VITAL SIGNS MONITOR
INSTRUCTION MANUAL

MODEL: VI-200A
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CHAPTER 1 Introduction .......................................................................................................................... 5
  1.1 About the Manual ................................................................................................................................. 5
  1.3 Safety Information ............................................................................................................................... 5
  1.4 Explanation of Symbols ....................................................................................................................... 7
  1.5 Description of Abbreviation ................................................................................................................ 7
  1.6 Intended Use ....................................................................................................................................... 8
CHAPTER 2 Overview Of Monitor ............................................................................................................... 8
  2.1 Special Feature ..................................................................................................................................... 8
  2.2 Appearance of Monitor ....................................................................................................................... 8
  2.3 Specification ......................................................................................................................................... 13
CHAPTER 3 Patient Safety .......................................................................................................................... 14
  3.1 Environment ......................................................................................................................................... 15
  3.2 Grounding ........................................................................................................................................... 15
CHAPTER 4 Getting Started ...................................................................................................................... 15
  4.1 Unpacking and Inspection .................................................................................................................. 15
  4.2 Connect the Power Cables .................................................................................................................. 15
  4.3 Power on the Monitor ......................................................................................................................... 16
  4.4 Connect with Patient Sensors ........................................................................................................... 16
  4.5 Check the recorder .............................................................................................................................. 16
CHAPTER 5 Measuring screen and Main menu ........................................................................................... 16
  5.1 measuring screens ............................................................................................................................... 16
  5.2 Main menu ......................................................................................................................................... 18
CHAPTER 6 Date & Time, ID Settings ......................................................................................................... 18
  6.1 Time settings ....................................................................................................................................... 19
  6.2 ID number setting .............................................................................................................................. 19
CHAPTER 7 Take a Measurement ............................................................................................................... 20
  7.1 SpO₂ and PR measurement ................................................................................................................ 20
  7.2 NIBP Monitoring and precautions .................................................................................................... 22
  7.2.1 Introduction ................................................................................................................................... 22
CHAPTER 8 Data Management .................................................................................................................. 25
  8.1 SpO₂ & PR Record Review ................................................................................................................ 25
  8.2 SpO₂ Trend Review ............................................................................................................................ 26
  8.3 PR Trend Review ................................................................................................................................ 27
8.4 NIBP Record Review ................................................................. 27
8.5 NIBP Trend Review ................................................................. 27
8.6 Erase Data ................................................................................. 28
CHAPTER 9 NIBP Measurement Set ................................................. 28
CHAPTER 10 Alarm Set ................................................................. 29
  10.1 Alarm priority ........................................................................ 29
  10.2 Alarm set .............................................................................. 30
CHAPTER 11 Real Print Set(Optional) .................................................. 31
CHAPTER 12 Network Set(Optional) ................................................... 32
CHAPTER 13 System Set ................................................................. 32
CHAPTER 14 System Configuration ................................................... 33
CHAPTER 15 NIBP Operation .......................................................... 34
CHAPTER 16 System Info ................................................................. 34
CHAPTER 17 Trouble shooting .......................................................... 34
CHAPTER 18 Maintenance And Cleaning .......................................... 35
  18.1 Maintenance ......................................................................... 35
  18.2 Warranty and Repair .............................................................. 37
Appendix ......................................................................................... 38
CHAPTER 1 Introduction

1.1 About the Manual
Before using the vital signs monitor, the user must carefully read this manual so that the user can operate the monitor properly and make it reach the specific safety standard and performance index.

This manual explains how to set up and use the monitor. Important safety information relating to general use of the monitor appears after this introduction. Other important safety information is located throughout the text where appropriate.

Note: There requires no routine calibration, safety maintenance or in-service during the monitor’s life.

1.2 Contraindications
- Active patients.
- Intravascular dyes such as indocyanine green or methylene blue.
- Significant levels of dysfunctional hemoglobins (such as carbonxy-hemoglobin or methemoglobin).
- The presence of high ambient light. Shield the sensor area (with a surgical towel, or direct sunlight, for example) if necessary.
- Venous pulsations may cause erroneous low readings (e.g. tricuspid value regurgitation)
- Venous congestion may cause under reading of actual arterial oxygen saturation. Therefore, assure proper venous outflow from monitored site. Sensor should be not below heart level (e.g. sensor on hand of a patient in a bed with arm dangling to the floor)
- Avoid placing the sensor on any extremity with an arterial catheter, intravascular line or blood pressure cuff.
- Do not use the monitor when the patient is in cardiac arrest or is in defibrillation.
- Excessive caution with low perfused patient; skin erosion and/or pressure necrosis may occur.

1.3 Safety Information
Warnings alert the user to potential serious outcomes, such as death, injury, or adverse events to the patient or user.
Cautions alert the user to excess care necessary for the safe and effective use of the monitor.
Notes contain important information that may otherwise be overlooked or missed.

⚠️ Warnings
- The medical equipment must be manipulated by personnel who have already got relative training of operation.
- Do not use the monitor in a MRI or CT environment
- The monitor must be able to measure the pulse properly to obtain an accurate SpO₂ measurement. Verify that nothing is hindering the pulse measurement before taking the SpO₂ measurement.
- Explosion hazard: Do not use the monitor in the presence of flammable anesthetic or in an explosive atmosphere.
- Do not make any clinical judgments based solely on the monitor. The monitor is intended only as an adjunct in patient assessment. It must be used in conjunction with clinical signs
and symptoms.

- Prolonged use or the patient’s condition may require changing the sensor site periodically. The site must be checked at least every four hours to ensure adequate adhesion, circulation, skin integrity and correct optical alignment. If the circulatory condition or skin integrity is compromised, the sensor should be applied to a different site.

- Use only the battery, power cable and accessories appointed by manufacturer, as other accessories may cause improper performance or dangers.

- Connect the monitor to a three-wire, grounded, hospital-grade receptacle if necessary.

- By replacing the fuse, please use the safety fuse of the same type and rated fuse.

- When connecting the monitor to any instrument, verify proper operation before clinical use. Refer to the other device’s manual for full instructions. Accessory equipment connected to the monitors data interface must be certified according to IEC Standard 60601-1 for electro medical equipment. All combinations of equipment must be in compliance with IEC Standard 61601-1 systems requirements. To avoid potentially hazardous leakage currents, always check the summation of leakage currents when several item of equipment are interconnected.

- Before use the equipment, inspect whether all the cables are in good condition, the damaged cables and connectors must be replaced. Operator should examine whether the system is in correct working state and operating condition.

- As with all medical equipment, carefully route patient cabling to reduce the possibility of patient entanglement or strangulation.

- Do not use a tape to secure the sensor to the site; this can restrict blood flow and cause inaccurate readings. Use of additional tape can cause skin damage or damage the sensor.

- Dispose of the device and its accessories according to applicable local regulations.

- To avoid an electrical hazard, never immerse the unit in any liquid or attempt to clean it with liquid cleaning agents. Always disconnect monitor from AC Main Power before performing cleaning of maintenance.

- If the monitor becomes accidentally wet during use, discontinue operation of the monitor until all affected components have been cleaned and permitted to dry completely. Contact our local representative if additional information is required.

- The signal output part can be only connected to the computer complying with the requirements of IEC60950.

**Cautions!**

- Do not sterilize by irradiation, steam, autoclave or ethylene oxide.

- Operation of the monitor may be affected by the use of an electrosurgical unit (ESU).

- The system may not conform to all performance specifications if stored or used outside the environmental specification identified in specification.

- Alarm must be set up according to different situation of individual patient. Make sure that audio alarm can be activated when alarm occurs.

- Do not only depend on the alarm system, the doctor and nurse will not draw attention when an alarm turn down or turn off.

- Single-use accessories should never be reused.

- If the accuracy of any measurement does not seem reasonable, first check the patient’s vital signs by alternate means and then check the monitor for proper functioning.
For proper equipment maintenance, perform the service procedures at the recommended intervals as described in the manual.

If the monitor needs to be used continuously long-term, please note to connect the monitor with the main power supply by the alarm of battery, otherwise, the monitor will automatically shut down, which leads to the break-off of the monitoring.

The monitor can monitor only one patient synchronously.

Do not place the monitor in any position that might cause it to fall on the patient. Do not lift the monitor by the power supply cord or patient connections.

As to the other points for attention, please carefully read the relevant chapter in this instruction.

FEDERAL LAW (U.S.A) restricts this device to sale by or on the order of a physician.

1.4 Explanation of Symbols

<table>
<thead>
<tr>
<th>Icon</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Type BF applied part</td>
</tr>
<tr>
<td></td>
<td>Power on/off button</td>
</tr>
<tr>
<td></td>
<td>Alarm sound on indicator</td>
</tr>
<tr>
<td></td>
<td>Alarm sound off indicator</td>
</tr>
<tr>
<td></td>
<td>Alarm Inhibition indicator</td>
</tr>
<tr>
<td></td>
<td>Battery power indicator</td>
</tr>
<tr>
<td></td>
<td>Fuse</td>
</tr>
<tr>
<td></td>
<td>Up/increase button</td>
</tr>
<tr>
<td></td>
<td>USB transmission state indicator; When it is colored, it indicates USB transmission is activated; While it is grey, it indicates the USB transmission is off.</td>
</tr>
<tr>
<td></td>
<td>Pulse beep indicator; when it is colored, it indicates the beeper on, while it turns to grey, it indicates the beeper is off.</td>
</tr>
<tr>
<td></td>
<td>NET state indicator; when it is colored, it indicates Network function is activated, While it is grey, it indicates Network function is off.</td>
</tr>
<tr>
<td></td>
<td>Print icon; When print is set on, it is colored, while off, it is grey.</td>
</tr>
<tr>
<td></td>
<td>Down/decrease button</td>
</tr>
<tr>
<td></td>
<td>Alarm inhibition/off button or Return button</td>
</tr>
<tr>
<td></td>
<td>Confirm button</td>
</tr>
<tr>
<td></td>
<td>Equipotential grounding terminal</td>
</tr>
</tbody>
</table>

1.5 Description of Abbreviation

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>SpO₂</td>
<td>Arterial oxygen saturation</td>
</tr>
<tr>
<td>PR</td>
<td>Pulse rate</td>
</tr>
<tr>
<td>NIBP</td>
<td>Non-invasive measurement of blood pressure</td>
</tr>
</tbody>
</table>
### 1.6 Intended Use

The vital signs monitor is a portable device indicated for use in non-invasively measuring and displaying functional oxygen saturation of arterial haemoglobin (SpO₂), pulse rate (PR), Non-invasive measurement of blood pressure (NIBP) of adult and pediatric patients in hospitals, medical facilities, and sub-acute environments. The vital signs monitor is intended for spot-checking and/or continuous monitoring of patients.

### CHAPTER 2 Overview Of Monitor

#### 2.1 Special Feature
- Portable, compact structure
- LED & LCD display
- Display: SpO₂, Pulse rate, Pulse bar, SpO₂ plethysmogram, NIBP data and so on
- Convenient clinical operation
- Up to 99 patients information and 72-hour records storage
- Visible & 3-level audio alarm
- Battery-low indication
- Built-in rechargeable battery or AC power
- Built-in thermal recorder (optional)
- Probes suitable for adult
- Oscillometry method to measure NIBP and one hose cuff system.
- Probes suitable for pediatric or neonatal (optional)

#### 2.2 Appearance of Monitor

2.2.1 Front panel
Description of Fig.1:

1. **LCD displaying screen**: Display SpO₂ waveform, Pulse rate, Pulse amplitude indicator (blip bar), NIBP status indicator, USB indicator, the date & time, alarm limits, system icon and ID number.

2,8. **Pressure unit indication lamp**: Indicate the use of mmHg or kPa when the corresponding indication lamp on.

3,4,7. **LED displaying screen**: Display SYS, DIA, MAP of Non-invasive measurement of blood pressure (NIBP).

5. **SpO₂ sensor port**

6. **NIBP cuff port**

9. **NIBP button**: Press this button in any time, the cuff starts inflating and press it again, the cuff switches to deflate.

10. **Down button**: Press this button you can select different item and decrease the number.

11. **CHG indication lamp**: Battery charge indicator. If the battery is charging, the lamp will flash. When the battery is charged to full, the lamp will off.

12. **Power button**: Press the button for 3 seconds you can power on the monitor, or press the button for 4 seconds you can turn off the monitor.

13. **Internal battery indication lamp**:
   - Battery power enough: The lamp is always on.
   - Battery low: The lamp will be flashing.
   - Without battery: The lamp will be closed.

14. **UP button**: Press this button you can select different items and increase the number.

15. **MENU&OK button**: Press this button to confirm your setting and enter the menu submenu.

16. **Alarm silence/Return button**: ONLY in the measuring screen, Press this button to silence the audio alarm for 2 minutes, and repress to restore the audio alarm; Press it for over 2 seconds to turn off the audio alarm and repress it for over 2 seconds to restore the alarm. In main menu or submenu, press this button to return to the previous screen.

2.2.2 Rear panel
Description of Fig.2:
1 USB Interface: The monitor can be used as the USB device (The function is reserved.).
2 NET: The NET socket is connected with the center monitor system of our company (The function is reserved).
3 I/O: for programme upgrade (The function is reserved.).
4 Speaker
5 Power socket: AC Power supply socket.
6 Ground terminal

2.2.3 Battery installation

Description of figure 3:
1—Fixing hole of fixing screw for fixing battery cover.
2,3,4,5—Electrodes for battery.
6—Battery cover.

Unscrew the fixing screw in the battery cover of the monitor bottom panel, open the battery cover, and then place the battery into the battery box with the polarity correctly. The battery is shown in the fig.4.

Make sure that the poles of the battery are playing as the following position: 2 to 2', 3 to 3'; 4 to 4', 5 to 5'.
Notes:
- Make sure that the polarity of the battery is correctly inserted. Otherwise the unit cannot operate normally.
- If the monitor is not used for a long time, please charge and discharge the Ni-MH battery thoroughly once a month.
- If the battery low indication icon displays during measuring, please connect the monitor to AC power socket timely, avoiding the current measuring is affected.

⚠️ Warning!
Do not use the battery not appointed by our company.

2.2.4 Ground wire connection
Connect the monitor to the ground system with ground wire (refer to Fig.5), as the following steps:
Firstly, plug the part 1 to the monitor’s grounding terminal as Fig.6 shown.

Secondly, press the clap on the other terminal of the ground wire to connect with the user’s ground system. Refer to figure 7.

2.2.5 Printer (Optional)
The monitor can print the stored records and SpO₂ waveforms with the configured SP-B6J printer.
Description of Fig.8:
1—Open button, pull the jut of cassette door, you can open the paper cassette door.
2—No paper indicator, the lamp will be lighted when there is no paper.
3—Printer power indicator, when the printer is printing records the lamp will be lighted.
4—Output of printing paper.

Paper replacement

Note : Record paper requirement
Only standard 50(+0/-1) mm thermosensitive record paper can be used, otherwise the recorder may not function, the recording quality may be poor, and the thermosensitive print head may be damaged.

Steps for paper replacement:
1. Press the open button as the Icon indication and then pull the paper cassette door towards you until it is completely open.
2. Remove the spent paper core.
3. Place a new roll of paper into the paper cassette door with a few inches of paper being unrolled. And then push slightly the roll of paper to the paper cassette (refer to the Fig.10). Ensure proper orientation of paper roll.
4. Close the paper cassette door with a few inches of paper being kept outside of the door.

NOTE
- When the recorder is working, the record paper goes out steadily. Do not pull the paper, or the recorder may be damaged.
- Do not operate the recorder without record paper.
- Make sure that the paper surface with the heat sensitive material towards to the heat sensitive
2.3 Specification

**Power supply**

<table>
<thead>
<tr>
<th>Power supply</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Main power</td>
<td>100-230V AC</td>
</tr>
<tr>
<td>Internal power</td>
<td>7.2V DC</td>
</tr>
<tr>
<td>Rated power input</td>
<td>Less than 45VA</td>
</tr>
<tr>
<td>Fuse</td>
<td>250V I max 1 A</td>
</tr>
</tbody>
</table>

**Environment**

<table>
<thead>
<tr>
<th>Environment</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Operating Temperature</td>
<td>5°C to 40°C</td>
</tr>
<tr>
<td>Relative Humidity</td>
<td>≤ 80%, non-condensing</td>
</tr>
<tr>
<td>Standard Atmosphere Pressure</td>
<td>86kPa~106kPa</td>
</tr>
</tbody>
</table>

| Storage Temperature  | -20°C to 55°C                                      |
| Relative Humidity    | 0 to 93%, non-condensing                          |
| Standard Atmosphere Pressure | 50kPa~106kPa                                    |

**Display**

| Display              | LCD & LED                                         |

**Displayed Parameters**

<table>
<thead>
<tr>
<th>Alarms</th>
<th>High and low limits selectable on patient parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pulse &amp; SpO₂</td>
<td>Pulse Rate, SpO₂ plethysmograph, and oxygen saturation of blood hemoglobin</td>
</tr>
<tr>
<td>NIBP</td>
<td>SYS, DIA, MAP</td>
</tr>
</tbody>
</table>

**SpO₂**

<table>
<thead>
<tr>
<th>Display range</th>
<th>0 to 100%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Measurement Range</td>
<td>70 to 100%</td>
</tr>
<tr>
<td>Accuracy</td>
<td>80-100%:± 2%; 70-79%:± 3%; 0-69% : Unspecified</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LED Specifications</th>
<th>Wavelength</th>
<th>Radiant Power</th>
</tr>
</thead>
<tbody>
<tr>
<td>RED</td>
<td>660±3nm</td>
<td>1.8mW</td>
</tr>
<tr>
<td>IR</td>
<td>940±10nm</td>
<td>2.0mW</td>
</tr>
</tbody>
</table>

| Display Update       | <5s        |
| Resolution           | 1%         |

**Pulse Rate**
### NIBP measurement

<table>
<thead>
<tr>
<th>Measurement Range</th>
<th>Pediatric</th>
<th>Adult</th>
</tr>
</thead>
<tbody>
<tr>
<td>SYS</td>
<td>30-135</td>
<td>30-255</td>
</tr>
<tr>
<td>DIA</td>
<td>15-110</td>
<td>15-220</td>
</tr>
<tr>
<td>MAP</td>
<td>20-125</td>
<td>20-235</td>
</tr>
<tr>
<td>overvoltage protection of software</td>
<td>150mmHg</td>
<td>280mmHg</td>
</tr>
<tr>
<td>overvoltage protection of hardware</td>
<td>145 ± 5mmHg</td>
<td>280 ± 10mmHg</td>
</tr>
<tr>
<td>Resolution</td>
<td>1mmHg</td>
<td></td>
</tr>
<tr>
<td>Leakage rate</td>
<td>Less 6 mmHg/min</td>
<td></td>
</tr>
</tbody>
</table>

### Alarm Setting Range

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Upper alarm range</th>
<th>Low alarm range</th>
</tr>
</thead>
<tbody>
<tr>
<td>SpO₂(%)</td>
<td>71-100</td>
<td>70-99</td>
</tr>
<tr>
<td>PR (bpm)</td>
<td>31-254</td>
<td>30-253</td>
</tr>
<tr>
<td>SYS (bpm)</td>
<td>61-255</td>
<td>60-220</td>
</tr>
<tr>
<td>DIA (bpm)</td>
<td>31-220</td>
<td>30-180</td>
</tr>
<tr>
<td>MAP (bpm)</td>
<td>31-220</td>
<td>30-200</td>
</tr>
</tbody>
</table>

### Outline

<table>
<thead>
<tr>
<th>Dimension</th>
<th>296mmX166mmX96mm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weight</td>
<td>1.8kg (including the battery)</td>
</tr>
</tbody>
</table>

### Classifications

The device is BF type.

**Note:** the device might not meet its performance specifications if stored or used outside the manufacturer’s specified temperature and humidity ranges.

---

### CHAPTER 3 Patient Safety

The vital signs monitor is designed to comply with the International Safety requirements for medical electrical equipment.
3.1 Environment

Follow the instructions below to ensure a completely safe electrical installation. The environment where the Monitor will be used should be reasonably free from vibration, dust, corrosive or explosive gases, extremes of temperature, humidity, and so on. For a cabinet mounted installation, allow sufficient room at the front for operation and sufficient room at the rear for servicing with the cabinet access door open.

The Monitor operates within specifications at ambient temperatures between 5°C and 40°C. Ambient temperatures that exceeding these limits could affect the accuracy of the instrument and cause damage to the modules and circuits. Allow at least 2 inches (5cms) space around the instrument for proper air circulation.

**Note:** Make sure that during operation, the instrument is free of condensation. Condensation can form when equipment is moved from one building to another, thus being exposed to moisture and differences in temperature.

3.2 Grounding

To protect the patient and hospital personnel, the cabinet of the monitor must be grounded. Accordingly, The Monitor is equipped with a detachable 3-wire cable which grounds the instrument to the power line ground (protective earth) when plugged into an appropriate 3-wire receptacle. If a 3-wire receptacle is not available, consult the hospital electrician. If the capacity of the protective ground wires is in doubt, the equipment must be operated with internal power supply.

**CHAPTER 4 Getting Started**

**NOTE**

To ensure that the monitor works properly, please read Chapter 3, and follow the steps before using the monitor.

4.1 Unpacking and Inspection

Open the package and take out the monitor and accessories carefully. Keep the package for possible future transportation or storage. Check the components according to the packing list.

- Check for any mechanical damage.
- Check all the cables, modules and accessories.

If there is any problem, contact the distributor immediately.

4.2 Connect the Power Cables

Connection procedure of the AC power cable:

Make sure the AC power supply complies with following specification: 100-230 (VAC), 50/60 (Hz). Ensure that the AC outlet is properly grounded.
(1) Apply the power cable provided with the monitor. Plug the power cable into the power socket on the rear panel.

(2) Connect the other end of the power cable to a grounded 3-line power output.

**NOTE**
- Connect the power cable to the jack special for hospital usage.
- The battery needs to be charged after transportation or storage. If the power supply is not properly connected before turning on the monitor, it may not work properly because of insufficient power. Connect the main power supply to charge the battery.

### 4.3 Power on the Monitor

After the probe is connected to its input cables, turn the monitor on by momentary pressure on the front-panel POWER button for 3 seconds. As audible beep feedback after pressing, the monitor is initiated.

**NOTE**
- Check for all the functions that may be used to monitor and make sure that the monitor is in good status.
- The battery must be recharged to the full electricity after the battery low indicator appears.
- It is suggested to recharge the battery to the full electricity after each use of monitoring so as to reserve sufficient power in battery.

**WARNING**

If any sign of damage is detected, do not use it on any patient. Contact the biomedical engineer in the hospital or the distributor immediately.

### 4.4 Connect with Patient Sensors

Connect the probe to the monitor (refer to 7.1 of chapter 7).

### 4.5 Check the recorder

If your monitor is equipped with a recorder, open the recorder door to check if paper is properly installed in the output slot. If no papers present, do not press “PRINT” function button.

---

**CHAPTER 5 Measuring screen and Main menu**

**5.1 measuring screens**

**5.1.1** Press the button for about 3 seconds to turn the monitor on, one of the following
measuring screens will display.

![Fig. 11(1)](image1)

![Fig. 11(2)](image2)

Description of Fig.11(1) & (2):

1: **The current date and time**: 2008-08-25 18:00;
   - Silence audio alarm indicator: the current audio alarm status is on.
   - Battery power indicator.

2: The current system status in fig.11:
   - **USB function indicator status**: On
   - **Net function indicator status**: On
   - **Print indicator status**: the current status is on printing
   - Pulse beep sound status indicator: On
   - ID number: the current ID is 99.
   - Bed number: the current bed number is 255.

3: Display the current measuring information of user
   - **NIBP measuring mode**: the current mode is Auto and the auto mode cycle is 120 minutes.
   - **Patient type**: Pediatric and the initial pressure is 70 mmHg.
   - **Remaining time for next pressure measurement in Auto mode**: 50 minutes
   - **NIBP measuring mode**: Auto
   - **Patient type**: Pediatric
   - **Remaining time for next pressure measurement in Auto mode**: 50 minutes
4: The trend graphic of SpO₂ in Fig.11(1):
The list of the 7 NIBP records lastly measured in Fig.11(2).
5: Display of SpO₂ alarm limits, PR alarm limits indication, and measurement information.
The current SpO₂ alarm limits: Upper alarm: 100; Lower alarm: 95.
The current PR alarm limits: Upper alarm: 120; Lower alarm: 60.
The current measurement status: "SpO₂ measuring...".
Pulse bar.

5.2 Main menu
5.2.1 Menu 1
In measuring screen, short press the button for about 1 second to enter into menu 1, and press the button and button to look through the items. Please refer to Fig12.

5.2.2 Menu 2
In measuring screen, press the button for over 2 seconds to enter into menu 2 and then press the button and button to look through the items. Refer to Fig.13.

CHAPTER 6 Date & Time, ID Settings
The Monitor features flexible configurations. You can configure various aspects of the monitor, including TIME, ID number, printer and so on. Always set the date and time before using the unit for the first time. Set different ID numbers for different user.
Check whether the date and time are correct before using the unit, reset them if necessary. The date and time are important indicators when a measurement is taken.
6.1. Time settings

After entering into menu 2, press the button to select “Date And Time” item and press the button to confirm your selection, you will see the following picture.

![Fig.14](image)

Press the button and button to select the different data items, and press the button to enter the selected item. Then increase or decrease the data by pressing the button or the button and press the button to confirm your setting. Press the button to return to the previous screen.

The setting ranges of the date and time are as follows:

Year: 2000-2020
Month: 1-12
Day: 1-31
Hour: 0-23
Minute: 0-59
Second: 0-59

6.2. ID number setting

After entering into menu 1, press the button to select “Patient Info Set” item and press the button to confirm your setting, (refer to the Fig.15)

![Fig.15](image)

Press the button or button to set the suitable ID number, and confirm your selection by pressing button. Press the button to return to the previous screen. The setting
CHAPTER 7 Take a Measurement

7.1 SpO₂ and PR measurement

After the time and the ID number settings, connect the sensor and take a measurement according to the following pictures:

Firstly, select the suitable sensor in terms of type and dimension; secondly, connect the sensor with the monitor (refer to fig.16). Clip the sensor to the rational position of the patient finger (refer to fig.17).

NOTE: The sensor meets biology compatibility request.

Measuring display

- In standard measurement condition, the monitor will display “Searching for pulse” and “SpO₂ measuring…” successively on information column. Review the measuring result displayed on the measuring screen when taking a measurement.

- Display “Finger Out” when signal is inadequate or finger off.

⚠️ Note! DO not read the values during the signal inadequate indicator.

⚠️ Warnings!

1. The test methods used to establish the SpO₂ accuracy is clinical testing. The monitor used to measure the arterial haemoglobin oxygen saturation levels and these levels are to be compared to the levels determined from arterial blood sampling with a CO-oximeter.

2. The measurement would not be performed if the following instances come across in operation:

range of the ID number is 1~99.
- Shock
- Low temperature of hand
- Have taken vascular activity medicine
- Anemia
- carboxyhemoglobin
- methemoglobin
- methylene blue
- Indigo carmine

3. Use only SpO₂ sensors provided by manufacturers for SpO₂ measurements. Other SpO₂ sensors may cause improper performance.

4. Do not use a SpO₂ sensor with exposed optical components.

5. Excessive patient movement may cause inaccurate measurements.

6. Tissue damage can be caused by incorrect application or use of sensor, for example by wrapping the sensor too tightly. Inspect the sensor site to ensure skin integrity and correct positioning and adhesion of the sensor. More frequent inspection should be taken depend on different patients if necessary.

7. Set the upper limit of SpO₂ alarm to 100% means cut off the upper alarm. High density of oxygen will cause adverse affection to the neonate's. So the upper limit of SpO₂ alarm must be selected prudently according to the knowledge of clinical practice.

8. Inaccurate measurements may be caused by:
   - Incorrect sensor application or use
   - Significant levels of dysfunctional hemoglobins (such as carboxyhemoglobin or methemoglobin)
   - Intravascular dyes such as indocyanine green or methylene blue
   - Exposure to excessive illumination, such as surgical lamps (especially ones with a xenon light source), bilirubin lamps, fluorescent lamps, infrared heating lamps or direct sunlight
   - High-frequency electrosurgical interference
   - Venous pulsations
   - 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
   - There is arterial occlusion proximal to the sensor
   - The patient is in cardiac arrest or in shock

Loss of pulse signal can occur in any of the following situations:
   - The sensor is wrapped too tightly.
   - There is excessive illumination from light sources such as the surgical lamp, bilirubin lamp, or sunlight.
   - A blood pressure cuff is inflated on the same extremity as the one to which an SpO₂ sensor is attached.

Note
   ✦ Pulse sensor should obviate the light source, e.g. radial lamp or infrared lamp.
   ✦ The measured record is stored automatically every four seconds. The monitor can store 72 hours records.
   ✦ The new record will be stored with the initial records being erased when the
stored records are full.
✧ For more information on the section, please SpO₂ operator’ manual.

7.2 NIBP Monitoring and precautions

7.2.1 Introduction
1. The Non-invasive Blood Pressure (NIBP) module measures the blood pressure using the oscillometry method.
2. It is applicable for adult and pediatric usage.
3. There are three modes of measurement available: manual, Auto and STAT. Each mode displays the diastolic, systolic and mean blood pressure.

WARNING
● You must not perform a NIBP measurement on patients with sickle-cell disease or under any condition in which the skin is damaged or expected to be damaged.
● For a thrombasthemia patient, it is important to determine whether measurement of the blood pressure shall be done automatically. The determination should be based on the clinical evaluation.
● Ensure that the correct setting is selected when performing a NIBP measurement on children. It may be dangerous for the children to use an over pressure level.

◆ Measurement limitations
To different patient conditions, the oscillometry measurement has certain limitations. The measurement is in search of regular arterial pressure pulse. In those circumstances when the patient’s condition makes it difficult to detect, the measurement becomes unreliable and measuring time increase. The user should be aware that the following conditions could interfere with the measurement and avoid these to happen.
✧ Patient movement
Measurements will be unreliable or may not be possible if the patient is moving, shivering or having convulsions. These motions may interfere with the detection of the arterial pressure pulses. In addition, the measurement time will be prolonged.
✧ Cardiac arrhythmia
Measurements will be unreliable or may not be possible if the patient’s cardiac arrhythmia has caused an irregular heart beat. The measuring time thus will be prolonged.
✧ Heart-lung Machine
Measurements may not be possible if the patient is connected to a heart-lung machine.
✧ Pressure changes
Measurements will be unreliable or may not be possible if the patient’s blood pressure is changing rapidly over the period of time during which the arterial pressure pulses are being analyzed to obtain the measurement.
✧ Severe shock
If the patient is in severe shock or hypothermia, measurements will be unreliable since reduced blood flow to the peripheries will reduce pulsation of arteries.
✧ Heart rate extremes
Measurements can not be made at a heart rate of less than 40 bpm and greater than 240bpm.

7.2.2. Preparation for Cuff
WARNING
● Before starting a measurement, verify that you have selected a setting appropriate for your patient (adult or pediatric).
Do not apply the cuff to an upper arm that has an intravenous infusion or catheter in place. This could cause tissue damage around the catheter when infusion is slowed or blocked during cuff inflation.

Make sure that the air hose is connected with the blood pressure cuff and the monitor is neither blocked nor tangled.

It is recommended to only use the cuff and extended tube applied or appointed by our company.

If you are not familiar with this monitor and NIBP parameters, then follow this chapter’ sections:

1. Plug in the extended tube and switch on the system.
   - Ensure the luer lock to plug into the corresponding socket firmly if it is used. And when unplugging, drag its hoop backwards and pull out the luer lock.

2. Apply the blood pressure cuff to the patient’s arm.
   - Ensure that the cuff is completely deflated.
   - Apply the cuff of appropriate size to the patient, and make sure that the symbol “Ф” is over the appropriate artery. Ensure that the cuff is not wrapped too tightly around the limb. Excessive tightness may cause discoloration and eventual ischemia of the extremities.

   **NOTE**
   - The width of the cuff should be either 40% of the limb circumference (50% for paediatrics) or 2/3 of the upper arm length. The inflatable part of the cuff should be long enough to encircle 50~80% of the limb. The wrong size of cuff can cause erroneous readings. If the cuff size is in question, then use a larger cuff.
   - Make sure that the cuff edge falls within the range of mark `<->`. If it does not, use a larger or smaller cuff that fits better.

3. Connect the air hose to the extended tube. The limb chosen for taking the measurement should be placed at the same level as the patient’s heart. If this is not possible you should apply the following corrections to the measured values.
   ① If the cuff is placed higher than the heart level, add 0.9 mmHg (0.10kPa) for each inch of different.
   ② If it is placed lower than the heart level, deduct 0.9 mmHg (0.10kPa)for each inch of different.

   **⚠️ ALARM:** Please ensure not to inadvertently connect the extended tube to intravascular fluid systems, which allows air to be pumped into a blood vessel and threaten the patient’s life.

4. Check whether the patient mode is appropriately selected. For the detailed operation please refer to CHAPTER 9 NIBP Measurement set.

5. Select a measurement mode in the NIBP mode item. For the detailed operation please refer to CHAPTER 9 NIBP Measurement set.

6. Press the NIBP button on the front panel to start a measurement.

   **NOTE:** Once you press the NIBP button, the devices will startup the NIBP measurement after a delay of about 3 seconds, regardless of whichever screen the device is displaying, and if stopping the measuring, just press the NIBP button again. Please DO NOT press the NIBP button frequently during the delay time.

**7.2.3 Blood pressure measurement**
◆ The placement of cuff
Firstly select a suitable cuff in terms of type and dimension, secondly take off your coat, baring upper arm, then wrap the cuff closely to your upper arm. Make sure to keep the lower edge of cuff at 2.5 cm above elbow joint.

◆ NIBP measurement
Body blood pressure is of volatility character, therefore, you should take several measurements in several days to judge whether or not the blood pressure is duratively increased. Detailed measurement requirements are as follows:

1, Taking NIBP measurement at the same time everyday. And then repeat the measurement every 2 minutes, calculating the average of the two datum. If the difference between the two measurement results for systolic pressure(SYS) or diastolic pressure (DIA) is >5mmHg, you should take another measurement to get the average of the three measurement results.

2, Precautions
- Before and after taking exercise;
- In 1 hour after meals;
- Before and after drinking beer, coffee or red tea;
- Before and after bathe;
- Before and after smoking

3, Relax for at least 5 minutes and keep off smokes and coffee within 30 minutes before beginning with measurement. The patient had better sit on the backed chair with upper arm bared. Keep the elbow at the same level with heart. For standing measurement, the patient should stand for 2 minutes before measuring when he/she has been ever lying. For whichever method of measurement, be sure to place the sphygmomanometer at the same level with heart.

Note: The LED displays the results of the last blood pressure measurement until another measurement is completed.

WARNING
- Prolonged non-invasive blood pressure measurement in AUTO mode may be associated with purpuric, ischemia and neuropathy in the limb wearing the cuff. When monitoring a patient, examine the extremities of the limb frequently for normal color, warmth and sensitivity. If any abnormality is observed, stop the blood pressure measurements.
- If you are in doubt about the accuracy of any reading(s), check the patient’ vital signs by an alternative method before checking the functioning of the monitor.
- The blood pressure cuff should not be applied to the limb attaching the SpO₂ sensor, since cuff inflation will affect SpO₂ monitoring.
- Do not place the cuff on an extremity being used for intravenous infusion or any area where circulation is compromised or has the potential to be compromised.
- During use on patients, ensure that heavy objects are not placed on the hose. Avoid crimping or undue bending, twisting, or entanglement of the hose.
- When the measurement is performed on the pediatric patients. Make sure that correct mode setting has been selected (refer to NIBP Measurement Set section). Higher adult NIBP doesn’t fit for pediatric patients, wrong selection of patient mode may be dangerous to them.
Inaccurate measurements may result from such causes:

a. Limb’s twitch and tremble will cause inaccuracy or prolonged cycling of measurement; serious tremble will lead to the failure of measure.
b. Placing the cuff too loosely or tightly on the patient.
c. Leaky cuff or hose
d. Insure the NIBP and pulse rate within the range of this monitor.
e. Excessive patient motion will cause the inaccuracy, patient should be relax and avoid movement.

7.2.4 Pressure Safety Protection

- Automatic deflation will be activated when the cuff pressure exceed 280 mmHg under the adult mode and exceed 150 mmHg under the pediatric mode.
- You can press the NIBP button to cancel a NIBP measurement when necessary.
- When the monitor detect the pre-inflation pressure is not enough during measuring, the device will intellectually perform the air supplementing. The supplements will perform at most 3 times.
- The device applies the luer lock connector to the extended tube for tightness.

7.2.5 Maintenance and Cleaning

**WARNING**

- Do not squeeze the hose of cuff.
- Do not allow liquid to enter the connector socket when cleaning the monitor.
- Do not wipe the inner part of the connector socket when cleaning the monitor.
- When the reusable cuff is not connected with the monitor, or being cleaned, always place the cover on the rubber tube to avoid liquid permeation.

**NIBP cuff disinfection**

The cuff can be sterilized by means of conventional autoclaving, gas, or radiation sterilization in hot air ovens or disinfected by immersion in decontamination solutions, but remember to remove the rubber sleeve if you use this method. The cuff should not be dry-cleaned.

The cuff can also be machine-washed or hand-washed, the latter method may prolong the service life of the cuff. Before washing, remove the latex rubber sleeve, and for machine-washing, close the Velcro fastening. Allow the cuff to dry thoroughly after washing, and then reinsert the rubber sleeve.

### CHAPTER 8 Data Management

The monitor contains an internal memory that can store 72-hour data records. You can review, print or erase records. If all the records are cleared, they are not available to print.

**8.1 SpO₂&PR Record Review**

In “SpO₂&PR Record Review” screen, press the ↑ or ↓ button to page up or page down. After entering the window by pressing the  button, you can print, delete or review any ID’ SpO₂&PR records. Press the  button to return to the previous screen.
Select ID: with 1-99 IDs selectable for data reviewing.

Start Print: By the item, you can print the records of the selected ID.

Stop Print: By the item, you can stop the print action which is running, and several seconds later, the printer power will power off automatically. If you do not press the button, the printer will print all the records of current ID.

Delete: Delete all record data of the selected ID. A prompt will appear for enquiring your confirmation.

Review: Return to the record review screen of the selected ID.

Note:
1. If you do not perform the “Stop Print” command, the printer will continue printing data belonging to the current ID.
2. Display of 10 records as per 1 page.

8.2 SpO₂ Trend Review
You can review the trend of measured SpO₂ data.
In “SpO₂ Trend Review” screen, press the ▲ or ▼ button to page up or page down. After entering the window by pressing the ▼ button, you can review or delete the SpO₂ Trend chart of any ID. Refer to fig.19(1) or fig.19.(2). Press the ▼ button to return to the previous screen.

Select ID: with 1-99 IDs selectable for data reviewing.
Delete: Delete the trend data of the selected ID. A prompt will appear for enquiring your confirmation.
Review: Return to the trend review screen of the selected ID.
8.3 PR Trend Review
You can review the PR trend of the current ID or a different ID’s PR trend by setting manually. In menu 1, press the ▼ button to select the “PR Trend review” item and then press the ▶ button to enter into the submenu. The detailed operation is similar to the 8.2 SpO2 Trend Review.

8.4 NIBP Record Review
In menu 1, press the ▼ button to select the “NIBP Record Review” item and press ▶ to enter into this submenu. The detailed operation is similar to the 8.1 SpO2&PR Record Review.

8.5 NIBP Trend Review
You can review the NIBP trend chart of the current ID or other ID by setting manually. In menu 1, press the ▼ button to select the “NIBP Trend Review” item and then press the ▶ button to enter into the submenu. The detailed operation is similar to the 8.2 SpO2 Trend Review (refer to section 8.2).
8.6 Erase Data

Select the “Erase Data” item in menu 2 and then press the button to enter into the Erase Data screen (refer to Fig.23). Press the or button to select “Erase NIBP Data”, “Erase SpO₂ Data” or both of them for deleting. Then press “OK” to delete all the data records, or, move to “Cancel” item by pressing the button and press the button to return to previous screen. You will be taken to the previous screen by pressing the button.

![Fig.23]

CHAPTER 9 NIBP Measurement Set

Select “NIBP Measurement Set” item in menu 1 and then press the button to enter into submenu. Refer to the following figure. Press the or button to select item to set and press to pitch on the item (the background color of the selected item will be reversed color). Press the or button to set the selected item and then press the button to confirm.
Description of Fig.24

NIBP mode: Auto, STAT or Manual;

- **Auto mode:** press the NIBP button on the front panel to start the first auto measuring. The device will successively measure patient’s BP after taking break for the time you set in Interval item(refer to "Auto mode cycle"). If you press the NIBP button during Automatic mode, the measurement in progress will stop.

- **Manual mode:** under the mode, press the NIBP button, you can start a single measuring.;

- **STAT mode:** the monitor will measure the blood pressure continuously in 5 minutes. After completion of last measurement it waits 5 seconds, and then measures the pressure again.

*Note:* During the measurement in process, the change for mode is non-effective. You should cancel the measuring currently and then reset the mode.

**Auto mode cycle:** ONLY applied for auto mode, selected values contain 1, 2, 3, 5, 10, 30, 45, 60, 90, 120, 240, 480 minutes;

**Patient Type:** The blood pressure type monitored by the device, with two choices of Adult and Pediatric.

**Initial Pressure:** Cuff Pre-inflation value, with the range of 140-180 mmHg in Adult type or 70-120 mmHg in Pediatric type.

*NOTE:* Due to the air supplement function of the device, initial pressure is only indicated for the first inflation value.

**Pressure Unit:** The blood pressure unit: mmHg or kPa, and the matching indication lamp will turn on once you make a choice.

### CHAPTER 10 Alarm Set

#### 10.1 Alarm priority

There are three-level priorities for alarm.

High priority: the highest level alarm, indicate the patient is in the very dangerous situation.

Medium priority: indicate the warning should be paid attention.

Low priority: indicate the finger off or finger sensor off.

Alarm of this monitor includes technical and physiological alarm. All three priorities are divided by built-in module and can not be changed by user.

**Assignment of priority:**
High priority: NIBP (SYS, DIA and MAP) and SpO₂
Medium priority: PR and battery low
low priority: Other information
Note: 1. The alarm sound will go on for a cycle.
2. After silencing the alarm, the corresponding icon will indicate this.
3. The power low alarm: the corresponding indication lamp will be flashing.

**VISUAL ALARM:**
If the alarm is activated through over limitation of physiological alarm, corresponding data will be flashing with striking color (red for SpO₂ and yellow for PR), meanwhile, the alarm information is displayed in information column. If the alarm is activated by more than one physiological alarm, each parameter will be displayed, flashing with striking color, meanwhile, the alarm information is displayed in information column circularly.

**AUDIO ALARM:**
Audio alarms can be heard if there is no silence. The audio alarm has different tone pitch and on-off beep patterns for each alarm priority.

Medium priority: "du-du-du", beeps every 8 seconds.
Low priority: "du-", beeps every 20 seconds.
NOTE: The audio alarm issues according to the highest priority when several alarm events happens simultaneity, and these visual alarm information are displayed by turns.

**AUDIO ALARM INHIBITION/OFF:**
On the measuring screen, short press the button to silence the audio alarm for 2 minutes, the audio alarm indicator icon will be covered with dashed “ x” ; long press the button, a yellow prompt window will pop up, enquiring you whether to turn off audio alarm. Press the button to turn off the audio alarm or press any other button to exit from the operation, After pressing the button, the audio alarm indicator icon will be covered with real linear “ x” . And long press the button again, you can activate the audio alarm again.

**Note: for the information on alarm printing, please refer to CHAPTER 11 Real Print Set.**

**10.2 Alarm set**
In menu 1 screen, press the or the button to select the “ Alarm Set” item, and then press the button to enter into the submenu. Refer to the Fig.25. Press the or the button to select the item and press the to pitch on the item (the background color of the selected item will be reversed color), and then choose the switch status, increase or decrease alarm value by pressing the or button. Press the button to returning to the previous menu.
**Alarm limits setting range:**

- **SpO₂:**
  - Upper alarm range: 71-100
  - Low alarm range: 70-99
  - The default SpO₂ upper alarm is 100.
  - The default SpO₂ low alarm is 95.

- **PR:**
  - Upper alarm range: 31-254
  - Low alarm range: 30-253
  - The default PR upper alarm is 120.
  - The default PR low alarm is 50.

- **SYS:**
  - Upper alarm range: 61-255
  - Low alarm range: 60-220
  - The default PR low alarm is 160.
  - The default PR low alarm is 90.

- **DIA:**
  - Upper alarm range: 31-220
  - Low alarm range: 30-180
  - The default PR low alarm is 95.
  - The default PR low alarm is 60.

- **MAP:**
  - Upper alarm range: 31-220
  - Low alarm range: 30-200
  - The default PR low alarm is 110.
  - The default PR low alarm is 60.

**SpO₂/PR/SYS/DIA/MAP switch:**

- **ON:** Turn on the alarm.
- **Off:** Turn off the alarm even though the value is out of limits.

**Note:**

1. When the monitor is turned off, the alarm limits of the latest set are restored, so after the monitor is turned on, please set alarm limits again if necessary.

2. The alarm set should be handled by the qualified personnel.

---

**CHAPTER 11 Real Print Set (Optional)**

The monitor can print data by configured printer such as B6 printer.

In the menu 2 screen, press the ▲ or the ▼ button to select the “Real Print Set” item, and then press the ▶ button to enter into the item. Refer to the Fig.26. Press the ▲ or ▼ button to select the sub-items to set up. And press the ▶ button to enter. Set the selected item by pressing the ▲ or ▼ button, then press the ▶ button to confirm.

Press the ▶ button to returning to the previous menu.
Print Duration: The time interval of printing data, with the range of 4-40s; Numbers for selection contain 4,8,12,16,20,24,28,32,36,40.

Auto Print
Start Print: By the button, the printer begins to print the records after about 3 seconds with the change of the button to “Stop Print”. Pressing “Stop Print” will end the real time print, and the button returns to “Start Print”.

Print Cycle: The cycle of printing automatically, with the range of 1-60 min; The default is 1 min.

Alarm Print: The function is activated automatically when one or more parameter alarm occur: SpO₂, PR, SYS, DIA, MAP.

Note:
1. It is suggested the monitor be connected to the main power supply when applying the printer for printing.
2. for the information on printing, please refer to section 2.2.5 Printer (Optional).

CHAPTER 12 Network Set (Optional)

The function is reserved currently.

CHAPTER 13 System Set

In the menu 2 screen, press the ↑ or the ↓ button to select the “System Set” item, and then press the OK button to enter into the submenu.
Brightness
Press the ▲ or ▼ button to pick the “Brightness” sub-item and press the □ button to confirm (refer to Fig.27). Press the ▲ or ▼ button to adjust the degree of brightness.

After that, confirm your setting by pressing the □ button. The □ button is used for returning to the previous menu.
The backlight brightness level: 1-8 for selection, and default brightness is 7.

USB Mode
The function is reserved currently.

Alarm Volume
Press the ▲ or ▼ button to pick the “Alarm Volume” sub-item and press the □ button to confirm. Press the ▲ or the ▼ button to increase or decrease the value of alarm volume. After that, confirm your selection by pressing the □ button. The □ button is used for returning to the previous menu.
Alarm volume range: 1-8 and the default is 7.

Beep Volume
The detailed operation is similar to Alarm Volume.
Beep Volume:0-8, off the beep when set to 0; the default is 7.

Display Mode
By it, you can choose the display mode for measuring screen.
NIBP Record: Display the lastly 7 NIBP measurement data on the lower of measuring screen.
SpO₂ wave: Display the SpO₂ wave on the lower of measuring screen.

Language Set
For setting language if necessary. Only English is available currently.

CHAPTER 14 System Configuration

In menu 2 screen, pick and enter “System Configuration” item. Two listed options are shown as Fig.28. Press the ▲ or the ▼ button to choose and then press □ to confirm.
The button is used for returning to the previous screen.

Note that only one of Factory Default and User Saved can be chosen.

Fig.28

Restore selected configuration: Restore configuration you have selected. 
Save Current configuration: Save the current settings made by the current user.

Factory Default: initial settings made by manufacturer.
User Saved: the settings made by user last time.

CHAPTER 15 NIBP Operation

This settings are applied for production. For customers, it is not necessary to use.

CHAPTER 16 System Info

In the menu 2 screen, pick and enter “System Info” item. The button is used for returning to the previous screen.

By this menu you can look through the software and hardware version and the remaining time for SpO\textsubscript{2} and NIBP records.

CHAPTER 17 Trouble shooting

<table>
<thead>
<tr>
<th>Problems</th>
<th>Possible reason</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>SpO\textsubscript{2}% or pulse rate can not be shown normally</td>
<td>1. Finger is not plugged correctly 2. Patient’s SpO\textsubscript{2} value is too low</td>
<td>1. Retry by plugging the finger 2. Measure more times, If you can...</td>
</tr>
</tbody>
</table>
### CHAPTER 18 Maintenance And Cleaning

#### 18.1 Maintenance

Customers should be responsible for periodic maintaining of the product and its accessories. It is very important for our company to warrant the service and repairs. We reserve the rights to change the time limit of warranty and replacement if the following steps are non-implemented:

1. An effective maintenance plan should be designed for the product and its reusable parts.
accessories. It includes periodic inspections and cleaning. It should be accord with the policy of local infection control department or health institution.

(2) Be sure to disconnect power line from the product before cleaning and inspecting.

(3) Periodic cleaning (accordance with the policy of local infection control department or health institution). Dampen a cloth with a commercial, nonabrasive cleaner and wipe the tip, bottom, and front surfaces lightly. The following admmissive liquor can be used:

- Ammonia (diluted),
- Glutaraldehyde,
- Sodium hypochlorite bleacher (diluted)
- Mildness suds (diluted).

Please obey the following rules to prevent from damaging the product:

- Always use diluted liquor recommend by the manufacturer.
- Always wipe up cleaning liquor after cleaning.
- Never use cleaning matter containing wax.
- Never spray water or cleaning liquor over the product, or allows any liquid to flow into power switcher, connector, or other intake.
- Never use the following cleanser:
  - any kinds of abrasive cleaner and menstruum
  - acetone
  - ketone
  - spirituous cleanser
  - lycine.

- In order to clean display, please use clean flexible cloth and make it wet with cleanser in the glass. Never spray cleanser in the glass on the screen, or use alcohol or medical disinfector, such as glutaraldehyde or lycine.

- Please use warm wet cloth and mildness suds to clean cables. Other cleaning ways may reduce the life of cables and lead wires.

Recommendation:

- Do not power on/off frequently.
- Take down and safekeep probes, extended tube after using product.
- Please hold the product in package if the product would not work for a long time.
- Do not make the product contact with chemical medicine and reagent.  
  
  Battery Maintenance

- Do not squeeze the hose of cuff.
- Do not allow liquid to enter the connector socket when cleaning the monitor.
- Do not wipe the inner part of the connector socket when cleaning the monitor.
- When the reusable cuff is not connected with the monitor, or being cleaned, always place the cover on the rubber tube to avoid liquid permeation.
18.2 Warranty and Repair

18.2.1 Warranty and repair content
(1) Repair response time: AM9:00 to PM17:30 on Monday to Friday except legal holiday.
   Repair time: AM9:00 to PM17:30 on Monday to Friday except legal holiday.
(2) Repair service: Including telephone support, field inspecting, fittings replacement.
   - Telephone support: we can give guidance to customer’s engineer to inspecting the
     instrument when you dial our service line. Professional repair engineer online
     provides technical support.
   - Field inspecting: we will send engineers to repair the instrument if necessary.
     Certified engineers of our company or local repair team trained by our company
     provide this service.
   - Fittings replacement: if necessary, we will replace the damaged fittings according to
     contract. The damaged fittings should be returned to us except for special reason.
(3) Spare machine for repair: it is used to replace the damaged machine for customer using,
   customer should send the damaged machine to us to repair.
(4) Repair for sponsoring and contributing machine: customer should send the machine to
   us to repair.
(5) Updating software is free.

18.2.2 Exemption and restriction:
(1) Warranty does not apply to the damage or loss sustained due to well-known act of god,
    such as fire, earthquake, flood, thunder, cyclone, hail, electrical storm, blast, building
    collapse, commotion, etc.
(2) Non-service items:
    ① The cost and insurance of dismantling and testing, overhauling, reinstall, transfer,
       moving the instrument or parts.
    ② Damage or loss sustained due to inspected or repaired by other institute that is not
       certified
    ③ Damage or alteration by anyone else who are not our company authorized service
       personnel.
(3) The damage or lose sustained due to connection to peripheral equipment (such as
    printer, computer etc.), that are not provided by our company are not covered by the
    warranty.
(4) Obligation restriction: In the duration of warranty, if the operators use other fittings that
    are not provided by us, we reserve the right to cancel warranty.

18.2.3 Customer guarantees:
(1) Read the user manual carefully before operation.
(2) Operation and maintenance according to the user manual, and guarantee the requests
    of power and environment.

18.2.4 Non-warranty and Non-replacement Policy
   ● The work environment is not eligible. For example, if the relative humidity exceeds
     70%, circuit boards of the instrument may be damaged due to condensate.
   ● If voltage of power supply is fluctuant and exceeds 240VAC, the power adapter may
be damaged.
- There is smear or marks that are not belong to the instrument and cannot be removed from the outside surface of the instrument.
- The instrument or its fittings are mechanically damaged.
- The circuit is short and damaged due to liquor or other stuff flow in the instrument or its fittings.
- All probes and accessories are not free replacement.
- Leakage of air cell of blood pressure sleeve due to improper storage or operation is not free replacement.
- The malfunction with result form improper repair by anyone other than our company authorized service personnel.
- The malfunction with result from improper use.

18.2.5 Special warranty period required by Customer

We stipulate the warranty period according to the relevant electronic regulation of country, the unit’s warranty period is one year and the accessory’s is three months. When customer requires to extending the warranty period, you should consider whether it is reasonable. AS the electronic product’s quickly replacing speed, if the warranty period over three years, purchased accessories may be out of stock. In this case, we will adopt to entirely upgrade or replace the old, you should pay the minimum acceptable cost of renewed device.

18.2.6 Repackaging

Remove all the detectors, leads and accessories and put them into the plastic bag. Try to use the original packaging case and materials. Any damage due to the improper packaging during the transportation shall be responsible by the user.

If you are still within the period of warranty, please present the warranty card and one copy of the invoice or receipt.

Please present a written note detailing all the troubles when repairing the instrument.

### Appendix

**List of Accessories**

The accessories list below is specified to be used in this device of our company. The user can order the various accessories according to the hospital requirements.

The standard accessories:

<table>
<thead>
<tr>
<th>No.</th>
<th>Accessories</th>
<th>Quantity</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>SpO₂ probe for adult(&gt;30Kg)</td>
<td>1 piece</td>
</tr>
<tr>
<td>No.</td>
<td>Accessories</td>
<td>Quantity</td>
</tr>
<tr>
<td>-----</td>
<td>----------------------------------------------------------------------------</td>
<td>----------</td>
</tr>
<tr>
<td>2</td>
<td>Power cord&lt;br&gt;Standard power cord (3× 0.75mm²)</td>
<td>1 piece</td>
</tr>
<tr>
<td>3</td>
<td>Ground wire (Φ6)</td>
<td>1 piece</td>
</tr>
<tr>
<td>4</td>
<td>Operator’s manual</td>
<td>1 piece</td>
</tr>
<tr>
<td>5</td>
<td>NI-MH Battery Pack</td>
<td>1 piece</td>
</tr>
<tr>
<td>6</td>
<td>Screwdriver</td>
<td>1 piece</td>
</tr>
<tr>
<td>7</td>
<td>Pressure cuff for Adult 25 cm to 35 cm</td>
<td>1 piece</td>
</tr>
<tr>
<td>8</td>
<td>10’ NIBP extension hose</td>
<td>1 piece</td>
</tr>
</tbody>
</table>

The optional accessories:

<table>
<thead>
<tr>
<th>No.</th>
<th>Accessories</th>
<th>Quantity</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>SpO₂ probe for adult&lt;br&gt;Model: M-50G19CSO</td>
<td>1 piece</td>
</tr>
<tr>
<td>2</td>
<td>SpO₂ probe for adult or pediatric(Pediatric 10-50 Kg)&lt;br&gt;Model: M-50D18CSO</td>
<td>1 piece</td>
</tr>
<tr>
<td>3</td>
<td>SpO₂ probe for infant&lt;br&gt;Model: M-50B16CSO</td>
<td>1 piece</td>
</tr>
<tr>
<td>4</td>
<td>SpO₂ probe for neonate(&lt;3Kg)&lt;br&gt;Model: M-50C17CSO</td>
<td>1 piece</td>
</tr>
<tr>
<td>5</td>
<td>SpO₂ probe for pediatric(Pediatric 10-50Kg)&lt;br&gt;Model: M-50H15CSO</td>
<td>1 piece</td>
</tr>
<tr>
<td>6</td>
<td>Pressure cuffs:&lt;br&gt;Adult 25 cm to 35 cm&lt;br&gt;Child 18 cm to 26 cm&lt;br&gt;Large Arm 33 cm to 47 cm&lt;br&gt;Thigh 46 cm to 66 cm</td>
<td>1 piece</td>
</tr>
<tr>
<td>7</td>
<td>Thermal array recorder (including one roll paper for recorder)&lt;br&gt;Model: SP-B6J</td>
<td>1 group</td>
</tr>
</tbody>
</table>