

Pulse Oximeter User Manual



Product Model:YM101/YM102/YM103

Date: 2020-07-08

YIMI Life (\(\xi_{0123} \)

Power indication PIX PIX PIX PRISON Index

Figure 1 Front View of YM101/YM102/YM103

1.2 Operation Method

- A. Open the battery cover, and put the two AAA batteries into the battery compartment in correct polarities, then B. Press the bottom of the equipment and open the
- probe, then insert one finger into the probe; . Press the button to turn the equipment on, and the measure interface will appear;
 D. After about 8 seconds, the measurement result can be read directly from the display screen:
- E. Before reading the parameters, make sure that stable numbers of the pulse oximeter interface has sustained more than 4 second; . The equipment will turned off automatically within 8

seconds when the finger left the probe.

WARNINGS:

use for a long time.

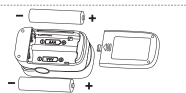


Figure 2 Battery Installation

A. Pass the thinner end of the lanvard through the hanging hole end and tighten the lanyard (Figure3).

1.3 Battery Installation

orrect polarities (Figure 2). B. Push the battery cover horizontally along the arrow hown as right.

· Battery polarities should be correctly installed, otherwise, damage may be caused to the equipment. Please remove the batteries if the equipment will not

1.4 Lanyard installation

- 1.5 Attention for Operation
- a) To avoid glare and direct sunlight exposure;
- ied with arterial canal or blood pressure cuff or receiving ntravenous injection;
- The ray between photo detector and light emitting

- size were applicable; B. Before use check and confirm that the environment

- A. Before use check and confirm that the people or finge hould be non-combustible material, as well as to avoid high or low temperature and humidity, but also need to
- b) To avoid radiation infrared or ultraviolet radiation;
 c) Avoid contact with the organic solvent, mist, dust,
- D. The equipment may not work normally on microcirculation barrier patients, Warm or rub the finge
- The patient should not use enamel or other makeup;

- corrosive gases; ... The equipment should not be used at a location or limb
- or re-position the equipment could improve the measurement
- Avoid to insert a wet finger into the probe.

should be noted that long-press means pressing the key for about 2 seconds, and short-press means pressing th A. The user should fully insert the finger into the probe. B. It is recommended to let the LED light shine directly 1.6.2 Menu Operation

Figure 4 Finger Placement Diagram

1.6 Functions and Menu Operation

item's submenu, confirming setting values, and

menu items and viewing the setting values of items. It

should be noted that long-press means pressing the key

1.6.1 The button operation rules

2. Don't shake the finger and try to keep the patient still After the oximeter is turned on, long-press the powe

outton to activate the menu, then short-press the button to view the setting values of each item. If the user wants to change the setting value of the item, long-press to enter the item's submenu, the parameter value starts to flash. parameter value required by the user is selected, long oress to confirm and exit the submenu.

tem1. Setup the LED display brightness The first item is to setup the display brightness. Long-1 to 3. The greater the value, the greater brightness of the



he second item is to setup the SpO2 alarm limits. For example: When Spo2 High limit is set to 96, an alarm will be issued when the spo2 value is higher than 96, and when

2.1 Classification

Degree of protection against electric shock:

Type BF-Applied part

Degree of protection against hazards of explosion: IP22 2.2 Power Requirements

Item4.Turn Alarm On/Off he forth item is long-press to turn Alarm on /off.

Spo2 low limit is set to 94, an alarm will be issued when the

he third item is to setup the PR alarm limits. For example

imit is set to 50, an alarm will be issued when the PR value

When PR High limit is set to 130, an alarm will be issued when the PR value is higher than 130, and when PR low

po2 value is lower than 94.

H. 100

s lower than 50.

tem3.Setup the PR Alarm Limits

tem5.Check the software version The fifth item is to view the software version.

2.5 Environmental Specifications

Type of protection against electric shock: II (Internally powered equipment)

Operating mode: Spot checking

2.3 Physical Specifications

SpO2 declared accuracy 70%~100%: ±2digits SpO2 Display Range: 30%~99% SpO2 Resolution: 1%.

PR declared accuracy : 25~250 bpm: ±3digits

Atmosphere Pressure

Specification of battery: Two AAA (LR03) Operating current: 25-50mA

Width*Height*Depth: 57×30×31 mm Weight: 28g (Bare machine)

2.4 Measurement Specifications

PR Resolution: 1 bpm

Temperature 450~+104°F / +10~+40°C Operating: +50~+104°F / +10~+40°C Storage/Transportation: -4~+140°F / -20~+60°C Humidity 15~95%, noncondensing Storage/Transportation: 15~95%, noncondensing 10~95%, noncondensing

Operating: 70~106kpa Storage/Transportation: 50~107.4kpa

2.6 Display Display Type: Display Color: YM101: Red: YM102: Green YM103: White SpO2%, Pulse Rate, PI%, Display content:

Bar Graph, Battery Indicator

supported by clinical studies covering the entire claimed ange. The fraction of inspired oxygen (FiO2) delivered to steady-state saturation periods over the specified SpO2 accuracy range (e.g. 70 % to 100 %), then the SpO2

pulse oximeter to the values of SaO2 determined with a 2)The clinical trial included 11 subjects, including 6 male and 5 females, with an age range of 18 to 46 years, the subjects skin color included dark black, medium black,

The equipment's design life expectancy is about 2 years, keep your equipment and accessories free of dust and dirt.

A. Please clean the equipment before use according to hapter 3.2:Remove the batteries insidethe battery assette if the equipment will not be operated for a long

3.1 Maintenance

damage the equipment.

- B. Replace the batteries in time when the battery voltage indicate lamps were empty;
 C. It is recommended that the equipment should be kept in a dry environment with no corrosive gases and good ventilation anytime. The moisture and high-light environments will affect its lifetime and even might
 - e) Dry your equipment in a ventilated, cool place.
- temperature is between -20 to 60°C and the relative humidity is less than 95%.
- E. The packed equipment can be transported by ordinar conveyance. The equipment not be transported mixed with toxic, harmful, corrosive materials.

• No modification of this equipment is allowed.

icetone-based cleaners).

• If you spill liquid onto the equipment, contact us or

there is heavy pollution or lots of dust and sand in your place, the equipment should be cleaned more frequen Before cleaning the equipment, consult your hospital's regulations for cleaning the equipment. Recommended

- clean your equipment, follow these rules: a) Shut down the pulse oximeter
- b) Clean the display screen using a soft, clean cloth dampened with a glass cleaner;
 c) Clean the exterior surface of the equipment and
- robe using a soft cloth dampened with the cleaner: d) Wipe off all the cleaning solution with a dry cloth
- o avoid damage to the equipment, follow these rules:
- Always dilute according the manufacturer's instructions of ise lowest possible concentration. Do not immerse part of the equipment in the liquid.
- Do not pour liquid onto the equipment or Never use abrasive materials (such as steel wool or ilver polish), or erosive cleaners (such as acetone or
- your service personnel.





ate

3.3 Disinfection

Clean the pulse oximeter before disinfecting it. The recommended disinfectant is ethanol 70%. Disinfection

Never use ETO or formaldehyde for disinfection.

3.4 Disposal Dispose of the pulse oximeter in accordance with local environment and waste disposal laws and regulations.

One lanyard. Two AAA batteries(Optional).

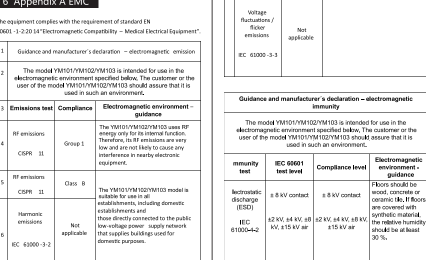
One user manual.

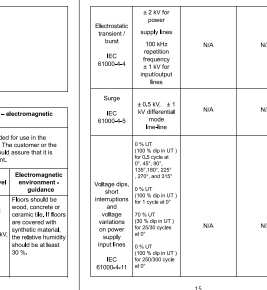
Trouble	Possible Reason	Solutions
The device	The batteries are drained away or almost drained away.	Replace batteries.
can not be turned on	The battery installation is incorrect.	Install the battery over again.
	The device works abnormally.	Please contact the product distributor

The Spo2 and PR are not displayed displayed normally User's blood perfusion Warm the finger is very low and try again The device was set to shut he display seconds when there is no Normal correct physiological signals suddenly The battery is almost Replace batteries drained away The finger is not inserted deep enough Replace the finger and try again The finger is shaking or displayed Not used in the work Please use in normal environment required | working environme by this manual Please contact the product distributor abnormally.

Trouble Possible Reason Solutions 50601 -1-2:20 14 "Electromagnetic Compatibility – Medical Electrical Equipment Guidance and manufacturer's declaration - electromagnetic emission The model YM101/YM102/YM103 is intended for use in the electromagnetic environment specified below. The customer or the user of the model YM101/YM102/YM103 should assure that it is used in such an environment. 3 Emissions test Compliance Electromagnetic environment guidance CISPR 11 Try to keep still RF emissions

IEC 61000 -3-2





				1
Electrostatic transient / burst IEC 61000-4-4	± 2 kV for power supply lines 100 kHz repetition frequency ± 1 kV for input/output lines	N/A	N/A	m
Surge IEC 61000-4-5	± 0.5 kV, ± 1 kV differential mode line-line	N/A	N/A	
Voltage dips, short interruptions and voltage variations on power supply input lines IEC 61000-4-11	0 % UT (100 % dip in UT) for 2.5 dip in UT) for 1.5 cycle at 0°, 45°, 90°, 135°, 180°, 225°, 270°, and 315° 0 % UT (100 % dip in UT) for 1 cycle at 0° 70 % UT (30 % dip in UT) for 2.5	N/A	N/A	C 6
		15		

frequency (50/60 Hz) magnetic fiel IEC 61000-4-8			ı, 50/60Hz	Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.	
NOTE [:] UT i	s the a. c. mai	ns voltage prio	or to applica	tion of the test level.	
Guidan	ce and manuf	facturer's dec		lectromagnetic	Re
electroma	he model YM1	nment specifie	d below. Th	for use in the e customer or the assure that it is	
			vironinent.		
Immunity test	IEC 60601 test level	Compliance level	Electroma	gnetic environment - guidance	61

50/60Hz	magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.					Recommended separation distance $d\left[\frac{3.5}{V_{1}}\right]\sqrt{P}$ $d\left[\frac{3.5}{E_{1}}\right]\sqrt{P} \text{ 80MHz}$
to applicat	tion of the test level.					
				10 V/m		800MHz
ration – e	lectromagnetic		Radiated	80 MHz to 2.7 GHz	10 V/m	$d \left[\frac{7}{E_1} \right] \sqrt{P} 800MH$ 2.7GHz
intended for use in the below. The customer or the 03 should assure that it is onment.			RF IEC 61000-4-3			where P is the ma output power rating transmitter in watts
Portable communication should be part of the cables, that separation	gnetic environment - guidance and mobile RF attions equipment used no closer to any the device, including an etwork distance calculated quation applicable to				17	according to the tran- manufacturer and d recommended sep- distance in metres(m). Field strengths from fix transmitters, as determi an electromagnetic survey, * should be les the compliance level in frequency range b
					17	

				distance
				$d\left[\frac{3.5}{V_1}\right]\sqrt{P}$
				$d \left[\frac{3.5}{E_1} \right] \sqrt{P}$ 80MHz to
		10 V/m		800MHz
=	Radiated RF IEC 61000-4-3	80 MHz to 2.7 GHz	10 V/m	$d \left[\begin{array}{c} 7 \\ E_1 \end{array}\right] \sqrt{P} \text{800MHz} \text{to}$ 2.7GHz where P is the maximum output power rating of the transmitter in 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 electromagnetic site survey, ^a should be less than the compliance level in each frequency range ^b
1			17	

		Recommended separation distance	
V/m //Hz to GHz	10 V/m	$d \left \lfloor \frac{3.5}{V_{\rm f}} \right \rfloor \sqrt{P} \text{80MHz} \text{to} \\ \text{800MHz} \\ d \left \lfloor \frac{7}{E_{\rm f}} \right \rfloor \sqrt{P} \text{800MHz} \text{to} \\ \text{2.7GHz} \\ \text{where P is the maximum output power rating of the transmitter in watts (W)} \\ \text{according to the transmitter manufacturer and d is the recommended separation distance in metres(m).}$	N N N ppi ol a ai ai 6,44 ai ai ai ai
		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	M to M 2
	17		<u> </u>

			Interference may vicinity of equip with the following (((•)))		
NOTE 1 At 80 MHz and 800 MHz, the higher frequency ran NOTE 2 These guidelines may not apply in all situations. E propagation is affected by absorption and reflection frobiects and people.					
and 80 MHz 6,795 MHz;	are 6,765 MH	z to 13,567 MHz;	dical) bands between		
amateur radi		een 0,15 MHz	and 80 MHz are 1		
	7 MHz to 7,3 I MHz to 18,17 M		z to 10,15 MHz, 1		
	21,4 MHz, 24 z to 54,0 MHz		,99 MHz, 28,0 MH		

		Interference may occur vicinity of equipment r with the following symb $\left(\left(\begin{pmatrix} \bullet \\ \bullet \end{pmatrix}\right)\right)$					
NOTE 2 These guideli	NOTE 1 At 80 MHz and 800 MHz, the higher frequency range appll NOTE 2 These guidelines may not apply in all situations. Electrom propagation is affected by absorption and reflection from stru objects and people.						
and 80 MHz are 6,765	a The ISM (industrial, scientific and medical) bands between 0,1 and 80 MHz are 6,765 MHz to 6,795 MHz; 13,553 MHz to 13,567 MHz; 26,957 MHz to 27,283 MI						
40,66 MHz to 40,70 MHz. The amateur radio bands between 0,15 MHz and 80 MHz are 1,8 MHz							
MHz, 3,5 MHz to 4,0 MHz, 5,3 MHz to 5,4 MHz, 7 MHz to 7,3 MHz, 10,1 MHz to 10,15 MHz, 14 MHz MHz, 18,07 MHz to 18,17 MHz,							
21,0 MHz to 21,4 MHz and 50,0 MHz to 54,0		24,99 MHz, 28,0 MHz to 29					

b The compliance levels in the ISM frequency bands between 150 kHz and 80 MHz and in the frequency range 80 MHz to 2,7 GHz are intended to decrease the likelihood that mobile/portable communications equipment could cause interference if it is inadvertently brought into patient areas. For this reason, an additional factor of 10/3 has been incorporated into the formulae used in calculating the recommended separation distance for transmitters in these frequency ranges.

c 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 re-orienting or relocating the YM101/YM102/YM103.

d Over the frequency range 150 kHz to 80 MHz, field strengths should be

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