If you
 are unable to achieve stable readings, discontinue

use

2 Overview

The pulse oxygen saturation is the percentage of HbO2 in the total Hb in the blood, so-called the O2 concentration in the blood. It is an important bio-parameter for the respiration. A number of diseases relating to respiratory system may cause the decrease of SpO2 in the blood, furthermore, some other causes such as the malfunction of human body's self-adjustment, damages during surgery, and the injuries caused by some medical checkup would also lead to the difficulty of oxygen supply in human body, and the corresponding symptoms would appear as a consequence, such as vertigo, impotence, vomit etc. Serious symptoms might bring danger to human's life. Therefore, prompt information of patients' SpO2 is of great help for the doctor to discover the potential danger, and is of great importance in the clinical medical field.

The Pulse Oximeter features in small volume, low power consumption, convenient operation and being portable. It is only necessary for patients to put one of his fingers into a probe for diagnosis, and a display screen will directly show the measured value of pulse oxygen saturation with the high veracity and repetition.

2.1 Features

A. Operation of the product is simple and convenient.

B. The product is small in volume, light in weight and convenient in carrying.

C. Low power consumption

2.2 Major applications and scope of application

The Pulse Oximeter can be used in measuring the pulse oxygen saturation and pulse rate through finger. The product is suitable for being used in family, hospital, oxygen bar, community healthcare, physical care in sports (It can be used before or after doing sports, and it is not recommended to use the device during the process of having sport) and etc.

⚠ The problem of overrating would emerge when the patient is suffering from toxicosis which caused by carbon monoxide, the device is not recommended to be used under this circumstance.

2.3 Environment requirements

Storage Environment

- a) Temperature :-40°C ~+60°C
- b) Relative humidity :5%~95%
- c) Atmospheric pressure :500hPa~1060hPa

Operating Environment

a) Temperature:10°C ~40°C

- b) Relative Humidity :30%~75%
- c) Atmospheric pressure: 700hPa~1060hPa

3 Principle

Principle of the Oximeter is as follows: An experience formula of data process is established taking use of Lambert Beer Law according to Spectrum Absorption Characteristics of Reductive Hemoglobin (Hb) and Oxyhemoglobin (HbO2) in glow & near-infrared zones. Operation principle of the device is: Photoelectric Oxyhemoglobin Inspection Technology is adopted in accordance with Capacity Pulse Scanning & Recording Technology. so that two beams of different wavelength of lights can be focused onto human nail tip through perspective clamp finger-type sensor. Then measured signal can be obtained by a photosensitive element, information acquired through which will be shown on screen through treatment in electronic circuits and microprocessor.

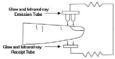


Figure 1.

4 Technical specifications

4.1 Main performance

- A. SpO₂ value display
- B. Pulse rate value display, bar graph display
- C. Pulse waveform display
- D. Low-voltage indication: low-voltage indicator appears before working abnormally which is due to low-voltage
- E. The display mode can be changed
- **F.** With SpO₂ value and pulse rate value of storage, the stored data can be uploaded to computers

4.2 Main Parameters

A. Measurement of SpO₂

Measuring range: 0%~100%

Accuracy:

When the SpO₂ measuring range is 70% \sim 100%,the permission of absolute error is $\pm 2\%$;

below 70% unspecified

B. Measurement of pulse rate

Measuring range:30bpm~250bpm

Accuracy: ± 2 bpm or $\pm 2\%$ (select larger)

C. Resolution

SpO₂: 1%, Pulse rate: 1bpm.

D. Measurement Performance in Weak Filling Condition

 SpO_2 and pulse rate can be shown correctly when pulse-filling ratio is 0.4%. SpO_2 error is \pm 4%, pulse rate error is \pm 2 bpm or \pm 2% (select larger).

E. Resistance to surrounding light

The deviation between the value measured in the condition of man-made light or indoor natural light and that of darkroom is less than ±1%.

F. Power supply requirement: $: 2.6 \text{ V DC} \sim 3.6 \text{V}$ DC

G. Optical Sensor

Red light (wavelength is 660nm, 6.65mW) Infrared (wavelength is 880nm, 6.75mW)

5 Installation

5.1 View of the front panel



Figure 2. front view

5.2 Battery installation

A. Refer to Figure 3. and insert the two AAA size batteries properly in the right direction.

B. Replace the cover.

A Please take care when you insert the batteries for the improper insertion may damage the device.



Figure 3.

5.3 Accessories

- A. One hanging rope
- B. Two dry batteries(AAA)
- C. A User Manual
- D. a data line
- E. a disk (PC software)

6 Operating Guide

6.1 Application method

Δ

- a) Insert the two batteries properly to the direction, and then replace the cover.
- b) Open the clip as shown in Figure 4.
- c) Let the patient's finger put into the rubber

۲.

.

cushions of the clip (make sure the finger is in the right position), and then clip the finger.

- d) Do not shake the finger and keep the patient in a stable state during the process.
- e) The data can be read directly from the screen on the measuring interface.

Fingernails and the luminescent tube should be on the same side.



Figure 4.
Put finger in position

B. Change display direction

On the measuring interface, short press the button to change the display.

C. Real-time data transmission setting

Firstly, please install the affiliated software into the computer, and two icons would appear on the desktop after installation. The icon of "SpO2 "is a program for receiving real-time data which is shown as figure 5; the icon of "SpO2 Review" is a program for receiving stored data which is shown as figure 6.

- a. Please connect the device to computer with the affiliated data line, then double click the "SpO2 "icon to start the program.
- b. When you unplug the data line from computer, there is a dialog box "Save data at view" appearing on the desktop, in which you can input some patient's basic information.



Figure 5. SpO₂ program



Figure 6. SpO2 Review program

⚠ If the users choose to turn on the synchronizing display function on computer, it would probably take several seconds for the data to appear on the computer screen.

D. Menu operations

When the device is under the measuring interface, Press the button for about 1 second in order to enter the menu interface shown as figure 7. Users can adjust the setting through the main menu, such as data storage. The specific operation methods are as the following:



Figure 7. Main Menu Interface

a) Data storage setting

This instrument has the ability to store 24 hours worth of data. It can store the measured pulse rate and SpO₂ value accurately, transfer the data to the computer, display the data and print reports (with the included SpO₂ Software - Green Heart)

- a. On the main menu interface, short press the button to select "Record", then long press the button to choose whether store the data or not, choose "on" to permit storing, choose "off" to forbid storing.
- b. If the data storage function is being turned on, when return to the measuring interface, a flashing yellow dot would appear on screen, which means the device is in a state of storing.
- c. In the state of storing, the device is on measuring interface, the sign "Recording" would

appear on the screen in 30 seconds, and then the screen will be automatically shut down, only a flashing yellow dot appear on screen. If short press the button at this moment, the sign "Recording" would appear on the screen, and then the screen will be automatically shut down again; if long press the button, the device would return to the measuring interface

- d. If turning on the data storage function, the former data storage will be automatically removed.
- e. In the state of data storing, after the screen is automatically shut down, the pulse sound indication would be off for saving power.
- f. When the storage space is full, it displays "Memory is full" on the screen, and then shut down in a few seconds. But it will still display "Memory is full" by the next time you turn on the device on the purpose of warning the user, and a few seconds later enter the measuring interface.

b) Uploading the data to the PC after recording

a. Please connect the device with computer by the data line which is equipped with the device, then double click "SpO2 Review" icon to open"SpO2 Review" program, click the 'New Session' Icon in the software, enter the patient data and then click 'ok'. The Software will then display "device

connected, waiting for data".

- b. on the main menu interface, the users to upload the stored date to computer when the symbol "Record" shows "off".
- c. In the state of storing, it is not applicable for the users to upload the stored date to computer.

c) Exit the main menu

On the main menu interface, Click the button to select "Exit", then long press the button to exit the main menu

E. Power off

If the measuring finger out of the device on the measuring interface, The device will power off automatically when it gets no signal within 5 seconds. The device can't being power off when it is in a state of storing.

6.2 Attention for operation

- A. Please check the device before using, and confirm that it can work normally.
- **B.** The finger should be in a proper position (see the attached illustration of figure 4 for reference), or else it may result in inaccurate measure.
- C. The SpO₂ sensor and photoelectric receiving tube should be arranged in a way with the subject's arteriole in a position there between.
- D. The SpO₂ sensor should not be used at a location

- or limb tied with arterial canal or blood pressure cuff or receiving intravenous injection.
- E. Do not fix the SpO₂ sensor with adhesive or else it may result in venous pulsation and inaccurate measure of SpO₂ and pulse rate.
- F. Excessive ambient light may affect the measuring result. It includes fluorescent lamp, dual ruby light, infrared heater, direct sunlight and etc.
- G. Strenuous action of the subject or extreme electrosurgical interference may also affect the accuracy.
- H. Testee can not use enamel or other makeup.
- I. Please clean and disinfect the device after operating according to the User Manual (7.1).

6.3 Clinical restrictions

- A. As the measure is taken on the basis of arteriole pulse, substantial pulsating blood flow of subject is required. For a subject with weak pulse due to shock, low ambient/body temperature, major bleeding, or use of vascular contracting drug, the SpO₂ waveform (PLETH) will decrease. In this case, the measurement will be more sensitive to interference.
- B. For those with a substantial amount of staining dilution drug (such as methylene blue, indigo green and acid indigo blue), or carbon monoxide hemoglobin (COHb), or methionine (Me+Hb) or

thiosalicylic hemoglobin, and some with icterus problem, the SpO₂ determination by this monitor may be inaccurate.

 C_{\bullet} The drugs like dopamine, procaine, prilocaine, lidocaine and butacaine may also be a major factor blamed for serious error of SpO₂ measure.

D. As the SpO₂ value serves as a reference value for judgement of anemic anoxia and toxic anoxia, some patients with serious anemia may also report good SpO₂ measurement.

7 Maintain transportation and storage

7.1 Cleaning and Disinfecting

Using medical alcohol to disinfect the device, nature dry or clean it with clean soft cloth.

7.2 Maintain

A. Please clean and disinfect the device before using according to the User Manual (7.1).

B. Please change the battery when the screen shows \bigcap

C. Take out the battery if leave the equipment unused for long time.

D. The device needs to be calibrated once a year (or according to the calibrating program of hospital). It also can be performed at the state-appointed agent or just contact us for calibration.

7.3 Transportation and storage

A. The packed device can be transported by ordinary conveyance or according to transport contract. The device can not be transported mixed with toxic, harmful, corrosive material.

B. The packed device should be stored in room with no corrosive gases and good ventilation. Temperature: -40°C~60°C; Humidity: ≤95%

8 Troubleshooting

Trouble	Possible Reason	Solution
The SpO ₂ and Pulse Rate can not be displaye d normall y	The finger is not properly positioned. The patient's SpO ₂ is too low to be detected.	Place the finger properly and try again. Try again; Go to a hospital for a diagnosis if you are sure the device works all right.

The SpO ₂ and Pulse Rate are not displaye d stably	The finger is not placed inside deep enough. The finger is shaking or the patient is moving.	Place the finger properly and try again. Let the patient keep calm
The device can not be turned on	1. 1. The battery is drained away or almost drained away. 2. The battery is installed incorrectly. 3. The malfunction of the device.	1.Please change batteries. 2.Please Install the battery again. 3. Please contact the local service center.
The display is off suddenl y	The device's malfunction. The battery is drained away or almost drained away.	Please contact the local service center. Please change batteries.

9 Key of Symbols

Signal	Description
\triangle	Warning – See User Manual
%SpO ₂	The pulse oxygen saturation(%)
PRbpm	Pulse rate (bpm)
Û	Low-voltage
E-0-0	menu button/power button/function button
☀	Type BF
SN	Serial number
	the finger clip falls off (no finger inserted)] Probe error Signal inadequacy indicator

18 19

+	battery positive electrode
1	battery cathode
•	USB
IPX1	Ingress of liquids rank
Z	WEEE (2002/96/EC)

10 Function Specification

Information	Display Mode	
The Pulse Oxygen Saturation (SpO ₂)	2-digit digital OLED display	
Pulse Rate (PR)	3-digit digital OLED display	
Pulse Intensity (bar-graph)	bar-graph OLED display	
SpO ₂ Parameter Specification		
Measuring range	0%~100%, (the resolution is 1%).	
Accuracy	70%~100%:±2%,Below 70% unspecified.	

Average value	Calculate the Average value in every 4 measure value. The deviation between average value and true value does not exceed 1%.	
Pulse Parameter Specification		
Measuring range	30bpm~250bpm, (the resolution is 1bpm)	
Accuracy	±2bpm or±2% (select larger)	
Average pulse rate	Moving calculate the Average pulse rate every 4 cardio-beats cycle. The deviation between average value and true value does not exceed 1%	
Safety Type	Interior Battery,BF Type	

Pulse Intensity		
Range	Continuous bar-graph display, the higher display indicate the stronger pulse.	
Battery Requirement		
1.5V (AAA size) alkaline batteries × 2 or rechargeable battery		
Battery working life		
Two1.5V (AAA size)600mAh alkaline batteries can work continually for 24 hours		
Dimensions and Weight		

Dimensions	58.5(L) × 31(W) × 32 (H) mm
Weight	About 50g (with the batteries)