# Section 1 Safety

internal servicing

Instructions for the Safe Operation and Use of the Fingertin Pulse Oximeter A310 Fingertin Pulse Oximeter Do not attempt to service the Fingertip Pulse Oximeter, Only qualified service personnel should attempt any needed

> Prolonged use or the patient's condition may require changing the sensor site periodically. Change sensor site and check skin integrity, circulatory status and correct alignment at least every 2 hours. SpO2 measurements may be adversely affected in the

presence of high ambient light. Shield the sensor area (with a surgical towel, or direct sunlight, for example) if necessary. The following reason will cause interference to the testing accuracy of the Fingertip Pulse Oximeter. · High-frequency electrosurgical equipment.

· Placement of a sensor on an extremity with a blood pressure cuff arterial catheter, or intravascular line The patient has hypotension severe vasoconstriction severe

anemia or hypothermia. The patient is in cardiac arrest or is in shock. · Fingernail polish or false fingernails may cause inaccurate

SpO2 readings. 1.2 Warnings

WARNING: EXPLOSION HAZARD — Do not use the Fingertip Pulse Oximeter in a flammable atmosphere where concentrations of flammable anesthetics or other materials nav occur

them to explode WARNING: Do not attempt to recharge normal dry-cell

The Oximeter can 1. Power of

The screen are

suddenly off

batteries, they may leak. And may cause a fire or even

nadequate or not finger

be there at all

he installed

3.The Oximeter

might be damaged

3.The Oximeter

incorrectly

WARNING: .Do not use the Fingertip Pulse Oximeter in an MDI or CT anvironment

> WARNING: Do not modify this equipment without thorization of the manufactures WARNING: If this equipment is modified, appropriate inspection and testing must be conducted to ensure continued

safe use of equipment. CAUTION: Keep the operating environment free of dust, vibrations, corrosive, or flammable materials, and extremes of temperature and humidity.

CAUTION: Do not operate the unit if it is damp or wet because of condensation or spills. Avoid using the equipmen immediately after moving it from a cold environment to a warm humid location

CAUTION: Never use sharp or pointed objects to operate the front-panel switches CAUTION: The batteries must be taken out from the battery compartment if the device will not be used for a long time.

CAUTION: The device shall only be used if the battery cover is closed.

CAUTION: The batteries must be proper disposed accord o local regulation after their use. CAUTION: The device should keep away from the children

pets and pests to avoid swallowing.

WARNING: Do not throw batteries in fire as this may causes

1.3 Definitions and Symbols

Symbol Description Type BF Equipment

Batch code<sup>4</sup>

LOT Date of manufacture\*

SN

mation of manufacture, including name nerature limitation

When the end-user wishes to discard this roduct, it must be sent to separate collection scilities for recovery and recycling

The information you should know to protect

The important information you should know

he equipment from possible damage

Follow instruction for use

Anti-dust & Anti-water class

he information you should know to protect patients and medical staff from possible injury Visual & Sound reminder function. Real-time spot-checks Low Battery voltage indicator.

in sports and etc.

2.3 Product Features

value and nlethysmogram\*

Automatically switch off

suitable for family, hospital (including clinical use in

· Color OLED display, simultaneous display for testing

Lightweight for carrying and Easy-To-Use.

Manually adjust the direction of interface.

Standard two AAA 1.5V Alkaline Battery support more han 20 hours continuous work.

\*Batch code, Date of manufacturer and Serial No are printed CAUTION: The device can not be used to measure the child on the label on the battery cover. below 3 years as the test result is not guarantee to accurate.

Section 2 Introduction

2.1 General This chapter provides a general description of the Fingertip

Pulse Oximeter including Brief device description Product features 2.2 Brief Device Description

Clinical testing is used to establish the SpO2 accuracy. The The Fingertip Pulse Oximeter, based on all digital technology. measured arterial hemoglobin saturation value (SnO2) of the s intended for noninvasive spot-check measurement of sensor is compared to arterial hemoglobin oxygen(SaO2) functional oxygen saturation of arterial hemoglobin (SpO2). value, determined from blood samples with a laboratory Advanced DSP algorithm\* can minimize the influence of CO-oximeter. The accuracy of the sensors in comparison to notion artifact and improve measurement accuracy of low The Oximeter can be used to measure human Hemoglobin Saturation and heart rate through finger. The product is

the CO-oximeter samples measured over the SpO2 range of 70 -100%. Accuracy data is calculated using the root-mean-square(Arms value) for all subjects. Only about two-thirds of FINGERTIP PULSE OXIMETER EOUIPMENT measurements can be expected to fall within internist/surgery. Anesthesia, pediatrics, intensive care and ±Arms of the value measured by a CO-Oximeter. etc.) Oxygen Bar, social medical organizations, physical care Pulse simulator shall be used to assess Pulse rate Accuracy. The measured pulse rate is compared to the preset pulse rate

and eximptome

root-mean-square (Arms value) for all subjects. \*DSP algorithm: Digital signal processor algorithm \*Low Perfusion: In physiology, perfusion is the process of a

Low Perfusion: 0.2%. (Advanced DSP algorithm can body delivering blood to a capillary bed in its biological mnrove measurement accuracy, under the condition of low tissue. Under the condition of low perfusion, the measurement of non-invasive saturation of pulse-blood oxygen is low-accurate

Electro-

contact

\*Plethysmograph: is an instrument for measuring changes in volume within an organ or whole body (usually resulting from fluctuations in the amount of blood or air it contains). PI (Perfusion Index ) is the ratio of the pulsatile blood flow

Floors should be

wood concrete or

value in simulator. Accuracy data is calculated using the

CAUTION: The Fingertip Pulse Oximeter is intended only

conjunction with other methods of assessing clinical signs

CAUTION: A function tester cannot be used to assess the

accuracy of a Fingertin Pulse Oximeter monitor or sensor.

as an adjunct in nationt assessment. It must be used in

to the non-pulsatile static blood flow in a natient's peripheral tissue, such as finger tip, toe, or ear lobe. Perfusion index is an indication of the pulse strength at the sensor site.

3.3.1 Remind Setup 3.1 Description of the Front Panel (as figure 3.1.1)



Table 3.1.1 Part Definition and Description

Section 3 Installation, Setup, and Operation

### Name Description

Power button	Turn on the machine	Press the direction button for 1 second, move the "*"
Power button	Display the SPO2/PR data & Plethysmogram	to the back of Beep, long press the direction button to on/off. (Note: When Beep is turned on, the sound emitted du
Battery Compartment		test indicates the pulse rate sound)

### 3.2.Display

Voltage 0 % UT: N/A

After switch on, the OLED display of the Fingertip Pulse Oximeter is as follows:

Mains power quality

should be that of a



Figure 3.2.1 OLED display

## 3 Parameter setting

When the device is under measuring interface, press the direction button for 1 second in order to enter into menu page (figure 3.3.1 and figure 3.3.2.). There are two submenus for

Remind Setup. User can adjust the setting through moving the "\*" symbol to the back of the Sound Reminder. Been. Restore or Brightness. Sound Reminder Press the direction button for 1 second, move the "\*" symbol

to the back of Sound Reminder, long press the direction button to turn it on/off (Note: If the measured value exceeds the maximum or

minimum value of SPO2 or PR, there will give off sound when sound reminder is turned on.)

to turn it during the

When the "\*" symbol show behind "Restore", long press the

direction button can be changed to "OK", which causes the device to restore factory data setting. Press the direction button for 1 second, move the "\*" symbol

to the back of Demo, long press the direction button to turn it on/off When the "\*" symbol show on "Brightness", long press the

direction button to change the Brightness value from 1 to 5.

3.3.2 Limit Value Setting When the \* symbol show on the Remind Setup, long press

guidance

Portable and mobile

RF communication

 $d = \left[\frac{7}{E}\right] \sqrt{P}$  800 MHz

where p is the

ransmitter

maximum output

power rating of the

transmitter in watts

(W) according to the

manufacturer and d is

to 2.5 GHz

menu (figure 3.3.2). User can press the direction button to select the items. And press the direction button for 1 second to change the data you need On the Reminder Limit setup menu page (figure 3.3.2), when

### Press the direction button for 1 second and enter into the

the \* symbol show behind the "+/-". Press direction button for 1 second to change the "+" to "-" or change the "-" to "+" When "+" shows on the right side, press the direction button for 1 second, move the "\*" after the Spo2 Hi or PR Hi setting. can increase the value to a higher value (until it reaches to the When "-" shows on the right side, press the direction button

the direction button until enter into the Reminder Limit setup

for 1 second, move the "\*" after the Spo2 Lo or PR Lo value setting, can reduce the value to a lower value (untill it reaches to the lowest)...

Limit Setup Sp02 Hi Brightness Evit Figure 3.3.2 Figure 3.3.1

the recommended

metres (m),b

transmitters, a

less than the

range. b

each frequency

Interference may

occur in the vicini

of equipment marks

with the following

determined by an

survey, a should be

fixed RF

Field strengths fro

separation distance in

2. The plethymogram can been regarded as correct if the 1. The sound reminder have 1 second delay after the incorrect

wave is fluctuated regularly. Section 4 Cleaning and Disinfection The customer can preset the limit value to the 98 or 99 to

check whether it is normal for sound reminder setting. 3.4.1 Install hattery

Installing two AAA batteries into battery cassette in correct

result being detected

polarities and cover it.



WARNING: Do not attempt to recharge normal alkaline batteries, they may leak and may cause a fire or even evplode

3.4.2 Turn the Fingertin Pulse Oximeter on Put one of fingers into rubber hole of the Oximeter (it is best to put the finger thoroughly) with nail surface upward, then

Press power button to turn the Fingertin Pulse Oximeter or

The oximeter will be automatically powered off when no

. When hattery power is at lowest level, the hattery canac

3.4.3 Read correspondent data from display screen

indicates symbol of " Tin OLED, remind users of

Switch off the power and take out the batteries before

Keen the exterior surface of the device clean and free of dust

and dirt. Cleaning exterior surface (OLED display screen

included) of the unit with a dry and soft cloth Use 75%

density of medical alcohol to clean the surface and use dry

fabric with little alcohol to avoid alcohol permeates into the

Field strengths from fixed transmitters, such as base

tations for radio (cellular/cordless) telephones and land

and TV broadcast cannot be predicted theoretically with

nobile radios, amateur radio, AM and FM radio broadcast

finger in the device for longer than 16 seconds.

releasing the clamp.

replacement of battery.

4.1 Cleaning

4.2 Disinfection

CAUTION: Don't use strong solvent. For example, acetone. CAUTION: Never use an abrasive such as steel wool or metal polich

natient use the machine in the hospital.

CAUTION: Do not allow any liquid into the product, and do not immerse any parts of the device into any liquids.

Disinfecting the machine after using by the patient if multiple

Use 75% density of medical alcohol to clean the surface that

CAUTION: Avoid pouring liquids on the device while

contacting with the patient.

CAUTION: Don't remain any cleaning solution on the

surface of the device. Section 5 Troubleshooting and Maintenance

# 5.1 Maintenance

Replace the batteries timely when battery indication is low. Clean surface of the Fingertip Pulse Oximeter before it is

used in diagnosis for natients Remove the batteries inside the battery cassette if the Oximeter will not be operated for a long time.

It is better to preserve the product in a place where ambient temperature is -25°C to 55°C(-13 F -131 F) and humidity is 15%-85% Regular inspection to make sure that no obvious damage

existed to affect the safety and performance of device. No flammable substance, overtop or lower temperature and humidity existed in operation conditions.

# 5.2 Troubleshooting

Oxyhemoglobin 1. Finger might

of heart rate is

shown unstably

or heart rate is

abnormal and

cause sound

reminder

Table 5.2.1 Troubleshooting Possible Reason

USER MANUAL

LOCKEE ( \( \xi\_{0123} \)

HEALTH - TECHNOLOGY

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Oxyhemoglobin . Finger is not or heart rate can gged correctly. not be shown Patient's perfusion is too Try some mor low to be times, if you can make sure about no problem existing in the product Please of to a hospital timely for exact

not be plugged

en enough

ovement status

Finger is not

Patient's

SPO2&PR is

ged correctly

Finger is

1.Retry by

plugging the

hospital for furthe

2. Try not to mov et the natient atient's body is in keep calm. Physical Characteristics plugging the 2. Go to the

Weight approx: 50g±2g (including 2 x AAA battery) Classification Anti-electric Shock Type: Internally powered equipment Anti-electric Shock Degree: Type BF equipment EMC: Type B Mode of operation: Continuous Operation Enclosure Degree of ingress protection: IP22 IP22 means shell of this product can withstand the water dropping to the surface when the shell deviate 15 degree from

2. Go to the 2.Batteries might | hospital for further xamination

might be damaged .The product is 2.Replace the

tomatically powered off when batteries o signal is detected longer than 16 seconds 2.Power quantity

exhausted Section 6 Specification The Fingertip Pulse Oximeter Specifications:

Dimensions: 62 mm (L) x 34mm (W) x 31mm (D)

of the batteries is

horizontal surface.

Internal:

Air Pressure

not be powered on batteries might be plugging the Power Consumption | Smaller than 45mA(Normal)

Environmental

Operating Temperature: 5°C to 40°C Storage Temperature: -25°C to 55°C Relative Humidity: 15% to 93% non-condensing 86Kpa-106Kpa

2xAAA 1.5v alkaline battery

### Sound Reminder Limit default value:

Parameter	Value
Hemoglobin saturation	Upper limit: 100/ bottom limit:94
Pulse rate:	Upper limit: 130 /bottom limit:50

# Electronics Parameters:

Parameter		Value	
Hemoglobin saturation display		35-100%	
		30-250 BPM	
	Hemoglobin Saturation	1%	
	Pulse rate	1 BPM	
	Hemoglobin Saturation	±2% (90%-100%) ±3% (70%-90%), Unspecified (<70%)	

Pulse rate ±1 BPM

0-1%: 0.1% Accuracy 1-20%: 1%

## Probe LED Specification:

	Wave Length	Radiant Power
RED	660±2 nm	1.8 mW
Infra RED	905±2 nm	2.0 mW

### Manufacturer's Declaration of the EMC Guidance and manufacturer's declaration - electromagnetic emission -for all EQUIPMENT AND SYSTEMS

		Н		assure that it is used in such an envir			
	30-250 BPM		3	Emissions test		Electromagnetic environment - g	
obin n	1%			RF emissions	Group 1	The A310 Pulse RF energy only in function. Therefore	
e	1 BPM	П	4			emissions are ve	
obin n	±2% (90%-100%) ±3% (70%-90%),			CISPR 11		not likely to caus interference in no electronic equipr	

# Resolution 0.1%

electromagnetic emission

Display 0-20%

# Guidance and manufacturer's declaration -

The A310 Pulse Oximeter is intended for use in the electromagnetic environment specified below. The customer or the user of A310 Pulse Oximeter should

Electromagnetic		
environment - guidance		
The A310 Pulse Oximeteruses		
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.		

CISPR 11 The A310 Pulse Oximeter i for use in all establishments emissions including domestic establishments and those 61000-3directly connected to the Voltage public low-voltage power supplynetwork that supplies ouildings used for domestic flicker

### Guidance and manufacturer's declaration electromagnetic immunity -for all EQUIPMENT and SYSTEMS Guidance and manufacturer's declaration -

61000-3

electromagnetic immunity The A310 Pulse Oximeter is intended for use in the electromagnetic environment specified below. The customer or the user of the A310 Pulse Oximetershould assure that it is used in such an environment. Immunity IEC 60601 | Complian- | Electromagnetic test level | ce level | environment -

### discharge ceramic tile. If floors (ESD) ± 15 kV are covered with ± 15 kV synthetic material, tl relative humidity 61000-4-2 should be at least 30 Electro- ± 2 kV for N/A Mains nower quality static should be that of a transient / supply typical commercial burst environment. 61000-4-4 | input/out-Surge + 1 kV Mains power quality should be that of a typical commercial 61000-4-5 mode environmen + 2 kV

±8 kV

dips, short 0.5 cycle interrup- g Power (50/60 magneti 61000-4-8 the test level

typical commercial or tions and At 0°, 45 nospital environment. voltage 90°, 135 If the user of the variations 180°, A310 Pulse Oximeter on power 225°, requires continued supply 270° and operation during input lines 315° ower mains interruptions, it is cycle and ecommended that the 61000-4-1 70 % UT: A310 Pulse Oximeter 25/30 be powered from an uninterruptible power supply or a battery. 0 % UT: 250/300 30 A/m 30 A/m Power frequency nagnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment

NOTE UT is the a. c. mains voltage prior to application of Guidance and manufacturer's declaration electromagnetic immunity -for EQUIPMENT and

SYSTEM that are not LIFE-SUPPORTING Guidance and manufacturer's declaration electromagnetic immunity

lectromagnetic environment specified below. The customer or the user of the A310 Pulse Oximeter should assure that it is used in such an environment. Immunity IEC 60601 | Complian- | Electromagnetic test level | ce level | environment -

The A310 Pulse Oximeter is intended for use in the

		Conducted RF IEC 61000-4-6	3 Vrms 150 kHz to 80 MHz	N/A
		Radiated RF	6Vrms in ISM banda between 150 kHz	10 V/
d		61000-4-3	to 80 MHz	
-			80 MHz to 2.7 GHz	

equipment should be used no closer to any part of the A310 Pulse Oximeter, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter Recommended separation distance  $d = \left[\frac{3.5}{\sqrt{P}}\right]\sqrt{P}$  $d = \left(\frac{3.5}{E_0}\right) \sqrt{F}$  80 MHz to 800 MHz

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

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 electromagnetic sit compliance level in A 310 Pulse Ovimete strengths should be less than 3V/m.

LIFE-SUPPORTING

Recommended separation distances between portable and SYSTEM -for EQUIPMENT and SYSTEMS that are not

between portable and mobile RF communications equipment

in which the A310 Pulse Oximeter is used exceeds the applicable RF compliance level above, the A310 Pulse Oximeter should be observed to verify normal operation, I abnormal performance is observed additional measures For transmitters rated at a maximum output power not listed nay be necessary, such as re-orienting or relocating the Over the frequency range 150 kHz to 80 MHz, field

for the higher frequency range applies. RF communications equipment and the EQUIPMENT or

Recommended separation distances between portable and mobile RF communications equipment and the A310 Puls

The A310 Pulse Oximeter is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the A310 Pulse Oximeter can help prevent electromagnetic interference by maintaining a minimum distance (transmitters) and the A310 Pulse Oximeter as recommended below according to the maximum output power of the communications

above the recommended senaration 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

Separation distance according to frequency of transmitter

kHz to 80 MHz 80 MHz to 800 MHz 800 MHz to 2.7 GHz

 $d = \left[\frac{3.5}{4}\right]\sqrt{P}$ 

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

 $d = [\frac{3.5}{3.5}]_{1/P}$ 

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