Preface

Thank you for using patient monitor.

In order to enable you to skillfully operate Monitor as soon as possible, we provide this user's manual with delivery. When you install and use this instrument for the first time, it is imperative that you read carefully all the information that accompanies this instrument.

Based on the need to improve the performance and reliability of the parts and the whole instrument, we sometimes will make some amendments to the instrument (including the hardware and software). As a result, there might be cases of discrepancies between the manual and the actual situation of products. When such discrepancies occur, we will try our best to amend or add materials. Your comments and suggestions are welcome.

Our liaison-way:

Address: No.2 Innovation First Road, Technical Innovation Coast, Hi-tech Zone, Zhuhai, P.R.China

Post code: 519085

Fax: +86-756-3399919

Toll-free consultation hot line: +86-400-8818-233

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The contents and version contained in this manual are subject to amendments without notification.

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The version number of this manual: V5.0
The version number of software: V2.2.4
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Liabilities of the Manufacturer

Only under the following circumstances will manufacturer be responsible for the safety, reliability and performance of the instrument.

➡ All the installation, expansion, readjustment, renovation or repairs are conducted by the personnel certified by manufacturer.

■ The electrical safety status at the installation site of the instrument conforms to the national standards.

■ The instrument is used in accordance with the operation procedures.

CE mark



EC Representative Name:

Shanghai International Trading Corp. GmbH (Hamburg)

EC Representative Address:

Eiffestrasse 80, 20537 Hamburg Germany

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Chapter 1 General Introduction

1.1 Intended use

The Monitor is used to monitor patient's physiological parameters such as ECG RESP SpO₂, NIBP, IBP, TEMP, CO₂, AG and ICG continuously. It is intended to be used in various hospital rooms such as Coronary Care Unit, Intensive Care Unit, Neonatal Intensive Care Unit and Operating Room to provide additional information to medical and nursing staff about the physiological condition of the patient.

It is not intended to be used in outdoor transport applications.

1.2 About this Manual

This user's manual consists of the following chapters:

Chapter 1 gives an introduction to the content and the specific signs of this manual, the main features and appearance of the monitor, the basic operations of various buttons, the meanings of the signs on the monitor, specifications and performance criteria of the monitor, the ambient requirements for the working and storage of the monitor.

Chapter 2 gives important safety notes <u>Please do read this chapter before using the</u> <u>monitor!</u>

Chapter 3 gives an introduction to the preparatory steps before using the monitor.

Chapter 4 provides general operation instruction for the monitor, including illustrations of the screen display, normal selection for soft button on screen, details for entry of patient data and trend maps, also.

Chapter 5 gives details of specific parameter measurement, preparatory steps, cables or probes connection, setup of parameters, maintenance and cleaning of equipments and sensors.

Chapter 6 gives detailed description of system alarm, including level and mode of alarm, default setting and changing procedure of alarm parameters, prompt of specific alarms, and the general operation to carry out when an alarm occurs.

Chapter 7 gives detailed description of record function.

Chapter 8 gives general maintenance and cleaning methods of the monitor and its parts.

Signs in this manual:



Warning: Means it must be strictly followed so as to prevent the operator or the patient from being harmed.



Caution: Means it must be followed so as not to damage the instrument.



Note: Important information or indications regarding the operation or use.

C Note: This manual introduced the product that with full configuration. Some functions of the product you bought may not be provided.

1.3 Brief Introduction to the Monitor

The monitor has features as follows:

■ Multiple measuring functions include 3-lead, 7-lead, 12-lead ECG/HR, RESP, dual TEMP, SpO₂/Pulse, NIBP, dual IBP, CO₂, AG and ICG are optional.

■ Complete built-in module design ensures stable and reliable performance

■ Unique all-lead ECG on-one-screen display, which can facilitate the diagnosis and analysis of cardiac disease

 $rac{}$ Can store the trend data for 120 - 168 hours and has the function of displaying trend data and trend graphs

■ Function of alarm event reviewing, can store 1000 - 1800 pieces of alarm events

■ Function of NIBP measurement reviewing, can store 750 – 1000 pieces of NIBP measurement data

■ Function of reviewing 10 - 30 minutes one important lead's EGC waveform

Built-in recorder is optional and it supports real-time recording, trigger printout by alarm

- Parameter display with big character
- Optional function of Calculator of drug concentration
- Optional function of Display of oxyCRG
- Function of Display of short trend
- 12.1" or 10.4" authentic color high brightness TFT LCD monitor
- Portable design, stylish and convenient
- Support connecting to VGA display for video output
- Rechargeable maintenance-free battery, can continue working when AC power is off
- Can be connected with the central unit to realize centralized monitoring
- Is resistant to high-frequency electrotome and is protected against defibrillation effects



1.4 Appearance and Structure of the Monitor

Fig. 1-4-1 The appearance of M8000 patient monitor



Fig. 1-4-2 The appearance of M9000 patient monitor



Fig. 1-4-3 The appearance of M8000A patient monitor



Fig. 1-4-4 The appearance of M9000A patient monitor

Caution: The AC input socket at the back panel of the monitor can be connected with 100-240V AC power by electrical wires supplied with this instrument.

1.5 Sockets



Fig. 1-5-1 Various sockets on the side panel



Terminal

Fig. 1-5-2 Various sockets on the back panel

Note: The 9 Pin D type socket (RS-232) is only used for maintenance and upgrading of the monitor by manufacturer. If the monitor is configured with VGA function, the socket will be changed to 15 Pin D type for connecting to the standard VGA display.

P Note: The Network Connector is a standard RJ45 socket and being used for connection with the central monitoring system provided by manufacturer.

Warning: The sensor cable sockets on Monitor can only be connected with the sensor cables supplied with this instrument and no other cables shall be used.

Notes on the signs on the monitor

Signs	Notes on the signs	
4 • •	Defibrillator-proof type CF equipment (Refer to IEC 60601-2-27) The unit displaying this symbol contains a F-Type isolated (floating) applied part providing a high degree of protection against shock, and is defibrillator-proof.	
۱ ۲ ۲	Defibrillator-proof type BF equipment (Refer to IEC 60601-1:1995) The unit displaying this symbol contains a F-Type isolated (floating) applied part providing a high degree of protection against shock, and is defibrillator-proof.	
\triangle	Attention! Please refer to the document supplied with this instrument (this manual)!	
(((•)))	Non-ionizing radiation	
4	Dangerous voltage	
\bigtriangledown	Equipotentiality	
\sim	Alternating current (AC)	
C € 0123	CE mark	
	Symbol for the marking of electrical and electronics devices according to Directive 2002/96/EC. The device, accessories and the packaging have to be disposed of waste correctly at the end of the usage. Please follow local ordinances or regulations for disposal.	

Signs	Notes on the signs
Hospital Only	Symbol marked on a tag attached to the supply cord of the monitor to warn that the supply cord should be connected to the sockets which are Hospital Only to achieve grounding reliability.
ECG	Short for "Electrocardiogram"
RESP	Short for "Respiration"
SpO ₂	Short for "Pulse Oxygen Saturation"
ТЕМР	Short for "Temperature"
IBP	Short for "Invasive Blood Pressure"
NIBP	Short for "Non-invasive Blood Pressure"
CO2	Short for "Carbon dioxide"
AG	Short for "Anesthetic gas"
ICG	Short for "Impedance cardiography"

Solution Note: The two channels of ECG1, ECG2 for ECG are used for the realization of the input for all-lead electrocardiogram; plug the 5-lead ECG cable with limb lead (RA, LA, C, RL, LL) into ECG1 (Channel 1), and the chest lead cable (C1, C2, C3, C5, C6) into ECG2 (Channel 2), and at this moment, the 12-lead (all-lead) ECG monitoring can be realized. In case only 7-lead monitoring is needed, then it is only necessary to plug the 5-lead ECG cable of the limb lead (RA, LA, C, RL, LL) into ECG1 (Channel 1). Be careful not to plug it into ECG2 (Channel 2).

1.6 Function Buttons and Trim Knob on the Front Panel

M8000/M9000





The Trim Knob is used for: Turn left or turn right to move the cursor. Press down to perform an operation, such as open the menu dialog or selects one option.

Fig. 1-6-1 Function Buttons and Trim Knob on the Front Panel

1.6.1 The Signs and Operation Instructions Within the Function Button Zone

Signs	Notes on the signs	Operation instructions of function buttons
AC/BAT	Indicating light of AC/DC	When the monitor is connected to the AC power, this indicating light is green (it is unrelated to the ON/OFF state of the monitor). When the monitor is not connected to AC power and the battery is used as the power source, this indicating light is orange.
CHARGE E [↓]	Indicating light of CHARGE	When the monitor is connected to the AC power of charge, this indicating light is turn-on. When the monitor is full of charge, this indicating light is turn-off.
Ċ/⊙	Power button	Press this button once and the monitor starts up. Repress this button, then the monitor is switched off.
MAIN	Return to Main Screen	Press this button once to exit the present menu and return to main screen.
TREND	Trend Review	Press this button once to see the Trend Graph and the Trend Table
FREEZE /RECORD ©/ 🛐	Switching type button Freeze (or defreeze) the waveforms /Record the real-time waveforms	Press this button in 2 seconds to freeze waveform, press again to defreeze waveform. Press this button over 2 seconds can start real-time recording. In case the real-time recording is underway, pressing this button will terminate real-time recording.
SUSPEND /SILENCE 滋/英	Switching type button Suspend the sounding of Alarm /Close the sounding of Alarm	Press this button in 2 seconds to make the monitor alarm paused or cancel the pause. Press this button over 2 seconds can silence the monitor's audio system or cancel the silence.

Signs	Notes on the signs	Operation instructions of function buttons
NIBP /STAT &/ &	Switching type button Begin (or Stop) the measurement of NIBP /Begin the STAT	Press this button in 2 seconds to start or stop the NIBP measurement. Press this button over 2 seconds to make NIBP module working at STAT measurement mode and perform continuous NIBP measurement within 5 minutes.
MENU	Menu	Press this button to display menu option.

1.6.2 Basic Operations



Fig. 1-6-2 Flow chart of basic operations

Solution Note: The system menu is located at the left bottom corner. By operating the Trim Knob in the above flow chart, select the options or make them spring out, and for detailed item selection, please refer to Chapter 4.

Chapter 2 Important Safety Notes

Warning: For pacemaker patients, Rate meters may continue to count the pacemaker rate during occurrences of cardiac arrest or some arrhythmias. Do not rely entirely upon rate meter alarms. Keep pacemaker patients under close surveillance. See this manual for disclosure of the pacemaker pulse rejection capability of this instrument.

Warning: Only trained doctors and nurses can use the device.

Warning: The monitor is neither a therapeutic instrument nor a device that can be used at home.

2.1 General Safety

1. Safety precautions for safe installation

□ The input socket of monitor can be connected to the electrical wires and common electrical wire can be used.

 \blacksquare Only the power supply type of AC 100-240V 50/60Hz specified by monitor can be used.

■ Connect the electrical wire to a properly grounded socket. Avoid putting the socket used for it in the same loop of such devices as the air conditioners, which regularly switch between ON and OFF.

■ Avoid putting the monitor in the locations where it easily shakes or wobbles.

Enough space shall be left around the monitor so as to guarantee normal ventilation.

■ Make sure the ambient temperature and humidity are stable and avoid the occurrence of condensation in the work process of the monitor.

Warning: Never install the monitor in an environment where flammable anesthetic gas is present.

2. Monitor conforms to the safety requirements of IEC 60601-1:1988+A1:1991+A2:1995. This monitor is protected against defibrillation effects.

3. Notes on signs related to safety



Defibrillator-proof type CF equipment (refer to IEC 60601-2-27)

The unit displaying this symbol contains a F-Type isolated (floating) applied part providing a high degree of protection against shock, and is defibrillator-proof.

The type CF applied parts provide a higher degree of protection against electric shock than that provided by type BF applied parts.



Attention! Please refer to the documents accompanying this monitor (this manual)!



Defibrillator-proof type BF equipment (IEC 60601-1:1995) The unit displaying this symbol contains a F-Type isolated (floating) applied part providing a high degree of protection against shock, and is defibrillator-proof.

4. When a defibrillator is applied on a patient, the monitor may have transient disorders in the display of waveforms. If the electrodes are used and placed properly, the display of the monitor will be restored within 10 seconds. During defibrillation, please note to remove the electrode of chest lead and move the electrode of limb lead to the side of the limb. The electrode of the defibrillator should not come into direct contact with the monitoring electrodes. Please ensure the monitor is reliably grounded and the electrodes used repeatedly should be kept clean.

Warning: When conducting defibrillation, do not come into contact with the patient, the bed and the monitor. Otherwise serious injury or death could be resulted in.

5. To guarantee the safe operation of the monitor, Monitor is provided with various replaceable parts, accessories and consuming materials (such as sensors and their cables, electrode pads). Please use the products provided or designated by the manufacturer.

6. Monitor only guarantees its safety and accuracy under the condition that it is connected to the devices provided or designated by manufacturer. If the monitor is connected to other undesignated electrical equipment or devices, safety hazards may occur for causes such as the cumulating of the leakage current.

7. To guarantee the normal and safe operation of the monitor, a preventive check and maintenance should be conducted for the monitor and its parts every 6-12 months (including performance check and safety check) to verify the instrument can work in a safe and proper condition and it is safe to the medical personnel and the patient and has met the accuracy required by clinical use.

8. When THERMAL CUT-OUT operates, the monitor will function continually without any SAFETY HAZARD by using its rechargeable battery. However, the battery is not being charged anymore. While the battery is low the monitor will give both visible and audible warnings in 5 minutes.

9. The American mains plug whose ground is in the same plug with its other two cords is Hospital Only. The supply cord should be connected to the sockets which are Hospital Only to achieve grounding reliability.

Caution: The monitor does not contain any parts for self-repair by users. The repair of the instrument must be conducted by the technical personnel been authorized by manufacturer.

2.2 Some important notes for safety

PATIENT NUMBER

The monitor can only be applied to one patient at one time.

INTERFERENCE

Do not use cellular phone in the vicinity of this equipment. High level of electromagnetic radiation emitted from such devices may result in strong interference with the monitor performance.

ACCIDENTAL SPILLS

To avoid electric shock or device malfunction, liquids must not be allowed to enter the device. If liquids have entered the device, take it out of service and have it checked by a service technician before it is used again.

ACCURACY

If the accuracy of any value displayed on the monitor or printed on a printout paper is questionable, determine the patient's vital signs by alternative means. Verify that all equipment is working correctly.

ALARMS

Do not rely exclusively on the audible alarm system for patient monitoring. Adjustment of alarm volume to a low level or off during patient monitoring may result in a hazard to the patient. Remember that the most reliable method of patient monitoring combines close personal surveillance and correct operation of monitoring equipment.

The functions of the alarm system for monitoring the patient must be verified at regular intervals.

BEFORE USE

Before putting the system into operation, please visually inspect all connecting cables for signs of damage. Damaged cables and connectors must be replaced immediately.

Before using the system, the operator must verify that it is in correct working order and operating condition.

Periodically, and whenever the integrity of the product is in doubt, test all functions.

CABLES

Route all cables away from patient's throat to avoid possible strangulation.

TO CLEAR PATIENT DATA

When monitoring a new patient, you must clear all previous patient data from the system. To accomplish this, shut down the device, and then turn on it. Selecting $\langle New patient \rangle$ in $\langle main \ setup \rangle$ menu can also clear the previous patient data.

DISPOSAL OF PACKAGE

Dispose of the packaging material, please observe the applicable waste control regulations and keeping it out of children's reach.

EXPLOSION HAZARD

Do not use this equipment in the presence of flammable anesthetics, vapors or liquids.

LEAKAGE CURRENT TEST

When interfacing with other equipment, a test for leakage current must be performed by qualified biomedical engineering personnel before using with patients.

BATTERY POWER

The device is equipped with a battery pack. The battery discharges even when the device is not in use. Store the device with a fully charged battery and take out the battery, so that the service life of the battery will not be shortened.

DISPOSAL OF ACCESSORIES AND DEVICE

Disposable devices are intended for single use only. They should not be reused as performance could degrade or contamination could occur.

The service life of this monitor is five years. At the end of its service life, the product described in this manual, as well as its accessories, must be disposed of in compliance with the guidelines regulating the disposal of such products. If you have questions concerning disposal of products, please contact manufacturer or its representatives.

EMC

Magnetic and electrical fields are capable of interfering with the proper performance of the device. For this reason, make sure that all external devices operated in the vicinity of the monitor comply with the relevant EMC requirements. X-ray equipment or MRI devices are a possible source of interference as they may emit higher levels of electromagnetic radiation. Also, keep cellular phones or other telecommunication equipment away from the monitor.

INSTRUCTION FOR USE

For continuous safe use of this equipment, it is necessary that listed instructions were followed. However, instructions listed in this manual in no way can supersede established medical practices concerning patient care.

LOSS OF DATA

Should the monitor at any time temporarily lose patient data, close patient observation or alternative monitoring devices should be used until monitor function is restored.

If the monitor does not automatically resume operation within 60 seconds, restart the monitor using the power on/off switch. Once monitoring is restored, you should verify correct monitoring state and alarm function.

2.3 Classifications

The Monitor is classified, according to IEC 60601-1:1988+A1:1991+A2:1995 as:

Type of protection against electric shock:	I
Degree of protection against electric	BF: CO ₂ , AG
shock:	CF: ECG, RESP, TEMP, IBP, NIBP, SpO ₂ ,
	ICG
Degree of protection against harmful	Ordinary Equipment (enclosed equipment
ingress of water:	without protection against ingress of water)
Degree of safety of application in the	Not suitable
presence of a flammable	
anesthetic-mixture with air or with	
oxygen or nitrous oxide:	
Mode of operation:	Continuous operation

I: Class I equipment

BF: Type BF applied part

CF: Type CF applied part

Not suitable: Equipment is not suitable for use in the presence of flammable anesthetic mixture with air or with oxygen or nitrous oxide.

2.4 Safe Operating and Handling Conditions

Method(s) of sterilization or disinfection	Sterilization: not applicable	
recommended by the manufacturer:	Disinfection: See "The Maintenance and	
	Cleaning of the System->General Cleaning"	
Electromagnetic interference	No cellular telephone nearby	
Electro surgical interference damage	No damage	
Diathermy instruments influence	Displayed values and prints may be disturbed	
	or erroneous during diathermy.	
Defibrillation shocks	The monitor specifications fulfill the	
	requirements of IEC 60601-1, IEC 60601-2-27,	
	IEC 60601-2-49, IEC 60601-2-34	
A uniliarry autouts	The system must fulfill the requirements of	
Auxinary outputs	standard IEC 60601-1-1	

Chapter 3 Preparations Before the Use of the Monitor

3.1 Unpacking the Case

 \blacksquare Unpack the packaging case

Open the packaging case, accessories include: electrical wire, various patient sensors and user's manual (this manual), warranty card, certificate and particular paper and the lower foam case contains the monitor.

Remove the monitor and accessories

Caution: Please place the monitor on level and stable supporting plane, not on the places that can easily shock or wake. Enough room should be left around the monitor so as to guarantee normal ventilation.

E Keep all the packaging materials for future use in transportation or storage.

□ Check the monitor and accessories

Check the monitor and its accessories one by one in accordance with the particular paper. Check to see if the parts have any mechanical damages. In case of problems, please contact us or our agent.

3.2 Connecting to Power

3.2.1 AC Power

■ Confirm the rated AC current is: AC 100-240V 50/60Hz

■ Use the electrical wires provided along with the instrument, put its output end plug (round headed) into the AC current socket on the back of the monitor, and the plug of input end into a grounded socket of the mains (It must be a special socket of the hospital), connect the monitor through the earth one of electrical wires.

■ When the indicating light above the power switch on the panel of the monitor is green, it means the AC power is on. And when the monitor is not connected to AC power and the DC battery is used as the power source, the indicating light is orange.

Warning: The monitor must be connected to a properly installed power outlet with protective earth contacts only. If the installation does not provide for a protective earth conductor, disconnect the monitor from the power line and operate it on battery power. **Completely only by disconnecting the power supply from the wall socket.** The wall socket has to be easily accessible.

Note: For measurements in or near the heart we recommend connecting the monitor to the potential equalization system. Use the green/yellow potential equalization cable and connect it to the pin labeled with the \bigtriangledown symbol.

3.2.2 Battery Power

The monitor has a battery pack to provide power to the monitor whenever AC power is interrupted. The battery is generally referred to as the "battery".

You must charge the battery before using it. There is no external charger. The battery is charged when the monitor is connected to AC power. To assure a fully charged battery that is ready for use, we recommend that the monitor be plugged into AC power whenever it is not in use.

Run time of the batteries is according to the usage and configuration of monitor. NIBP and SpO₂ monitoring and the usage of the recorder will drain battery power faster than other parameters.

Note: When the monitor is connected to AC power, the battery is in a state of being recharged. When it is unable to be connected to the AC power, the battery can be used to supply power, and at this time it is unnecessary to use the electrical wires, and the instrument can be switched on directly.

Note: A "Battery Low" message displaying at the technical alarm information area of screen and an audible system alarm indicate approximate 5 minutes of battery life remaining. You should connect the monitor to an AC power source when the message is displayed.

Note: This monitor contains a rechargeable battery. The average life span of this type of battery is approximately three years. When replacement becomes necessary, contact a qualified service representative to perform the replacement.

Disposal Note: Should this product become damaged beyond repair, or for some reason its service life is considered to be at an end, please observe all local, state, and federal regulations that relate to the disposal of products that contain lead, batteries, plastics, etc.

Install Battery

The battery storage is located at the bottom of the monitor, following the steps to install a battery.

- 1. Open the battery gate according to the direction marked on the monitor.
- 2. Turn the baffle up clockwise.
- 3. Push the battery into the gate with the electrode point to the bottom of the monitor.
- 4. After pushing the battery inside the storage withdraw, the baffle turn back to the middle position.
- 5、 Close the gate.

Uninstall battery

- 1. Open the battery gate according to the direction marked on the monitor.
- 2、Turn the baffle up clockwise.
- 3、 Take out the battery. Then close the gate.

3.3 Connecting to the Central Monitor System

Warning: Accessory equipment connected to the analog and digital interface must be certified according to the respective IEC standards (e.g. IEC 60950 for data processing equipment and IEC 60601-1:1988+A1:1991+A2:1995 for medical equipment). Furthermore all configurations shall comply with the valid version of the system standard IEC 60601-1-1. Everybody who connects additional equipment to the signal input part or signal output part configures a medical system, and is therefore responsible that the system complies with the requirements of the valid version of the system standard IEC 60601-1-1. If in doubt, consult the technical service department or your local representative.

If the user intends to connect the monitor to the central monitoring system, plug its connecting electrical cable into the Network Connector at the back of the monitor.

Note: This monitor can only be connected to the central monitoring system provided by manufacturer, do not attempt to connect this monitor to other central monitoring system.

3.4 Power on the Monitor

■ Press the power switch on the front panel of the monitor.

 \blacksquare About 30 seconds after the monitor is switched on, after passing the self-examination of the system, the monitor enters the monitoring screen.

Warning: In case the monitor is found to be working abnormally or indication of errors appears, please do not use this monitor for monitoring and should contact the after-sale service center as soon as possible.

3.5 Connecting to Various Kinds of Sensors

Connect sensor cables to the relevant sockets on the monitor and put sensors on the monitored locations on the body of the patient. Refer to the relevant content of **Chapter 5** for details.

Warning: For safety reasons, all connectors for patient cables and sensor leads (with the exception of temperature) are designed to prevent inadvertent disconnection, should someone pull on the leads. Do not route cables in a way that they may present a stumbling hazard. Do not install the monitor in a location where it may drop to the patient. All consoles and brackets used must have a raised edge at the front.

3.6 Preparation of Recorder

If the monitor you use has been provided with a recorder, before starting of monitoring please check if the recorder has had recording thermal paper installed. The thermal side (that is the smoother side) should face upwards and a small section should be pulled out onto the outlet of the paper (on the right panel of the monitor).

If record paper has been used up, following the steps to install recording paper.

- 1. Push down and pull out of the recorder cover to open recorder.
- 2. Install the paper with the thermal side upwards.
- 3. Close the recorder with a section of paper outside of the storage.

For detailed operation information, refer to Fig. 3-6-1



Fig. 3-6-1 Install Recording Paper

3.7 Shutting off the Monitor

Please follow these steps to shut off the monitor:

- Confirm that the patient monitoring is finished.
- Disconnect the cables and sensors form patient.
- Confirm that the monitoring data is stored or cleared.
- Press the power switch, then a dialog will pop up to ask you make sure the shut-off operation. Select "OK" to shut off the monitor. If the monitor can't be switched off normally, forced close the monitor by pressing and holding the power switch more than 5s. This may cause some damages to the device.

Chapter 4 Operation Instructions for the Monitor

Previous Note: In each menu, press 〈Previous〉 to return to the previous menu and press the 〈Main〉 button to return to main screen. In all the dialogue windows, there is help info to indicate the current operation.

Note: The monitor configuration is consist of standard and non-standard parameter configuration, and their operation methods are basically the same, the standard configuration includes 5-lead ECG, RESP, SpO₂, Single TEMP and NIBP modules, and the non-standard parameter configuration includes Dual TEMP, IBP, CO₂, AG and ICG modules.

4.1 Screen mode

In the **Select Screen>** of the **Main Setup>**menu, 8 kinds of different screen display modes can be selected, namely: Standard, NIBP Review, Big Numerics, Short Trend, 7 leads, 12 leads, oxyCRG, Other Bed. They are respectively showed as follow:

1) Standard



The ECG waveform of one lead is displayed on the uppermost region above the waveforms (this lead is called key monitoring lead and is set by the **<ECG1>** option in **<ECG>**), and the waveforms below are displayed differently according to different configurations.

2) NIBP Review



The recent groups of NIBP measurement results are displayed below the waveforms and the measurement records can be browsed by turning the trim knob.

3) Big Numerics



The main parameters are displayed in big font, e.g. HR, SpO₂, NIBP, RESP and EtCO₂.

4) Short Trend



The short trend diagram relevant to the parameters is displayed on the upper-left corner of the waveform.

5) 7-Leads



The ECG waveforms of 7-lead are displayed in the waveform display zone, they are I, II, III, aVR, aVL, aVF, and V- respectively.

6) 12-leads



The 12-lead ECG waveforms are displayed in the waveform display zone, they are I, II, III, aVR, aVL, aVF, V1, V2, V3, V4, V5, V6.

In order to facilitate the diagnosis and analysis of heart disease, monitor is especially designed with displaying 12-lead (all-lead) ECG synchronously on screen. Select $\langle 12$ **leads** \rangle in the **<Select Screen>** of the **<Main Setup>**menu. The waveform is composed of left and right regions, and displayed in the left region are ECG waveforms of limb leads and on the right are the ECG waveforms of chest leads.





The trend diagrams of HR, SpO₂ and RESP within 16 minutes are displayed under the waveforms.

8) Other Bed



The info for other beds is showed below the waveforms, including one waveform and parts of parameters. Among them, through **<Bed NO>**, the number of online machine can be selected and through **<Bed wave>** the waveform display of other beds can be selected. Press **<Run>** to initiate monitoring of other beds, and press **<Stop>** to terminate the present monitoring of other beds. Switching from monitoring of other beds screen to other screens will automatically terminate the present monitoring of other beds.

4.2 Main menu



Screen Such eight display modes as Standard, NIBP Review, Big Numerics, Short Trend, 7 leads, 12 leads, oxyCRG and Other Bed can be selected. And the display mode varies according to different manufacturer configurations.

Monitor Click and open the dialog of monitor configuration. Conduct some configurations of the monitor.

Trend Review Click and open the dialog of trend browse. Browse trend tables or trend diagrams.

Alarm Review Click and open the dialog of alarm event review. Browse alarm events.

ARR Review Click and open the dialog of arrhythmia review. Browse the waveforms and events of arrhythmia.

Alarm Setup Click and open the dialog of alarm configuration. Conduct configuration of alarm parameters.

New Patient Terminate the monitoring of the current patient and initiate the monitoring of a new patient. Pressing the option will delete the monitoring data of the current patient and patient Info and initiate the monitoring of a new patient.

Patient info Click and open the dialog of patient info. It provides the input and browse of patient info.

Calculator Click and open the dialog of calculator.

Standby Click and enter the standby state.



4.2.1 Monitor Setup

Monitor Setup
Beep Volume 2
Alarm Volume <mark>2</mark>
Wave Setup 🛄
Select Modu. 📭
Trend store 🛄
Short Trend 🛄
SystemSetup 📑
System Info 🛄
Demo Demo
Previous
Start/stop DEMO.

Beep Volume Set the volume of BEEP and options are **Off**, **1**, **2** and **3**. After one selection is made, a testing beep will be produced.

Alarm Volume Set the alarm volume and options are Off, 1, 2 and 3. After one selection is made, a testing beep will be produced.

Wave Setup Click and open the dialog of waveform configuration. Conduct the customization of screen waveforms and relevant waveform displays can be selected according to needs.

Select Modu. Click and open the dialog of module configuration. Some of the modules not in current use can be switched off, and after switching-off, the relevant parameters and waveforms will not be displayed and no alarm will be made.

Trend Store Click and open the dialog of configuration of trend storage. It provides the configuration function on the mode of trend storage and several modes of trend storage can be defined.

Short Trend Click and open the dialog of short trend diagram. Some scales and time of short trend diagram can be defined.

System Setup Click and open the dialog of system configuration. Conduct the configuration and maintenance of systems.

System Info Click and open the dialog of system info. Some info of the system will be displayed, such as version info.

Demo Switch on or switch off demonstration function

■ Waveform Setup

Waveform 1	ECG1
Waveform 2	SpO2
Waveform 3	Resp
Waveform 4	IBP1
Waveform 5	IBP2
Waveform 6	OFF
Waveform 7	OFF
Waveform 8	OFF
Waveform 9	OFF
Waveform 10	OFF
Previous	
Set No.1 wa	veform.

Waveform 1 Select the waveform displayed in the first line, and according to the lead types, different ECG waveforms can be selected (Note: The lead must be the ECG waveform, and cannot be switched off). At 3-Leads mode, it is the key monitoring lead and it is defaulted as Lead II.

Waveform 2 Select the waveform displayed in the second line, and options are **Off**, **Cascade** and **random waveform**. When selecting *<***Cascade***>*, waveform 2 is the cascade of waveform 1.

Waveform 3 Select the waveform displayed in the third line. Select **Off** close the wave display or select certain waveform to display.

Waveform 4 Select the waveform displayed in the fourth line. Select **Off** close the wave display or select certain waveform to display.

Waveform 5 Select the waveform displayed in the fifth line. Select **Off** close the wave display or select certain waveform to display.

Waveform 6 Select the waveform displayed in the sixth line. Select **Off** close the wave display or select certain waveform to display.

Waveform 7 Select the waveform displayed in the seventh line. Select **Off** close the wave display or select certain waveform to display.

Waveform 8 Select the waveform displayed in the seventh line. Select **Off** close the wave display or select certain waveform to display.

Waveform 9 Select the waveform displayed in the seventh line. Select **Off** close the wave display or select certain waveform to display.

Waveform 10 Select the waveform displayed in the seventh line. Select **Off** close the wave display or select certain waveform to display.

Select Module



SpO₂ Module Enable/Disable the display of SpO₂ module. After switching-off, the SpO₂ parameters and relevant alarm will not be displayed and the current SpO₂ waveform will be automatically switched off. After it is open, the SpO₂ waveform will also be opened.

NIBP Module Please refer to **SpO₂ module** instruction

Resp Module Enable/Disable the display of Resp module. After switching-off, the Resp parameters and relevant alarm will no be displayed and the current Resp waveform will be automatically switched off. After it is open, if there is no CO_2 module, the Resp waveform will be opened automatically.

CO₂ Module Enable/Disable the display of CO_2 module. After switching-off, the CO_2 parameters and relevant alarm will no be displayed and the current CO_2 waveform will be automatically switched off. After it is open, the CO_2 waveform will be automatically open, if there is an RESP waveforms, the RESP waveform will be switched off.

Gas Module Please refer to SpO₂ module instruction.

ICG Module Please refer to SpO₂ module instruction.

Temp Module Click and open the dialog of Temp module setup.



Temp 1 ModuleEnable/Disable the display of Temp 1 moduleTemp 2 ModuleEnable/Disable the display of Temp 2 moduleIBP ModuleClick and open the dialog of IBP module setup



IBP1 Module Enable/Disable the display of IBP1 module. After switching-off, no IBP1 parameters and relevant alarm will be displayed and the current IBP1 waveform will be automatically switched off. After it is open, the IBP1 waveform will also be opened.

IBP2 Module Please refer to IBP1 module instruction

Trend Storage Setup

Trend Storage Setup	
Interval	1 min
NIBP storage	ON
ALM storage	OFF
Warn storage	OFF
Previous	
Set auto trend interval time.	

Interval Select the cycle intervals of trend storage and options are Off, 1min, 2min, 3min, 4min, 5min, 10min, 15min, 20min, 25min and 30min.

NIBP storage Enable/Disable the switch of NIBP storage. When it is enabled, it indicates after NIBP measurement completed, a record will be stored.

ALM storage Enable/Disable the switch of alarm storage. When it is enabled, it indicates if there is a high alarm of physiological parameters a record will be stored.

Warn storage Enable/Disable the switch of warning storage. When it is enabled, it indicates if there is a medium alarm of physiological parameters a record will be stored.

Short trend Setup

Short Trend Setun	
Time Scale	30 min
HR Scale	0~300/min
Soll2 Scale	60~100%
Doon Scale	0~50/min
RESP SCALE	0 50711111
ST SUGLE	
IBPI Scale	
IBP2 Scale	U BUUMMHg
Ettuz Scale	0 100mmHg
CIScale	276L/min/m2
Previous	
Set CI scale for short trend.	
Time Scale Select the time interval of short trend diagram. Options are 5min, 10min, 15min, 20min, 30min, 1h and 2h.

HR Scale Select the scale of heart rate for short trend diagram. Options are 0~160/min and 0~300/min.

SpO₂ Scale Select the scale of SpO₂ for short trend diagram. Options are 40~100%, 60~100% and 80~100%.

Resp Scale Select the scale of respiration rate for short trend diagram. Options are 0~8/min, 0~24/min, 0~50/min and 0~100/min.

ST Scale Select the scale of ST-segment for short trend diagram. Options are -2~+2mm, -5~+5mm and -9~+9mm.

IBP1 Scale Select the scale of IBP1 for short trend diagram. Options are **0~300mmHg**, **0~150mmHg**, **0~200mmHg**, **0~100mmHg**, **-20~50mmHg** and **-50~300mmHg**.

IBP2 Scale Select the scale of IBP2 for short trend diagram. Options are **0~300mmHg**,

0~150mmHg, 0~200mmHg, 0~100mmHg, -20~50mmHg and -50~300mmHg.

EtCO₂ Scale Select the scale of EtCO₂ for short trend diagram. Options are 0~30mmHg, 0~60mmHg and 0~100mmHg.

C.I. Scale Select the scale of C.I. for short trend diagram. Options are $2\sim4$ L/min/m², $2\sim6$ L/min/m² and $2\sim8$ L/min/m².

System Setup



Language The categories of languages can be selected. To change the language, it is necessary to restart the monitor.

Recorder Click and open the dialog of recorder configuration.

Time Setup Click and open the dialog of time configuration. After the time of the system has been configured, please restart the monitor.

Mode Config Click and open the dialog of mode configuration.

Alarm Level Click and open the dialog of alarm level configuration.

MachineClick and open the dialog of machine maintenance. Enter the interfaceof machine maintenance and it is necessary to enter the password (password is 125689)

Brightness Set the brightness of monitor screen, the options are 1, 2, 3, 4 and 5.

Recorder Setup

R	ecorder Setup
Record Wave1	п
Record Wave2	Sp02
Record Wave3	002
Record Time	
Interval	OFF
Delay Time	4 S
Record Grid	ON
Alarm Record	OFF
Narn Record	OFF
Manual Time	OFF
Previous	
Set No.1 r	ecord waveform.

Record Wave1 Select the waveform recording in the first line. Select certain waveform to record. **It cannot be switched off.**

Record Wave2 Select the waveform recording in the second line. Select **Off** close the wave display or select certain waveform to record.

Record Wave3 Select the waveform recording in the third line. Select **Off** close the wave display or select certain waveform to display.

Record Time Select the time duration of the waveform for each recording. Options are **8s, 12s** and **16s**.

Interval Select the time interval for cycle recording. Options are Off, 1min, 2min,

3min, 4min, 5min, 10min, 15min, 20min, 25min and 30min.

Delay Time Delayed recordings start documenting on the recorder strip from a preset time before the recording is started. This interval is called the "Delay Time" and can be set to **Real time**, **4s** or **8s**.

Record Grid Enable/Disable recording of the grids when the recorder is producing waveforms.

Alarm Record Enable/Disable the alarm recording at the high level of physiological alarm.

Warn Record Enable/Disable the warn recording at the medium level of physiological alarm.

Manual Time Set the manual recording time. Options are off, 10s, 20s and 30s.

Time Setup



The user can configure system time. The user is advised to set system time before implementing monitoring. If the configuration is to be conducted during the process of monitoring, the user is advised to switch off the monitor after exiting the current window and then restart it. The time for the revision takes effect after the current window is exited.

Mode Setup



Factory Select the default configuration defined by the manufacturer and options are **Cancel, Adult, Children** and **Neonatal**, select $\langle Cancel \rangle$ to abort it.

User Config Select the mode of user saving. Select the previous custom configuration, select $\langle Cancel \rangle$ to abort it.

Save Config Save the current configuration info as custom configuration, enter the name of the user custom configuration, select $\langle OK \rangle$ to save the current mode and select $\langle OR \rangle$ to save the current mode and select

 $\langle Cancel \rangle$ to cancel saving.

Delete Delete the previous data of custom configuration, select the custom configuration that needs to be deleted; press the selected mode to delete the mode, and press $\langle Cancel \rangle$ to cancel deleting.

Q Caution: The mode name cannot be black when saving current configuration, otherwise, the custom configuration will not be save.

Alarm level Setup

Alarm Level	Alarm Level
Alarm Level	Exit
Default	HR Sp02 NIBP Resp
Previous	MED MED MED
	TEMP IBP EtCO2 FiCO2
	MED MED MED
	EtAA FIAA EtN20 FIN20
	OFF OFF OFF OFF
	EtO2 FiO2 ICG
	OFF OFF MED
Set alarm level.	

Alarm levels of all the parameters can be configured. Press **Set Alarm level** > option, the cursor will move to the region of configuring alarm levels. If the alarm level of a certain parameter is to be configured, first move the cursor to the alarm level of that parameter, press the option and then select the alarm level, Options are **low, med** and **high**.



Machine Setup

Maintenance Click and open the dialog of system maintenance

- **Factory** Manufacturer maintenance is not an operation option for users and it must be operated by the technical and maintenance personnel authorized by manufacturer.
- **Upgrade** Click and open the dialog of updating monitor's software.
- Nurse Call Click and open the dialog of nurse call.
- CO2 Function Click and open the dialog of function selection. Options are CO2 Cal Mode, CO2 Gain Cal, CO2 Flow.
- Fun. Select Click and open the dialog of function selection. Options are Alarm Limit, ShortCut Key, ARR Level, ECG BaseLine, ECG Auto and AlarmSound.
- Alarm Setup Click and open the dialog of alarm setup. You can check or set the alarm limit and alarm state.
- **HUM** Select the frequency of the industrial frequency and options are **50Hz** and **60Hz**. It is mainly configured according to the frequency of local power supply.
- **ECG Scale** Select $\langle On \rangle$ or $\langle Off \rangle$ to enable or disable ECG scale.

System Maintenance



Trend Setup Click and open the dialog of trend display configuration. Conduct configurations of trend diagrams and trend tables.

Color Click and open the dialog of color configuration and configure colors of parameters and waveforms.

Network Click and open the dialog of network configuration. Conduct network

configurations.

OverPress Initiate NIBP over-pressure test

Manometer Initiate NIBP manometer test.

NIBP Reset Reset NIBP module.

IBP set Click and open the dialog of IBP setup.

NIBP set Click and open the dialog of NIBP setup.

Recor. Cali. Conduct speed calibration of the recorder. This operation must be conducted when the recorder is changed.

Trend Setup

The user can define various trend display info according to needs or use the display configuration for default trend.



Trend Graph1 Configuration of trend diagram.

There are a total of three pages of trend diagrams and on each page trend diagram can be configured for six regions, and options are Off, HR, SpO₂, NIBP, PR, Resp, CO₂, T1, T2, AA, N₂O, O₂, IBP1, IBP2, ST, HR+SpO₂, SpO₂+PR, Resp+CO₂, PR+CO₂, T1+T2, IBP1+IBP2, AA+CO₂, N₂O+O₂ and C.I. It is possible to have self-configurations on the contents of the trend diagrams and at least one page of trend diagrams shall be configured.

Treed Oriental
Trenu Graphi
Area1 HR
Area2 SpO2
Area3 P1
Area4 P2
Area5 Resp
Area6 OFF
Previous
Set Area i treno gragn parameters.

Trend Table Configuration of trend tables

There are a total of three pages of trend tables and on each page trend table can be configured for six regions, and options are HR, SpO₂, NIBP (S/D), NIBP (M), IBP1 (S/D), IBP1 (M), IBP2 (S/D), IBP2 (M), Resp, PR, T1, T2, CO₂, AA, N₂O, O₂, ST and C.I.. It is possible to have self-configurations on the contents of the trend tables and at least one page of trend tables shall be configured.

	Trend Table1
Area1	HB
Area2	Sn02
Area3	IBP1(S/D)
Area4	IBP2(M)
Area5	Besn
Area6	OFF
Previous	
Set Area1 tre	nd table parameters.

Color Setup



Enter the interface of color configuration, the colors of various parameters and waveforms can be configured.

Network Setup

Network Setup						
IP	192	168	0	134		
Net Mask	255	255	255	0		
Gateway	192	168	0	1		
MacNO	38					
RcvAlmLmt	ON					
Set WID	4					
Previous						
Set IP address.						

In the interface of network configuration, such items as **IP address**, **Net mask**, **Gateway**, **Machine number** can be configured. The configuration is mainly necessary when the monitor connecting to the Central Unit.

Rev Alm Lmt Set the switch of receiving alarm limit from the central unit. It can be

set to ON or OFF.

Set WID Set the wireless ID of the bedside unit.

System Info



Version	It displays the version number of software.
Module SN	It displays the product serial number of module.
Serial Number	It displays the serial number of the machine.
New Board	It displays the type of the new board.

4.2.2 Trend Review

Trend Graph



						PAGE 1/2
Irend Review	TIME	HR	Sp02	IBP1	IBP2m	RR
Page						
Cursor						
Record						
Scale 1 h						
Graph						
Table						
Main						
Return to main screen.						

Trend Table

Page Press this option and turn the trim knob to conduct the paging operation. Press it again to restore the initial status. If more than one page of trend diagrams or trend tables are configured, then the paging is switched between the trend diagrams or trend tables between different pages.

Cursor Press this option, turn the trim knob and move the cursor in the trend diagrams or trend tables. Press it again to restore the initial status. It is possible to move the cursor in the trend diagrams and trend tables. In the trend tables, it is possible to browse the trend records by moving the cursor, and if it moves to the left side or the right side of trend diagram , continue moving can roll the trend diagram by 1/4 screen to the left or right.

Record Press this option to record the trend tables of the current page, but the trend diagram does not support recording.

Scale Press this option and the time intervals for one page of trend diagrams can be selected. Options are 1h, 2h, 4h, 6h, 8h, 10h, 12h, 24h, 48h and 72h and 168h.

Graph Press this option to switch to the display of trend diagram.

 Table
 Press this option to switch to the display of trend tables

4.2.3 Alarm Review

Hiarm Recall	22.1			
2007-05-24 14:21:42	HE LOW			
2067-05-24 14:21:53	SpO2 Low			
ima 2007-05-24 14:21)	05			
IP: 0				
-114				
Sepoll pecond		1		

<</>>> Select this button, turn the trim knob to roll the records back and forth.

1/1 Select this button, turn the trim knob to turn the pages back and forth.

Record Print the currently selected alarm events through the recorder; and if no recorder is configured, this option is invalid.

Exit Exit the dialog of alarm review

4.2.4 ARR Review

ARR RECALL		×
	hadrahadrah	
	~h~h~h~h~h~	
001/128 2007-05-24 1 BRADY	4:22:40	
Exit analys	sis replay (<<]

Click and open the dialog of arrhythmia review and the arrhythmia data for 8 seconds are displayed on each screen, i.e. the ECG waveforms 4 seconds before and after the occurrence of the event, and a maximum of 128 groups of abnormal data can be stored for search.

|<< Turn to the first abnormal waveform record.

>>| Turn to the last abnormal waveform record.

<</>>> Select this button and turn the trim knob to turn the records back and forth.

Record Print the ECG waveform of the current screen through the recorder. If no recorder is configured, this option is invalid.

Exit Exit the dialog of Arrhythmia Review.

4.2.5 Alarm Setup

Alarm Setup
Common Alarm 🗅
IBP Alarm 📑
GAS Alarm 🛄
ST Alarm 🗅
ARR Alarm 🚬
ICG Alarm 🔤
Alarm Record 🛅
Alarm Volume OFF
Previous
Set Common parameters alarm.

Common Alarm Click and open the dialog of common parameters alarm. It can setup the alarm limits of common parameters.

Common Alarm	Adjust Ala	rm			
Adjust Alarm	EXIT				
Default Setup	HR	Sp02	NIBP Sys	NIBP Dia	NIBP Mean
Enable All Previous	0N	0N	0N	OFF	OFF
	40	90	80	40	60
	Pulse	Resp	Temp1	Temp2	
	ON	OFF	OFF	OFF	
	160	60	38.0	38.0	
Alarm limit setup.	40	4	34.0	15.0	

IBP Alarm Click and open the dialog of IBP alarm. It can setup the alarm limits of IBP.



GAS Alarm Click and open the dialog of GAS alarm. It can setup the alarm limits of the GAS module.



ST Alarm Click and open the dialog of ST alarm. If the ST analysis is not configured, this option is invalid.

ST Alarm	Adjust Alarm					
Adjust Alarm	Exit					
Default	I	II	III	aVR		
Enable All	ON	ON	ON	ON		
Previous	0.20	0.20	0.20	0.20		
	-0.20	-0.20	-0.20	-0.20		
	aVL	aVF	V-			
	ON	ON	ON			
	0.20	0.20	0.20			
Alarm limit setup	-0.20	-0.20	-0.20			

ARR Alarm Click and open the dialog of ECG analysis alarm. It can setup the alarm limits of various Arrhythmia.

ARR Alarm	Set Alarn	Param		
Alarm Setup	EXIT			
Default Setup	ASYSTOLE	VENT FIB	PAC	MULTI PVCS
Enable All	OFF	OFF	OFF	OFF
Previous	COUPLET	BIGEMINY	TRI- GEMINY	R ON T
	OFF	OFF	OFF	OFF
	VPB	TACHY	BRADY	MISSED BEATS
	OFF	OFF	OFF	OFF
	PNP	PNC	MULTI PACS	ST ELEVATE
	OFF	OFF	OFF	OFF
Cat alorg lavel	ST EPRESS	NOISE		
Set alarm level.	OFF	OFF		

ICG Alarm Click and open the dialog of ICG alarm. It can setup the alarm limits of the ICG module.

ICG Alarm	Adjust Alarm		
Adjust Alarm	Exit		
Default	C.I.	TFC	
Enable All	ON	ON	
Previous	5.0	60	
	1.5	15	
setup.			

Alarm Record Click and open the dialog of alarm recording. Configure whether the alarm records of various modules are recorded. Only when the switch for alarm recording of the module and the switch for alarm record in the record setup have been switched on, the physiological alarm in the relevant modules will trigger the alarm recording.

Alarm Record	Alarm Re	ecord		
Alarm Record	EXIT			
All OFF	ECG	Sp02	NIBP	Temp
Default Setup	OFF	OFF	OFF	OFF
Previous				
	IBP	C02	GAS	ARR
	OFF	OFF	OFF	OFF
Set record on alarm event.				

Alarm volume Configure the volume of alarm and options are off,1,2 and 3. Once a level is selected, a testing beep will be produced.

Note: In each dialog of alarm configuration, press the button (Adjust Alarm) and the cursor moves to the adjustment region of alarm limits. Press the button (Enable All) and all the alarms will be opened. If the user desires to adjust the alarm parameter of a certain parameter, first move the cursor onto the label of that parameter, and then press the trim knob to move the cursor up and down to select the parameter to be adjusted for revision.

4.2.6 Patient info

Patient Setup
Case No.
Name
Height 165
Weight 50
Sex Female
Age 55
Room No. 1
Bed No. 1
Pace No
Previous
Set patient ID.

Case No. The case number of patients (It can be configured according to the actual status of the hospital and a maximum of 10 letters can be entered), press $\langle Del \rangle$ to delete and $\langle Clear \rangle$ to clear; enter $\langle OK \rangle$ to confirm.

Name Patient name (It can be selected among A-Z and 0-9 and a maximum of 10 letters can be entered) enter $\langle OK \rangle$ to confirm.

Height Body height of patient (Turn the trim knob with an increment or decrement of 1 cm)

Weight Body weight of patient (Turn the trim knob with an increment or decrement of 1 kg)

Sex Gender of patient (male or female)

Age Age of patient (Turn the trim knob with an increment or decrement of 1 year)

Room No. Number of patient's room. Patient's room number can be displayed in the central unit.

Bed No. Number of patient's bed. Patient's bed number can be displayed in the central unit.

Pace Change the paced status. For paced patients, set [Pace] to [Yes]. For non-paced patients, set [Pace] to [No].

4.2.7 Drug Dose Calc

Select $\langle MENU \rangle \rightarrow \langle Calculator \rangle \rightarrow \langle Drug Calc \rangle$, enter the Drug dose calculation window. As follows:



This calculation of drug concentration is mainly aimed at facilitating the work of physicians. It conducts concentration calculation on some commonly used drugs. A content of titration table can be output through recorder.

In the system, the following categories of drugs can be calculated:

AMINOPHYLLINE, DOBUTAMINE, DOPAMINE, EPINEPHRINE, HEPARIN, ISUPREL, LIDOCAINE, NIPRIDE, NITROGLYCERIN, and PITOCIN. In addition, it provides DRUG_A, DRUG_B, DRUG_C, DRUG_D and DRUG_E to displace any other drugs flexibly.

The following formulas are used for the calculation of drug dosage: Drug concentration equal to total amount of drug divided by liquid volume Liquid velocity equal to drug dosage divided by drug concentration Duration time equal to total amount of drug divided by drug dosage Drug dosage equal to velocity of IV drip multiply drug concentration In the window of drug calculation, the operator should first select the name of the drug to be calculated, confirm the patient weight and then enter other known values.

Drug name

Move the cursor to $\langle Drug name \rangle$, press the trim knob, then turn the trim knob to select drug, and only one kind of drug can be selected for calculation at one time.

DRUG_A, DRUG_B, DRUG_C, DRUG_D and DRUG_E are only codes for drugs rather than their real names. The units for these five kinds of drugs are fixed and the operator can select the appropriate units according to the habits of the drugs. The rules of the units are as follow:

DRUG_A, DRUG_B, DRUG_C are fixed at the serial units of gram (g), milligram (mg) and microgram (mcg).

DRUG_D is fixed at the serial units of **unit**, **k unit** and **m unit**.

DRUG_E is fixed at the unit of **mEq**.

Weight

The operator should enter the patient weight first, and as independent info the weight is only used in the function of the calculation of drug concentration.

Turn the trim knob to move the cursor to the positions of the various calculation items in the calculation formula respectively, turn the trim knob, and select calculation value, then press the trim knob and confirm the selected calculation value. When the calculation value is selected, the value of the calculated item will be displayed at relevant locations. There are range limits for the value adoption of each calculation Item, if the calculation results exceed the range, "---"will be displayed.

Regarding this function of drug calculation, the values for other individual items can only be entered after the weight and drug name have been entered. In the system, the values that are given initially are only a group of random initial values and the operator shall not take this value as the calculation standard and a group of values appropriate to the patient must be reentered according to the physicians' comments.

Each kind of drugs has a fixed unit or unit series and the operator must select the appropriate units according to the physicians' comments. In the unit series of the same unit, the addition of the units will be automatically adjusted in accordance with the current entered value. When the expressed range that can be expressed by this unit is exceeded, the system will display "---".

When the operator has entered the value of a certain item, the system will give a prompt in the menu so as to remind the operator to verify the correctness of the entered value. Only by ensuring the correctness of the entered values, the calculated values can be reliable and safe.

In case of neonatal, drip velocity and volume per drip are invalid.

The values in the table may not be related to the patient monitored on this bed. Therefore the weight of this menu and the weight in the patient info are two different values. The values in this menu item are not affected by the values in the patient info.

Titration table

Select (**Titration**) in the menu of drug calculation to enter the interface of titration table.

Titrat	tion Tab	le				>
DRUG AM Dose/hr WEIGHT	10UNT	400.00 150.00 70.0 ⊧	mg Mg (g	LIQUID INF RAT DRIP RA	VOL 25 E 93 TE 31	0.00 m1 .75 m1/hr .25 GTT/min
DOSE	INF	RATE	DOSE	INF RATE	DOSE	INF RATE
0.00 1.00 2.00 3.00 4.00 5.00 6.00 7.00 8.00 9.00	0.0 0.6 1.2 1.8 2.5 3.1 3.7 4.3 5.0 5.6	0 3 5 3 0 3 5 3 0 3	10.00 11.00 12.00 13.00 14.00 15.00 16.00 17.00 18.00	6.25 6.88 7.50 8.13 8.75 9.38 10.00 10.63 11.25 11.88	20.00 21.00 22.00 23.00 24.00 25.00 26.00 27.00 28.00 29.00	12.50 13.13 13.75 14.38 15.00 15.63 16.25 16.88 17.50 18.13
BASE	DOSE PAGE	UP/DOWN	STEP		E TYPE RECO	Dose/hr 👤 RD
Select	t one as	the inp	out and ca	loulate the o Exit	other one	

In the titration table, turn the trim knob to $\langle Base \rangle$, then press the trim knob to select the desired item. Options are **Dose, Trans speed** and **Drop speed**. After selecting, press the trim knob to confirm the selection.

Move the cursor to \langle **Step** \rangle and press the trim knob to select the step size; the selectable range is 1-10.

Move the cursor to $\langle Dose Type \rangle$ and press the trim knob to select the dosage unit.

Move the cursor to $\langle Page Up / Down \rangle$, press the trim knob, and then turn the trim knob to browse the previous page and next page.

Move the cursor to $\langle \mathbf{Record} \rangle$, press the trim knob to give the output of the data of the titration table on the currently displayed interface.

Move the cursor to \langle **Exit** \rangle , press the trim knob to return to the window of drug calculation.

4.2.8 Hemodynamic Calculation

Calculation Process

1. Select $\langle MENU \rangle \rightarrow \langle Calculator \rangle \rightarrow \langle Hemodynamic \rangle$, enter the Hemodynamic Calculation window. As follows:



Please enter the parameter correctly

a. If you are calculating the current patient, the monitor can get C.O., HR, Height and Weight automatically. You need to put in the other parameters. The way to input parameters is firstly pitching on the case of the parameter, then turn the knob to enter its value, lastly press the knob to finish. You can turn the knob to the next parameter input.

b. If you are not calculating the current patient, no parameter's value will be offered by the monitor, you need to input them by yourself.

(Calc) Press the button, you will obtain the result window.

 $\langle Exit \rangle$ Press the button, you will close the current window.

2. After you have finished the data input, please make sure they are correct. Then you can press the button $\langle Calc \rangle$ to get the following window

CI	4.4	L/min/m ²	BSA	1.757	m ²
SV	114.9	mL	SI	65.4	mL/m ²
SVR	747.7	dyn•s/cm5	SVRI	1313.8	dyn•s•m²/cm5
PVR	882.7	dyn•s/cm ⁵	PVRI	1551.0	dyn-s-m²/cm5
LCW	8.7	kg•m	LCWI	4.9	kg•m/m ²
RCW	9.84	kg•m	RCWI	5,60	kg•m/m ²
LVSW	129.7	g•m	LVSWL	73.8	g•m/m ²
RVSW	146.92	g•m	RVSWI	83.61	g•m/m ²
EF	100.81	%			

- $\langle 4/4 \rangle$ Press the button, then turn the knob to flip the page. The former "4" shows the current page, the back one shows the number of the total pages.
- (Range/Unit) Press the button, the unit of the parameter will disappear, and the unit of the parameter in red word will be changed into the logical range of them. Press the button again, the unit of them will be displayed.
- $\langle Record \rangle$ Press the button, the current page will be printed out.
- **(Show Input)** Press the button, the corresponding input of current result will be present.
- $\langle Exit \rangle$ Press the button to quit the current result window.

Note: The red words mean the parameters' values are beyond the logical range. After the output parameter, the sign of "---"will appear sometime, it means the parameter is invalid.

Abbreviation	Unit	Full Name
C.O.	L/min	Cardiac output
HR	bpm	Heart rate
PAWP	mmHg	Pulmonary artery wedge pressure
Art Mean	mmHg	Mean artery pressure
PA Mean	mmHg	Mean pulmonary artery pressure
CVP	mmHg	Central venous pressure
EDV	ml	End diastolic volume
Height	cm	Height
Weight	kg	Weight

Input Parameter

Output Parameter

Abbreviation	Unit	Full Name
C.I.	L/min/m ²	Cardiac index
BSA	m ²	Body surface area
SV	ml	Stroke volume
SI	ml/m ²	Stroke index
SVR	DS/cm ⁵	Systemic vascular resistance
SVRI	DS·m ² /cm ⁵	Systemic vascular resistance index
PVR	DS/cm ⁵	Pulmonary vascular resistance
PVRI	DS·m ² /cm ⁵	Pulmonary vascular resistance index
LCW	kg∙m	Left cardiac work
LCWI	kg·m/m ²	Left cardiac work index
LVSW	g∙m	Left ventricular stroke work
LVSWI	g·m/m ²	Left ventricular stroke work index
RCW	kg∙m	Right cardiac work
RCWI	kg·m/m ²	Right cardiac work index
RVSW	g∙m	Right ventricular stroke work
RVSWI	$g \cdot m/m^2$	Right ventricular stroke work index
EF	%	Ejection fraction

4.2.9 HRV analysis

Heart rate variability is the tiny difference between successive heart rate interphase. It comes into being from the modulation of autonomic tones on the sinus nodal autonomy. And it makes several milliseconds difference and fluctuation between heart rate interphase. HRV analysis is a sensitive and non-invasive method to estimate the function of autonomic tones.

Select $\langle MENU \rangle \rightarrow \langle Calculator \rangle \rightarrow \langle HRV Analyze \rangle$, enter the HRV analysis window. (1) R-R interphase value histogram: It reflects the distribution of R-R interphase value. The abscissa is the R-R interphase, and the ordinate is the numbers of R-R interphase. Histogram can reflect the range and extent of R-R interphase variety. The R-R interphase value histogram is as follows:



(2) R-R interphase difference histogram: It reflects the difference of R-R interphase, namely, the extent of sinus arrhythmia. The abscissa is the dispersion of R-R interphase of sinus beat. The midpoint is zero. The R-R interphase difference histogram is as follows:



(3) Scatter: The abscissa is the R-R interphase of previous heart rate. And the ordinate is the R-R interphase of next heart rate. The rest may be deduced by analogy. The X-axis is RRn. The Y-axis is RRn+1. Each heart rate points make the R-R interphase scatter. The scatter displays the linearity and nonlinear variety trend of HRV. According to the scatter, you can see the state of autonomic tones function. The scatter is as follows:



(4) R-R trend graph: It is the summarizing of all kinds of indexes after analyzing the data in a period of time. It reflects its variety. The abscissa is time. And the ordinate is the average of R-R interphase in a period of time. The R-R trend graph is as follows:



Prote: HRV analysis can not be used on patient with pacemaker.

4.2.10 Ambulatory blood pressure analysis

Ambulatory blood pressure monitoring is a technology that measures blood pressure in daily life automatically and discontinuously. It reflects the level, difference and rhythm of blood pressure.

Select $\langle MENU \rangle \rightarrow \langle Calculator \rangle \rightarrow \langle NIBP Analyze \rangle$, enter the ambulatory blood pressure analysis window:

NIBP Analyze
Time Length 24 h
Time Inter. 5 min
Start Anal.
Anal.Result 🖣
Previous
Time Length

- (Time Length) : Set the time length of blood pressure monitoring. Options are "24h, 48h";
- (Time Inter.) : Set the time interval of NIBP analysis, Options are "5 min,10 min,15 min,30 min";
- **(Start Anal.)** : Click it to begin ambulatory blood pressure analysis;
- 〈Anal. Result 〉: Click it to display the NIBP analysis result. The NIBP analysis window is as follows:



4.3 Screen display

This Monitor adopts color LCD screen with high brightness, which can display parameters, waveforms, system status and other prompt info. The main screen is mainly divided into three regions, they are respectively:

- Display zone of system info and alarm prompt info (the uppermost part)
- Waveform display zone (left, and It shall vary according to different screen types)
- Parameter display zone (right and lowest part)

4.3.1 System status

The system time and status of battery capacity are displayed on the upper right corner.



Notes on battery capacities:



Battery capacity is exhausted

Only when the monitor is powered by battery and is recharging the battery, the icon for battery capacity is displayed. If AC power in current use and the battery capacity is full, the icon will not be displayed.

Note: When the battery capacity is exhausted, the system produces an alarm sound, prompting the user to plug in the AC power for recharging; if it is not recharged in time, the monitor will be automatically switched off due to insufficient capacity more than 5 minutes.

Caution: When the energy level of the battery is exhausted, plug in the AC power to recharge, and then the battery indication may quickly return to "Full battery level"; the AC plug should be plugged in so as to ensure the full capacity of the battery.

4.3.2 Info display region

The upper region of the screen is the info display region, which is used to display the status of alarm sound, alarm suspension countdown and alarm info.

Status of alarm sound

The alarm sound is in "Off" status, and if a new alarm is generated, the "Off" status of alarm sound will be automatically cancelled.

Pause the alarm, and if a new alarm is generated, the "Pause" status of alarm sound will be automatically cancelled.

Alarm indicating zone



Alarm levels

Red base color is high alarm

Yellow base color is medium and low alarm

The order displayed by the physiological parameter alarm is displayed from left to right in turn according to the alarm levels.

Parameter alarm

The value of that parameter displayed on the upper part of the screen will flash to indicating the alarm of that parameter.

Chapter 5 Parameters Measurement

5.1 Measurement of ECG/HR

5.1.1 Principles of Measuring

Before the mechanical contraction, the heart will firstly produce electrization and biological current, which will be conducted to body surface through tissue and humors; the current will present difference in potential in different locations of the body, forming potential difference ECG, also known as body surface ECG or regular ECG, is obtained by recording this changing potential difference to form a dynamic curve. Monitor measures the changes in the body surface potentials caused by the heart of the patient, observes the cardioelectric activities, records the cardioelectric waveforms and calculates the HR through the multiple electrodes connected to ECG cable.

5.1.2 Precautions during ECG Monitoring

Warning: Before connecting the ECG cables to the monitor, please check if the lead wires and cables have been worn out or cracked. If so, they should be replaced.

Warning: It is imperative to only use the ECG cables provided with the instrument by manufacturer.

Warning: The equipment is capable of displaying the ECG signal in the presence of pacemaker pulses without rejecting pacemaker pulses.

Warning: To avoid burning, when the electrotome operation is performed, the electrodes should be placed near the middle between ESU grounding pad and electrotome and the electrotome should be applied as far as possible from all other electrodes, a distance of at least 15 cm/6 in is recommended.

Warning: When the electrotome operation is performed, the ECG leadwires should be intertwisted as much as possible. The main unit of the instrument should be placed at a distance from the operation table. Power wires and the ECG lead cables should be partitioned and should not be in parallel.

Warning: The monitor is protected against defibrillation effect. When applying defibrillator to the patient, the monitor will experience transient disorderly waveforms. If the electrodes are used and placed correctly, the display of the monitor will be restored within 5 seconds. During defibrillation, the chest leads such as V₁~V₆ should be removed and such limb electrodes as RA, LA, RL, LL should be moved to the side of the limbs.

Warning: All the electrodes and conducting part shall not be into contact with any other conductors including the ground. For the sake of patient safety, all the leads on the ECG cables must be attached to the patient.

Warning: When conducting defibrillation, it is imperative to only use the electrodes recommended by manufacturer.

Warning: Do not come into contact with the patient, bed and the monitor during defibrillation.

Warning: The monitor cannot be directly applied to heart and cannot be used for the measurement of endocardio ECG.

Solution Note: When several parts of equipment are interconnected, the total leakage current is limited to the safety range according to standards IEC 60601-2-27.

5.1.3 Preparatory Steps before the Measurement of ECG/HR

1) Plug the ECG cable into the ECG socket of the monitor.

2) Place the electrodes onto the body of the patient and connect them to the relevant lead wires of the ECG cables, and at this moment ECG waveforms will appear on the screen.

3) Set the parameters relevant to ECG monitoring.

5.1.4 Connecting the ECG Cables to the Monitor

Monitor is provided with three different ECG cables relevant to 3-Lead ECG module, 5-Lead ECG module and 12-Lead ECG module:



Fig. 5-1-4 Connect the ECG cable to the monitor

1) 3-lead ECG cable

- Including three limb leads: RA, LL, and LA.
- Relevant ECG socket ECG1 (It can only be connected to this socket and can not

be connected to ECG2).

- Realize 3-lead ECG monitoring.
- 2) 5-lead ECG cable
 - Including four limb leads: RA, RL, LL, LA and one chest-lead V (V₄).
 - Relevant ECG socket ECG1 (It can only be connected to this socket and can not

be connected to ECG2).

- Realize 7-lead ECG monitoring.
- 3) Chest-lead ECG cable
 - \blacksquare Including five chest leads: V₁, V₂, V₃, V₅, V₆.

■ Relevant ECG socket ECG2 (It can only be connected to this socket and can not be connected to ECG1).

■ Its joint use with 5-lead ECG cable can realize all-lead (12-lead) ECG monitoring.

5.1.5 Connecting the ECG Electrodes to the Patient

1) Connection steps

 \blacksquare Clean the patient's skin and wipe the oil stains, sweat stains on the skin with alcohol. If necessary, shave body hair at the locations where the electrodes are to be placed or grind off the stratum corneum and clean it with alcohol.

■ Check if the buttons on the electrodes are clean and free of damage.

■ Place the electrodes on the body of patient. Before attaching, smear some conducting cream on the electrodes if the electrodes are not electrolyte self-supplied.

■ Connect the cable leads to the electrodes through the buttons of the electrodes.

Solution Note: For patients who tremble a lot or patients with especially weak ECG signals, it might be difficult to extract the ECG signals, and it is even more difficult to conduct HR calculation. For severely burnt patients, it may be impossible to stick the electrodes on and it may be necessary to use the special pin-shape electrodes. In case of bad signals, care should be taken to place the electrodes on the soft portions of the muscle.

Solution Note: Check the irritation caused by each electrode to the skin, and in case of any inflammations or allergies, the electrodes should be replaced and the user should relocate the electrodes every 24 hours or at a shorter interval.

Proof Note: When the amplifier is saturated or overloaded, the input signal is medical meaningless, then the equipment gives an indication on the screen.

2) Location for electrode placement



Fig. 5-1-5 Indicative map of the placement of ECG electrodes

AHA Label	AHA Color	IEC Label	IEC Color	Location
RA	White	R	Red	Under the clavicle of the right shoulder.
LA	Black	L	Yellow	Under the clavicle of the left shoulder.
RL	Green	Ν	Black	Right lower abdomen.
LL	Red	F	Green	Left lower abdomen.
V1	Red	C1	Red	4th intercostal space on the right sternum side.
V2	Yellow	C2	Yellow	4th intercostal space on the left sternum side.
V3	Green	C3	Green	Center of the line connecting V_2 and V_4 .
V4	Blue	C4	Brown	Node of the left 5th intercostal space and the mid-clavicular line.
V5	Orange	C5	Black	Node with the left anterior axillary line at the same height with V_{4} .
V6	Purple	C6	Purple	Node with the left mid-axillary line at the same height with V_{4} .

The following table shows the lead name to identify each lead wire and its associated color of AHA and IEC standards.

When conducting 3-leads ECG monitoring, use 3-lead ECG cable. The three limb-leads of RA, LA and LL as shown in Fig. 5-1-5, will be placed on the relevant locations. This connection can establish the lead of I, II, III.

When conducting 7-leads ECG monitoring, use 5-lead ECG cable. The four limb-leads of RA, LA, RL and LL as shown in Fig. 5-1-5, will be placed on the relevant locations. This connection can establish the lead of I, II, III, aVR, aVL, aVF; according to actual needs, chest lead V can be placed on any of the locations between $V_1 \sim V_6$, respectively making one lead of $V_1 \sim V_6$ established.

When conducting 12-leads ECG monitoring, use the 5-lead ECG cable and chest-lead ECG cable at the same time, and all the leads are placed on the relevant locations respectively as indicated by Fig. 5-1-5, among which the chest lead V of the 5-lead ECG cable is placed on the location of V₄. This kind of connection realizes the establishment of such 12 leads as I, II, III, aVR, aVL, aVF, V₁~V₆.

5.1.6 Setup of ECG/HR parameters



ECG1 Select the ECG waveform, and this lead is the key monitoring lead.

ECG Gain Select the gain item of ECG waveform, and options are AUTO, 0.25x, 0.5x, 1.0x, 2.0x and 4.0x.

HR Source Select HR source item, and common options are AUTO, ECG, PLETH.

When select ART for IBP measurement, the option **ART** is appeared in HR source.

Beep Volume Select the volume of BEEP, and options are **Off**, **1**,**2** and **3**. Once an option is selected, a testing beep will be produced.

Alarm Setup Click and open the dialog of alarm setup.

ECG Setup Click and open the dialog of ECG setup.

ECG Replay Click and open the dialog of ECG replay.

ARR Setup Click and open the dialog of Arrhythmia analysis configuration, Configure some of the parameters of Arrhythmia analysis.

ST Adjust Click and open the dialog of ST adjustment. In the pop-up window you can adjust the position of ST point and ISO point.

• Alarm setup

ST Alarm ARR Alarm Previous	
ARR Alarm	
Previous	
Set HR alarm.	

ECG alarm Click and open the dialog of HR alarm



Adjust alarmSelect this option to enter the configuration of alarm limits and configurethe limits by turning the trim knob to select the high limits and low limits, and exit byselecting $\langle EXIT \rangle$. The upper part is the high limit and the lower part is the low limit.HR alarmSelect $\langle ON \rangle$ to enable HR over limit alarm; select $\langle OFF \rangle$ to disable HRover limit alarm.
ST Alarm	Adjust A	Alarm		
Adjust Alarm	Exit			
Default	I	ĪĪ	III	aVR
Enable All	ON	ON	ON	ON
Previous	0.20	0.20	0.20	0.20
	-0.20	-0.20	-0.20	-0.20
	avL	aVF	V-	
	ON	ON	ON	
	0.20	0.20	0.20	
Alarm limit setup	-0.20	-0.20	-0.20	

ST alarm Click and open the dialog of ST alarm.

ST Alarm Select **<ON>** to enable ST over limit alarm; select **<OFF>** to disable ST over limit alarm;

Adjust alarm Select this option to enter the configuration of alarm limits; by turning the trim knob, select the high limits and low limits for configuration and exit by selecting

 $\langle Exit \rangle$. The upper part is the high limit and the lower part is the low limit;

Default Select this option and configure the ST alarm as the manufacturer's default configuration;

Enable All Select this option and configure all the ST alarms as low level alarms.

ARR alarm Click and open the dialog of Arrhythmia analysis alarm.



In the interface, it is possible to configure the alarm levels of various Arrhythmia or switch off the Arrhythmia alarm.

Alarm Setup Select this option to enter the configuration of ECG alarm levels; select the alarm levels of different Arrhythmia by turning the trim knob and exit by selecting <EXIT>.

Default Setup Select this option and configure the Arrhythmia alarm as the manufacturer's default configuration.

Enable All Select this option and configure all the Arrhythmia alarms as low level alarms.

• ECG Setup



Lead Type Select the lead type of ECG input, and options are 5 leads, 3 leads, Auto and 12 leads.

Scan Speed Select the scanning speed of ECG waveforms and options are 12.5mm/s, 25mm/s and 50mm/s. The output speed of the recorder remains the same as the scanning speed of the ECG lead.

MODE Select monitoring mode, and options are USER, DIAG, MON and OPS.

Resp Lead Select the calculation methods of RESP lead, and options are **RA-LL**, **RA-LA**, **RL-LA** and **RL-LL**.

DRIFT Select the modes of drift filtrations, and options are **Off**, **Drift 1** and **Drift 2**.

EMG Select myoelectric filtration, and options are Off, 25Hz and 40Hz.

HUM Select hum frequency filtration, and options are **Off** and **on**. Specific frequencies (50HZ, 60HZ) are configured in 〈**Machine**〉 and they must be configured according to the

frequency of local power supply.

Display PR Select to simultaneity display pulse rate. If simultaneity display of PR is selected, PR will be simultaneity displayed at the lower left corner of the ECG parameter display region.

Filter ECG mode	Drift filter	HUM filter	EMG filter
DIAG	OFF	OFF	OFF
OPS	Drift 2	50Hz/60Hz	25Hz
MON	Drift 1	50Hz/60Hz	40Hz
USER	Optional	Optional	Optional

The states of the filter under various modes of ECG

Note: Under the mode of DIAG, OPS and MON, the state of the filter cannot be regulated. Only under the state of USER can the state be regulated.

Caution: When "3 Lead" is selected as <Lead Type>, ECG is in 3-lead input mode, and only Lead I, II or III can be measured.

Q Caution: When "5 Lead" is selected as <Lead Type>, ECG is in 5-lead input mode, and Lead I, II, III, aVR, aVL and aVF and one chest lead can be measured at the same time; if chest lead ECG cable is connected, V1~V6 can be measured at the same time.

• ECG replay

ECG Replay
Start Save
Hist.Replay 🛼
Curri Replay 🐚
Previous
Current ECG waveform replay.

Start Save: Click it to save the waveform. Click it again to stop saving the waveform.

Hist. Replay: Enter history replay window.

Curr. Replay: Enter current replay window showed as follows:

Current Replay	×
Time: 2007-06-07 10:27:36 2007-06-07 10:27:40	
Nove forward/backward waveform. ((/>) 1 / 1 Record 1	Exit

<</>>> Select this button and it is possible to roll the waveform block by turning the trim knob back and forth, with 5 seconds each block.

1/1 Select this button, and it is possible to turn the pages back and forth, and the number before "/" shows the current page and the number following "/" shows total page numbers.

Record Print the enlarged waveform in current selection through the recorder.

Exit Exit the dialog of ECG replay.

• ARR Setup

	ARR analysis
ARR	ON
ST	ON
PACEMAKER	OFF
PVCs	10
	10
ST High	0.10
ST Low	-0.10
ARR Review	
Arr Learn	
Previous	
Arr reLear	ning.

ARR Only when $\langle \mathbf{ON} \rangle$ is selected, the monitor will conduct Arrhythmia analysis **ST** Only when $\langle \mathbf{ON} \rangle$ is selected, the monitor will conduct ST-segment analysis.

Pacemaker Only when (ON) is selected, the monitor will conduct pace-making analysis on patient with pacemaker.

PVCs Only when the times of continuous occurrences is selected from 1 to 10, the monitor will set off the alarm for the frequent occurrences of ventricular premature contractions.

PACs Only when the times of continuous occurrences is selected between 1 and 10, the monitor will set off the alarm for the times of premature beat.

ST High ST alarm high limit of arrhythmia analysis, and the unit is **mV**.

ST Low ST alarm low limit of arrhythmia analysis, and the unit is mV. In arrhythmia analysis, only when the ST-segment exceeds the configured high and low limits will be regarded as the elevation or depression of the ST-segment. They are different from the high and low limit in the configuration of ST alarm limits.

ARR Review Click and open the dialog of arrhythmia review. The user can review the arrhythmia that have occurred and can browse the waveforms 4 seconds before and after the occurrence of arrhythmia alarm.

ARR Learn Click it and you can start the arrhythmia relearning manually.

• ST Adjust

The ST value for each beat complex is the vertical difference between the ISO point and the ST point, as shown in the figure below. The isoelectric (ISO) point provides the baseline, the ST point is at the midpoint of the ST segment. The J point is where the QRS complex changes its slope; as it is a fixed distance away from the ST point, it can be useful to help you position the ST point correctly.



Isoelectric point ST Based Point

In the ECG menu, select \langle **ST Adjust** \rangle , the QRS complex will be displayed on the popped up window, the two vertical lines indicate the position of ISO point and ST point. Set the R wave peak as the reference point for ST measurement.

The ST measurement points and ISO point need to be adjusted when you start monitoring, and if the patient's heart rate or ECG morphology changes significantly.

You can select $\langle ISO \rangle$ or $\langle ST \rangle$ and then adjust the point right and left.

Q Caution: When conducting ST analysis, the system will not consider abnormal QRS complex.

5.1.7 Functions of Arrhythmia Analysis

Provide an analysis software module may be an optional function in your monitor.

The function is consisting of ST segment analysis and arrhythmia analysis. Arrhythmia analysis can identify more than 18 kinds of abnormal ECG, as listed in the following table:

Arrhythmia Type	Applicable Patient Type	Alarm Level
ASYSTOLE	All	Default High, User selectable
VENT FIB/TACHY	Non-paced	Default High, User selectable
MULTI PVCS	Non-paced	Default medium, User selectable
COUPLET	Non-paced	Default medium, User selectable
BIGEMINY	Non-paced	Default medium, User selectable
TRIGEMINY	Non-paced	Default medium, User selectable
RON T	Non-paced	Default medium,User selectable
VPB	Non-paced	Default medium, User selectable
ТАСНҮ	All	Default medium, User selectable
BRADY	All	Default medium,User selectable
MISSED BEATS	Non-paced	Default medium,User selectable
ST HIGHER	All	Default medium, User selectable
ST LOWER	All	Default medium, User selectable
PNP	Paced	Default medium, User selectable
PNC	Paced	Default medium, User selectable
NOISE	All	Default medium, User selectable
VTACHY	Non-paced	Default High, User selectable
PVCS HIGHER	Non-paced	Default medium, User selectable

5.1.8 Maintenance and Cleaning

If there is any sign that the ECG cable may be damaged or deteriorated, replace it with a new one instead of continuing its application on the patient.

To avoid extended damage to the equipment, disinfection is only recommended when stipulated as necessary in the hospital maintenance schedule, disinfection facilities should be cleaned first.

■ Cleaning:

Use a piece of clean cloth moistened in water or mild soap solution to clean the ECG cable.

Disinfection:

Use a piece of clean cloth to wipe the surface of the cable with a 10% bleach solution or

2% Cidex®, clean with clear water and wipe it dry.

5.2 Measurement of RESP

5.2.1 Principles of Measuring

Monitor measures RESP with the method of impedance. When a patient exhales and inhales, changes will take place in the size and shape of the thoracic cavity, causing consequent changes in the impedance between the two electrodes installed at the patient's chest. Based on the cycle of impedance changes, the respiration rate can be calculated.

5.2.2 Preparatory Steps of the Measurement of RESP

1) Plug the 5-lead ECG cable into the ECG socket of the monitor.

2) Place the various pads of the electrodes onto the body of patient and connect them to the relevant lead cables. At this moment, the screen will show RESP waves and the RESP rate will be calculated.

3) Set the parameters relevant to RESP monitoring.

5.2.3 Connect the ECG Cable with Patient and the Monitor

To measure RESP parameters, it is unnecessary to use other cables and it is only necessary to use the two RA and LL leads in the 5-lead ECG cable. So please refer to Fig. 5-1-4 to plug the 5-lead ECG cable into the CH1 ECG socket and refer to Fig. 5-1-5 to place the RA and LL leads onto the body of patient.

Warning: For the sake of safety, all the leads on the 5-lead ECG cable must be connected to the body of patient.

Q Caution: In order to get the best RESP waveforms, when selecting lead II for measuring RESP, it is advised to place RA and LL electrodes cornerways.

Caution: For reducing the influence of rhythmic blood flow on Resp electrode pickup impedance changes, avoid the liver area and ventricles of heart in the line between RA and LL electrodes. This is particularly important for neonates.

Caution: The measurement of RESP is not applicable for patient with excessive motion, otherwise it may cause the mistake of RESP alarm.

5.2.4 Setup of RESP parameters



Scan Speed Select the scanning speed of RESP waveform, and options are 6.25mm/s, 12.5mm/s and 25mm/s.

Resp Gain Select the waveform gain, and options are **0.25x**, **1x**, **2x** and **4x**.

Resp Source When the system is configured with CO_2 module, RESP source can be selected as **AUTO**, **ECG** and **EtCO**₂. Only when the monitor that user has bought has CO_2 module, **EtCO**₂ of RESP source is valid, otherwise the RESP source is defaulted as **ECG**.

Apnea Alarm Suffocation alarm occurs when the time of zero RESP rate has reached this time scale, the alarm will be set off. Options are **Off**, **10s**, **20s**, **40s** and **60s**.

Resp Alarm Click and open the dialog of Resp alarm configuration.

Resp Lead Select the calculation methods of Resp lead, and options are RA-LL,

RA-LA, RL-LA and RL-LL.

Resist Drift Select **<ON>** to enable resist drift; select **<OFF>** to disable resist drift.

• Resp Alarm



Adjust alarm Select this option to enter the configuration of alarm limits; conduct the configurations by turning the trim knob to select high or low limits and exit by selecting $\langle EXIT \rangle$. The upper part is the high limit and the lower one is the low limit.

RESP alarm Select **<ON>** to enable RESP over limit alarm; select **<OFF>** to disable RESP over limit alarm.

5.2.5 Maintenance and Cleaning

No special operation demanded. Please refer to chapter 5.1.8.

5.3 Measurement of SpO₂/Pulse

5.3.1 Principles of Measuring

The measurement of degree of blood oxygen saturation (also known as pulse oxygen saturation, usually shortened as SpO_2) adopts the principles of light spectra and volume tracing. The LED emits lights with two specific bandwidths, which are selectively absorbed by hemoferrum and desoxyhemoglobin. The optical receptor measures the changes in the light intensity after the light passes the capillary network and estimates the ratio of hemoferrum and the total hemoglobin.

Degree of pulse oxygen saturation %= $\frac{\text{hemoferrum}}{\text{hemoferrum + desoxyhemoglobin}} \times 100\%$

The sensor measurement wavelengths are nominally 660nm for the Red LED and 940nm for infrared LED.

Abnormal hemoglobin, carboxyhemoglobin, oxidative hemoglobin are not directly measured, for they are not the affecting factors in the measurement of SpO_2

Monitor adopts FFT filter and signal correlation techniques to deal with SpO₂ module's pulse waveform signals. Before the measurement of SpO₂, the noise produced in the false trace is smoothed so as to the eliminate disturbance in the measurement of saturation. In case of weak blood pulse, the noise produced by some confinements of electrical properties is greatly reduced.

The monitor is designed for measurement and recording of functional saturation.

When you use Masimo SpO₂ module, the Perfusion Index (PI) and Pleth Variability Index (PVI) can be measured and displayed on the screen.

■ Perfusion index (PI): PI is a value that indicates arterial pulse signal strength as the percentage of pulsatile signal to non-pulsatile signal. The perfusion index allows clinicians to place sensors on optimal sites.

■ Pleth Variability Index (PVI): PVI is a measure of peripheral perfusion changes secondary to respiration, or the PI amplitude modulation over a respiration, and can be closely related to intrathoracic pressure changes.

5.3.2 Preparatory Steps before the Measurement of SpO2/Pulse

1) Plug the SpO₂ sensor cable into the SpO₂ socket of the monitor.

2) Put the SpO₂ sensor onto the finger of the patient, and the screen should display SpO₂ waveforms, and the SpO₂ value and pulse rate should be calculated.

3) Set up the parameters relevant to SpO_2 and pulse monitoring.

5.3.3 Connecting to Patient and Monitor

Refer to Fig.1-5-1 and plug the SpO_2 sensor cable into the socket marked with SpO_2 , then put the sensor onto the finger of the patient, as shown in Fig. 5-3-3.



Fig. 5-3-3 Connection of SpO₂ sensor with the patient

After the SpO_2 sensor is connected to the patient, the screen shall display SpO_2 waveforms and then it shall calculate the SpO_2 value and pulse rate value.

Caution: In case it is necessary to add a clip to fix the fingertip sensor, the cable instead of the sensor itself should be clipped. Please note that the cable of sensor should not be pulled with force.

Solution Note: Frequent movements of the sensor may result in errors in the readings of the monitor.

Warning: In case NIBP and SpO₂ are measured at the same time, please do not place the SpO₂ sensor and the NIBP cuff on the same end of the limb, for the measurement of NIBP will block blood flow, affecting the measurement of SpO₂.

Warning: Do not conduct SpO₂ measurement on the finger smeared with fingernail oil, otherwise unreliable measurement results might be produced.

P Note: When using SpO₂ sensor, care should be taken to shield external light sources, such as light of thermo therapy or ultraviolet heating light, otherwise the measurements may be disturbed. Under such conditions as shock, hypothermia, anemia or the use of blood vessel-activating drugs, and with the existence of such substances as carboxyhemoglobin, methemoglobin, methylene blue the result of the SpO₂ measurement will be possibly not accurate.

P Note:

- Make sure the nail faces to the light window.
- The wire should be on the backside of the hand.
- SpO₂ waveform is not proportional to the pulse volume.

Warning: Do not use the sterile supplied SpO₂ sensors if the packing or the sensor is damaged and return them to the vendor.

Warning: Prolonged and continuous monitoring may increase jeopardy of unexpected change of dermal condition such as abnormal sensitivity, rubescence, vesicle, repressive putrescence, and so on. It is especially important to check the sensor placement of neonate and patient of poor perfusion or immature dermogram by light collimation and proper attaching strictly according to changes of the skin. Check per $2\sim 3$ hours the sensor placement and move it when the skin deteriorates. More frequent examinations may be required for different patients.

5.3.4 Setup of SpO2/Pulse parameters



Beep Volume Select the BEEP volume and options are **Off**, **1**, **2** and **3**. Once an option is selected, a testing beep will be produced.

HR Source Select the option of HR source, and options are **AUTO**, **ECG** and **PLETH**. When selecting **AUTO**, the HR source is ECG with the priority; and if there is no current ECG, the system automatically derives HR from SpO₂.

Scan Speed Select the scanning speed of the SpO₂ waveform, and options are 6.25mm/s, 12.5mm/s, 25mm/s and 50mm/s.

Alarm Setup Click and open the dialog of SpO₂ alarm configuration.

Average Time Select the average time for SpO_2 . The shorter the averaging time is, the quicker the monitor responds to the change in the patient's oxygen saturation level. (It is invalid while use Nellcor SpO_2 module).

SpO₂ Mode Select the response time mode for SpO₂, and options are **Common mode** and **Fast mode**. It is valid only use the Nellcor SpO₂ module.

■ When you use Masimo SpO₂ module, the below settings will be displayed.

FastSat The FastSat mode is clinically applicable during procedures when detecting rapid changes in oxygen saturation is paramount such as induction, intubation and sleep studies. You can select **On** or **Off** to enable or disable the FastSat mode.

Sensitivity Select the sensitivity level of Masimo SpO₂ module according to use condition, the options are **Max**, **Normal** and **Apod**. The **Apod** mode is the least sensitive in picking up a reading on patients with low perfusion but has the best detection for probe-off conditions. The Apod mode is useful for patients that are at particular risk of the sensor becoming detached (pediatric, combative, etc.)

Wave mode Select whether Resp wave includes in the SpO₂ wave or not, the options are **Resp filter** and **Resp Inclu.**

Alarm Setup



Adjust alarm Select this option to enter the configuration of alarm limits; conduct the configurations by turning the trim knob to select high and low limits and exit by selecting

 $\langle EXIT \rangle\,$. The upper part is the high limit and the lower one is the low limit.

SpO₂ alarm Select \langle **ON** \rangle to enable SpO₂ over limit alarm; select \langle **OFF** \rangle to disable SpO₂ over limit alarm.

PR alarm Select **<ON>** to enable PR over limit alarm; select **<OFF>** to disable PR over limit alarm.

■ When you use Masimo SpO₂ module, the below settings will be displayed.

Delay Time Select the delay time in the alarms as required in the options. If you select a delay time, when the SpO2 value is within the threshold of alarm limit, it will not trigger the alarm until reaching the delay time.

Threshold Select the threshold of the alarm limit.

5.3.5 Signal strength prompt

The signal strength prompt is used to indicate if the SpO_2 signal strength measured is adequacy.

Prompt	Description
Weak Signal	The invalidation weak signal
*	The low intensity signal
**	The medium intensity signal
***	The high intensity signal

5.3.6 Measurement Limitation

If you doubt the SpO_2 measurements, check the patient's vital signs first, then check the monitor and SpO_2 sensor. The following factors may influence the accuracy of measurements:

- -----Incorrect sensor application or use;
- —Significant levels of dysfunctional hemoglobins. (such as carboxyhemoglobin or methemoglobin);
- ——Intravascular dys 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 lights, infrared heating lamps, or direct sunlight (exposure to excessive illumination can be corrected by covering the sensor with a dark or opaque material)
- -----Excessive patient movement;
- -----Venous pulsations;
- ——Placement of a sensor on an extremity with a blood pressure cuff, arterial catheter, or intravascular line.
- -----Low perfusion;
- ——Electromagnetic interference, such as MRI device;
- ——Electrosurgical units.
 - Loss of pulse signal can occur in any of the following situation:
- ——The sensor is too tight;
- ——There is excessive illumination from light sources such as a surgical lamp, a brilirubin lamp, or sunlight;
- ——A blood pressure cuff is inflated on the same extremity as the one with a SpO2 sensor attached;
- ——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 is in shock.

5.3.7 Maintenance and Cleaning

Warning:

- **Do not sterilize by irradiation, steam, or ethylene oxide.**
- Do not subject the sensor to autoclaving.
- Do not immerse the sensor into any liquid.
- Do not use any sensor or cable that may be damaged or deteriorated.

Note: When disposing the disposable SpO₂ probe or useless SpO₂ probe, please observe all local, state, and federal regulations that relate to the disposal of this products or similar products.

For reusable SpO_2 sensor

Please unplug the sensor from the monitor before cleaning or disinfection.

Clean or disinfect the sensor before attaching to a new patient.

■ Cleaning:

Use a piece of clean cloth moistened in water or mild soap solution to clean the sensor and patient contact surfaces.

■ Disinfection:

Use a piece of clean cloth to wipe the sensor and patient contact surfaces with a 10% bleach solution or 70% isopropyl alcohol, clean with clear water and wipe it dry.

5.3.8 Masimo Information

Masimo SET.

Masimo Patents:

This device is covered under one or more of the following U.S.A. patents: 5,758,644, 5,823,950, 6,011,986, 6,157,850, 6,263,222, 6,501,975 and other applicable patents listed at: <u>www.masimo.com/patents.htm</u>.

No Implied License:

Possession or purchase of this device does not convey any express or implied license to use the device with unauthorized sensors or cables which would, alone, or in combination with this device, fall within the scope of one or more of the patents relating to this device.

5.3.9 Nellcor Information



This is the trademark of Nellcor Puritan Bennett Inc.

5.4 Measurement of TEMP

5.4.1 Brief Introduction to Measurement of TEMP

Monitor measures TEMP with TEMP sensors. The TEMP module of Monitor uses TEMP cable compatible with YSI-400. The minimum time to get accurate temperature measuring value is 3 minutes.

The monitor has two ports for body TEMP measurement, and can measure the temperature of two channels at the same time.

5.4.2 Preparatory Steps of the Measurement of TEMP

1) Plug the TEMP cables into the TEMP sockets of the monitor.

2) Place the TEMP sensors on body of patient and the screen will show the value of TEMP measurement.

3) Set the parameters relevant to TEMP.

5.4.3 Connecting Patient and Monitor

Refer to Fig.1-5-1 and plug the TEMP cable into the sockets marked with TEMP (either of TEMP1 and TEMP2), and then stick the TEMP sensor securely onto the body of patient.

Q Caution: The TEMP sensor and cables should be handled with care. When not in use, the sensor and the cable should be rounded into loose ring shape.

5.4.4 Setup of TEMP Parameters

	Total Paters
	remp setup
Unit	* C _
T1 Label	т1
T2 Label	T2
Alarm Setup	
Back to Main	
Set unit of TE	EMP.
10	

Unit Select the unit of TEMP, and options are $^{\circ}C$ and $^{\circ}F$.

T1 Label Select the labeling name for TEMP 1, and options are **T1, Eso, Naso, Tymp, Rect, Blad** and **Skin**.

T2 Label Select the labeling name for TEMP 2,and options are **T2**, **Eso**, **Naso**, **Tymp**, **Rect**, **Blad** and **Skin**.

Label	Meanings	Label	Meanings
Eso	Esophageal temperature	Rect	Rectal temperature
Naso	Nasopharyngeal temperature	Blad	Bladder temperature
Tymp	Tympanic temperature	Skin	Skin temperature

Alarm Setup Click and open the dialog of configuration for TEMP alarm.

TEMP Alarm	Adjust Alarm
Adjust Alarm	EXIT
T1 Alarm OFF T2 Alarm OFF Previous	Temp1 38.0
	34.0 Temp2 38.0
	15.0
Alarm limit setup.	

Adjust alarm Select this option to enter the configuration of alarm limits; conduct the configurations by turning the trim knob to select high or low limits and exit by selecting

 $\langle \text{EXIT} \rangle \,$. The upper part is the high limit and the lower one is the low limit.

T1 alarm Select **<ON>** to enable T1 over limit alarm; select **<OFF>** to disable T1 over limit alarm.

T2 alarm Select **<ON>** to enable T2 over limit alarm; select **<OFF>** to disable T2 over limit alarm.

5.4.5 Maintenance and Cleaning

Reusable temp probes

1. The temp probe should not be heated above 100° C. It should only be subjected briefly to temperatures between 80° C and 100° C.

2. Only detergents containing no alcohol can be used for disinfection.

3. The rectal probes should be used, if possible, in conjunction with a protective rubber cover.

■ Cleaning:

Use a piece of clean cloth moistened in water or mild soap solution to clean the probe.

■ Disinfection:

Use a piece of clean cloth to wipe the surface of the cable with 70% isopropyl alcohol, a 10% bleach solution or 2% Cidex®, clean with clear water and wipe it dry.

Warning: Disposable TEMP probes must not be re-sterilized or reused.

Note: For protecting environment, the disposable TEMP probe must be recycled or disposed of properly.

Disposal Note: Should the TEMP probe become damaged beyond repair, or for some reason its useful life is considered to be at an end, please observe all local, state, and federal regulations that relate to the disposal of this products or similar products.

Warning: The calibration of temperature measurement is necessary for every two years (or as frequently as dictated by your Hospital Procedures Policy). When you need calibrate the temperature, contact the manufacture please. **CP** Note: The self-test of the temperature measurement is performed automatically once every 10 minutes during the monitoring. The test procedure lasts about one second and does not affect the normal measurement of the temperature monitoring.

Note: If Temperature to be measured beyond probe's measuring range, over measuring range alarm will display on the screen. Check out if probe is on the corresponding patient body site, or change it to other site on the patient.

Note: If "TEMP self-check error" display on the screen, it is possibly that something is wrong with the temperature capture circuit, the operator should stop using the monitor and contact with the company.

5.5 Measurement of NIBP

5.5.1 Brief Introduction to Measurement of NIBP

Monitor automatically conducts measurement of NIBP with the method of shockwave. The method of shockwave indirectly estimates the systolic and diastolic pressures within the blood vessels by measuring the change of the pressure within blood pressure cuff along with the volume of the arteries and calculates the average pressure.

The measurement time of BP on a calm patient is less than 40s, and when each measurement ends, the cuff automatically deflates to zero.

The monitor applies to any standards of the cuffs for neonate, child and adult (including the cuffs used for arms and legs).

The monitor measures the blood pressure during the time of deflation. Monitor automatically conducts the second and third inflation measurements in case during the first inflation it is unable to measure the value of BP, and gives out the information for measurement failures.

The longest cuff pressure maintaining duration is 120 seconds (90 seconds in neonate mode), and when the time is exceeded, the air will be deflated automatically. The monitor has been designed with hardware protection circuit regarding overpressure, errors of microprocessors, and the occurrence of power failure.

5.5.2 Preparatory Steps of Measurement of NIBP

1) Plug the air hose of cuff into the NIBP socket of the monitor and tighten it clockwise to ensure secure contact of the plug and the socket (Please note that the plug should be loosened by turning counterclockwise first before unplugging).

- 2) Tie the cuff on the arm of patient.
- 3) Set the parameters and modes relevant to NIBP.

C Note: Make sure that the air conduit connecting the blood pressure cuff and the monitor is neither blocked nor tangled, and avoid compression or restriction of air conduit.

5.5.3 Connecting to Patient and the Monitor

Refer to Fig. 1-5-1 to plug the connector of air hose on cuff into the socket marked with NIBP and wrap the cuff onto the arm of patient. Make sure the mark of Φ on the cuff is placed on the femoral artery of the arm and the air hose should be below the cuff so as to ensure the air hose is not snarled after coming out of the cuff. The white line on the cuff should be within the range of " \iff ", otherwise it will be necessary to replace it with a more suitable cuff (smaller or bigger one). The cuff should be placed on the same plane with the heart so as to prevent the errors in readings caused by the effects of hydrostatics of the blood column between the heart and the cuff. If the position of the cuff is higher than the plane of heart, the measured BP readings tend to be smaller; in case the position of the cuff is lower than the plane of the heart, the measured BP readings tend to be higher.

P Note: The accuracy of measurement of BP depends on the suitability of the cuff. Select the size of the cuff according to the size of the arm of patient. The width of the cuff should be 40% of the circumference of the upper arm or 2/3 of the length of the upper arm.

Warning:

- Vou must not perform NIBP measurements on patients with sickle-cell disease or under any condition that the skin is damaged or expecting 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.
- Prolonged non-invasive blood pressure measurements in Auto mode are associated with purport, 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.

5.5.4 Setup of NIBP Parameters



Auto Time Configure the cycle intervals of BP measurement and options are 1min,2min, 3min, 4min, 5min, 10min, 15min, 30min, 60min, 90min, 2Hour, 4Hour and8Hour. During measurements, it cannot be altered.

Mode Configure the measurement mode of NIBP and options are Manual, Auto and STAT.

If STAT mode is configured, after measurement, the system will be automatically configured as the previous measurement mode. There is no STAT mode for neonatal. If STAT is selected, the rapid measurement will be initiated once it is confirmed.

Object Objects of measurements shall be configured, and options are **Adult**, **Children**, **Neonatal** and **Hyperpiesia**. The selection of objects of measurements during the measuring process will terminate the ongoing measurement.

Init_Inflate Select an appropriate initial cuff inflation pressure according to patient category and requirement.

Unit Select the unit for the NIBP measurement, and options are kPa and mmHg.

NIBP Alarm Click and open the dialog of alarm configuration of NIBP.

Leakage Air Leakage test

Vein Pun. You can use the NIBP cuff to cause sub-diastolic pressure, and block the venous blood vessel to assist venous puncture. Select $\langle Vein Pun. \rangle$, and set to a proper value.

Start Pun. After you set a proper value for $\langle Vein Pun. \rangle$, and select $\langle Start Pun. \rangle$ to start it.

• NIBP Alarm

NIBP Alarm	Adjust Alarm
Adjust Alarm	Exit
Alarm Source S/D/M	NIBP Sys
Alarm Switch <mark>ON</mark>	160
Previous	90
	NIBP Dia
	90
	50
	NIBP Mean
	110
Alarm limit setup.	60

Adjust alarm Select this option to enter the configuration of alarm limits; conduct the configurations by turning the trim knob to select high or low limits and exit by selecting

 $\langle EXIT \rangle$. The upper part is the high limit and the lower one is the low limit. Alarm Source Select the alarm source for NIBP parameters. When the selected parameter or one of the parameter exceed alarm limit, the monitor will give out alarm signal. The options are as follow:

 $\langle S \rangle$: Only Systolic pressure exceeds the alarm limit will trigger the alarm system.

 $\langle \mathbf{D} \rangle$: Only Diastolic pressure exceeds the alarm limit will trigger the alarm system.

 $\langle M \rangle$: Only Mean pressure exceeds the alarm limit will trigger the alarm system.

(S/M): Systolic pressure or mean pressure exceeds the alarm limit will trigger the alarm system.

 $\langle D/M \rangle$: Mean pressure or diastolic pressure exceeds the alarm limit will trigger the alarm system.

 $\langle S/D \rangle$: Systolic pressure or diastolic pressure exceeds the alarm limit will trigger the alarm system.

 $\langle S/D/M \rangle$: Mean pressure, systolic pressure or diastolic pressure exceeds the alarm limit will trigger the alarm system.

Alarm Switch Select <ON> to enable parameter over limit alarm; select <OFF> to disable parameter over limit alarm.

5.5.5 Precautions during Measurement

■ If the BP of the patient is above 180mmHg, <HYPER> measurement mode is recommended.

■ When using the STAT measurement or AUTO measurement, if the time duration is relatively long, care must be taken to check such abnormalities as purple spots, coldness and numbness at the limb end. If there are such phenomena, the cuff should be relocated or the measurement of NIBP should be halted. **To neonate mode, STAT measurement is unavailable.**

■ The presence of factors that change the properties of the cardiovascular dynamics of patient will adversely affect the measurement value of the monitor, and shock and hypothermia will also affect the accuracy of the measurement.

■ When the built-in main artery balloon pump is applied on the patient, the measurement value of NIBP will be affected.

■ For the limb that is on an intravenous drip or in a catheter insertion, or if the patient is connected to the heart-lung machine, or the patient is experiencing shiver or convulsions, the measurement of NIBP cannot be conducted.

■ When errors occur in the measurement of NIBP, the error codes will appear in the parameter display zone of the NIBP, and for the cause of the errors, please refer to **chapter 6.8.6**.

Measurement Limitations

Measurements are impossible with heart rate extremes of less than 40 bpm or greater than 240 bpm, or if the patient is on a heart-lung machine.

The measurement may be inaccurate or impossible:

1) If a regular arterial pressure pulse is hard to detect

2) With cardiac arrhythmias

3) With excessive and continuous patient movement such as shivering or convulsions

4) With rapid blood pressure changes

5) With severe shock or hypothermia that reduces blood flow to the peripheries

6) With obesity, where a thick layer of fat surrounding a limb dampens the oscillations coming from the artery

7) On an edematous extremity.

5.5.6 Periodic Check

□ Calibration

Warning: The calibration of the NIBP measurement is necessary for every two years (of as frequently as dictated by your Hospital Procedures Policy). The performance should be checked according to the following details.

Procedure of the Pressure Transducer Calibration:

- 1) Replace the cuff of the device with a rigid metal vessel with a capacity of 500 ml $\pm 5\%$.
- 2) Connect a calibrated reference manometer with an error less than 0.8 mmHg and a ball pump by means of a T-piece connector and hoses to the pneumatic system.
- 3) Access the NIBP menu.
- Turn the trim knob to the (Manometer) option and press. Then the NIBP module has started performing calibration.
- 5) Inflate the pneumatic system to 0mmHg, 50mmHg and 200 mmHg by ball pump separately. The difference between the indicated pressure of the reference manometer and the indicated pressure of the monitor will not exceed 3 mmHg. Otherwise, please contact our customer service.



6) Press the $\langle NIBP/STAT \rangle$ button on front panel can stop the calibration.

Fig. 5-5-6 Diagram of NIBP calibration

Air Leakage check

Procedure of the air leakage test:

- 1) Connect the cuff securely with the socket for NIBP air hole.
- 2) Wrap the cuff around the cylinder of an appropriate size.
- 3) Access the NIBP setup window.
- 4) Select the 〈Leakage〉 option and press. Then the prompt "Air Leakage test" will appear on the NIBP parameter area indicating that the system has started performing Air Leakage test.
- 5) The system will automatically inflate the pneumatic system to about 180mmHg.
- 6) After 20 seconds or so, the system will automatically open the deflating valve, which marks the completion of an air leakage test.
- 7) If no error information displays on NIBP parameter area, it indicates that the airway is in good situation and no air leaks exist. However if the prompt "AIR

SYSTEM LEAK" appears in the place, it indicates that the airway may have air leaks. In this case, the user should check for loose connection. After confirming secure connections, the user should re-perform the air leakage test. If the failure prompt still appears, please contact the manufacturer for repair.

8) Press the $\langle NIBP/STAT \rangle$ button on front panel can also stop the test.



Fig. 5-5-7 Diagram of air leakage check

5.5.7 Maintenance and Cleaning

Warning: Do not squeeze the rubber hose on the cuff. Do not allow liquid to enter the connector socked at the front of the monitor. Do not wipe the inner part of the connector socked when cleaning the monitor.

Warning: If liquid is inadvertently splashed on the equipment or its accessories, or may enter the conduit or inside the monitor, contact local customer service center.

Warning: Disposable blood pressure cuff must not be re-sterilized or reused.

C Disposal Note: Should the blood pressure cuff become damaged beyond repair, or for some reason its useful life is considered to be at an end, please observe all local, state, and federal regulations that relate to the disposal of this products or similar products.

For Reusable Blood Pressure Cuff:

■ Cleaning:

1. Please clean the cuff termly.

2. Take down the cuff from the connector, take out the bladder from the cover of the cuff, as the figure 5-5-8 shows.

3. Use a piece of clean cloth moistened in water or mild soap solution to clean the bladder and the tube.

4. Clean the cover of the cuff with the mild soap solution.

5. Dry the cover and the bladder. Roll the bladder and insert it into the large opening, as the figure 5-5-9 shows. Hold the hose and the cuff and shake the complete cuff until the bladder is in position. Put the hose out through the small hole under the internal flap.



Fig. 5-5-8 Take out the bladder

Fig. 5-5-9 Insert the bladder

S[™] Warning:

- Clean the bladder frequently, will cause the bladder scathed, except the necessary, do not clean the bladder.
- **Do not dry the bladder and cover with high temperature.**
- If need the high level disinfecting, please selecting the disposable cuff.

5.6 Measurement of IBP

5.6.1 Brief Introduction to Measurement of IBP

The method of IBP measurement is direct measuring the BP of artery or veins on the pressure sensor mainly through liquid coupling so as to obtain the pressure curve of the continuous BP.

The IBP parameters of Monitor can select Arterial Pressure (ART), Pulmonary Artery pressure (PA), Left Atrium Pressure (LAP), Right Atrium Pressure (RAP), Central Venous Pressure (CVP), Intracranial Pressure (ICP).

Monitor has two measurement channels for IBP, and the IBP of two channels can be measured at the same time.

5.6.2 Preparatory Steps for Measurement of IBP

1) Plug the cable of IBP into the IBP socket (either CH1 or CH2), and connecting cable to the pressure transducer. Fill the pressure transducer and extension tube with saline water mixed with heparin. Press the flexible valve to expel the saline water from the air outlet to expel air bubbles, and then reset it to zero.

Note: The method of touching test is to touch slightly the surface with finger. Waveforms should appear on the screen of the main unit. The blue ball cover should be put on the surface immediately when the energy converter is not used.

Note: Anytime the user applies a new transducer, it should be verified or periodically verified according to the hospital operating rules.

Warning: Disposable pressure transducer should not be reused. And it must be used before expired data. Do read the expired data on the IBP accessory package bag.

Warning: When the monitor is used with HF surgical equipment, the transducer and the cables must be avoided conductive connection to the HF equipment to protect against burns to the patient.

The specified transducer is designed to protect against the effects of a discharge of a cardiac defibrillator. When the patient is in the defibrillation, the waveform of IBP maybe distorted temporarily. After the defibrillation, the monitoring will go on normally, the operation mode and the user configuration are not affected.

Warning: The operator should avoid contact with the conductive parts of the appurtenance when being connected or applied.

- 2) Plug the cable of IBP into the IBP socket on the right panel of the monitor. Connect the extension tube of the transducer and blood vessel with the artery needles and secure them, then make sure three-way valve 1 and three-way valve 2 (See Fig. 5-6-4) are in a state of ON. At this moment, BP waveforms should appear on the screen of the monitor.
- 3) Set up parameters and modes relevant to IBP.

5.6.3 Setup of IBP Parameters

	IBP2 Setup
IBP Label	IBP2
Unit	mmHg
Scan Speed	25mm/s
Wave Scale	0~200mmHg
Display	S/D(M)
IBP Alarm	
IBP Zero	
Back to Main	
Set label.	

IBP Label Select the names of IBP labels. Options are **IBP1**, **IBP2**, **ART**, **CVP**, **PA**, **RAP**, **ICP** and **LAP**.

Unit Select the units of IBP, and options are mmHg, kPa and cmH₂O.

Scan speed Select the scanning speed of IBP waveforms, and options are 12.5mm/s, 25mm/s and 50mm/s.

Wave scales Select the scale of IBP waveforms and options are AUTO, 0~200mmHg, 0~300mmHg, -10~20mmHg and -50~300mmHg.

Display Select the format of IBP display, and options are S/D (M), S/D, Mean and M (S/D).

IBP Zero Conduct zero-calibration on IBP.

IBP Alarm Click and open the dialog of IBP alarm limit configuration.

IBP1 Alarm	Adjust Alarm
Adjust Alarm	Exit
Alarm Source S/D/M	IBP1 Sys
Alarm Switch OFF	160
	90
	IBP1 Dia
	90
	50
	IBP1 Mean
	110
Set IBP1 alarm source.	60

Adjust alarm Select this option to enter the configuration of alarm limits; conduct the configurations by turning the trim knob to select high and low limits and exit by selecting

 $\langle EXIT \rangle$. The upper part is the high limit and the lower one is the low limit. Alarm Source Select the alarm source for IBP parameters. When the selected parameter or one of the parameter exceed alarm limit, the monitor will give out alarm signal. The options are as follow:

 $\langle S \rangle$: Only Systolic pressure exceeds the alarm limit will trigger the alarm system.

 $\langle \mathbf{D} \rangle$: Only Diastolic pressure exceeds the alarm limit will trigger the alarm system.

 $\langle \mathbf{M} \rangle$: Only Mean pressure exceeds the alarm limit will trigger the alarm system.

(S/M): Systolic pressure or mean pressure exceeds the alarm limit will trigger the alarm system.

 $\langle D/M \rangle$: Mean pressure or diastolic pressure exceeds the alarm limit will trigger the alarm system.

 $\langle S/D \rangle$: Systolic pressure or diastolic pressure exceeds the alarm limit will trigger the alarm system.

 $\langle S/D/M \rangle$: Mean pressure, systolic pressure or diastolic pressure exceeds the alarm limit will trigger the alarm system.

Alarm Switch Select <ON> to enable parameter over limit alarm; select <OFF> to disable parameter over limit alarm.

5.6.4 Calibration of Zero-point

Start the unit and preheat it for 3 minutes. If it is in a stable state, turn off three-way valve 2 and turn on three-way valve 1, and then select option in *<***IBP Zero***>* of *<***IBP Setup***>*, then it can be seen on the screen that the scanning baseline has returned to zero baseline.

Solution Note: In the course of zeroing, should turn off the three-way valve near artery needle, don't connect artery needle with patient and make sure there is no air inside the whole tube.

5.6.5 Connecting to Patient As shown in Fig. 5-6-4

C Note: The pressure measuring side of the transducer should be on the same plane as the heart of the patient in the process of zero-setting and measurement and the user should make sure there is no air inside the whole tube in order to assure the correctness of the measured results. If air is found in tube or in pressure transducer, they must be rinsed by physiological salt solution.



Warning: If liquid (not the liquid which used to douche the tubes and pressure transducers) spills on equipment or accessories, especially when the liquid is likely to enter the equipment or transducer, contacting with the maintenance department of the hospital immediately.



Fig. 5-6-4 Schematic diagram for installation of IBP sensor

5.6.6 Setup of Range

The setup of IBP module range can provide you with the best waveforms and the best measurement results. Based on different contents of measurement, there are two ranges for selection, and each group has 5 options:

• Arterial Pressure (ART):

AUTO, 0-50mmHg, 50-150mmHg, 100-240mmHg, 0-300mmHg

• Pulmonary Artery pressure (PA), Left Atrium Pressure (LAP) ,Right Atrium Pressure (RAP) ,Central Venous Pressure (CVP), Intracranial Pressure (ICP)

AUTO, 0-20mmHg, 0-30mmHg, 0-50mmHg, 0-80mmHg

Note: AUTO will adjust the scale on which the pressure waveform is displayed on the screen automatically for the best observation status.

5.6.7 IBP Transducer Zero and Calibration

■ IBP Transducer Zero

Warning: It is the responsibility of the user to ensure that a zero procedure has recently been done on the transducer, otherwise there will be no recent, valid zero value for the instrument to use, which may result in inaccurate measurement results.

Procedure of the IBP Transducer Zero:

- 1) Turn off patient stopcock before you start the zero procedure.
- 2) The transducer must be vented to atmospheric pressure before the zero procedure.
- 3) The transducer should be placed at the same height level with the heart, approximately mid-axially line.
- 4) Access the Set IBP menu.
- 5) Turn the dial to pick the Zero1 item (Pick the Zero2 item when zeroing channel 2 IBP) and press will start zero the transducer.
- 6) Wait 3 seconds for the Zeroing procedure end and the pressure value that is displayed on screen will approximately return to zero.

Q Caution: Zero procedure should be performed before starting the monitoring and at least once a day and whenever after each disconnect-and-connect of the cable.

IBP Calibration

Caution:

- Mercury calibration should be performed by the biomedical engineering department either whenever a new transducer is used, or as frequently as dictated by your Hospital Procedures Policy.
- The purpose of the calibration is to ensure that the system gives you accurate measurements.
- > Before starting a mercury calibration, a zero procedure must be performed.
- If you need to perform this procedure yourself you will need the following pieces of equipment:
 - Standard sphygmomanometer
 - 3-way stopcock
 - Tubing approximately 25 cm long

The Calibration Procedure:

Warning: You must never perform this procedure while patient is being

monitored.

- 1) Close the stopcock that was open to atmospheric pressure for the zero calibration.
- 2) Attach the tubing to the sphygmomanometer.
- 3) Ensure that connection that would lead to patient is off.
- 4) Connect the 3-way connector to the 3-way stopcock that is not connected to the patient catheter.
- 5) Open the port of the 3-way stopcock to the sphygmomanometer.
- 6) Inflate to make the mercury bar rise to 0, 50 and 200 mmHg separately. The difference between the indicated pressure of the sphygmomanometer and the indicated pressure of the monitor will not exceed $\pm 4\%$ or ± 4 mmHg, whichever is greater. Otherwise, please contact the manufacturer.
- 7) After calibration, disassemble the blood pressure tubing and the attached 3-way valve.

5.6.8 Maintenance and Cleaning

Warning: The disposable transducers must not be re-sterilized or re-used.

Note: For protecting environment, the disposable transducers must be recycled or disposable of properly.

Disposal Note: When disposing the disposable transducers and tubing, please observe all local, state, and federal regulations that relate to the disposal of this products or similar products.

5.7 Measurement of CO₂ (Sidestream, CPT)

Use the CO_2 measurement to monitor the patient's respiratory status and to control patient ventilation. The measurement principle is primarily based on the fact that CO_2 molecules can absorb special infrared light, where the intensity of infrared light passing the respiratory gas is measured with a photo detector. As some of the infrared light is absorbed by the CO_2 molecules, the amount of light passing the gas probe depends on the concentration of the measured CO_2 .

5.7.1 Brief Introduction to Measurement of Sidestream CO2

• According to the Fig. 5-7-1, snap slantways dehydration flask on the receptacle fixed on the right panel of monitor. It will click into place when properly seated.



Fig.5-7-1 The installation sketch map of dehydration flask

• According to the Fig. 5-7-2, nip slantways the dehydration flask and disconnect from the receptacle fixed on the right panel of monitor. It will be remove the dehydration flask.



Fig.5-7-2 Remove sketch map of dehydration flask

According to the Fig. 5-7-3, one end of the sampling tube has been connected with screw thread interface of the dehydration flask, and the other end of the sampling tube has been connected with the screw thread interface tube (Φ 10mm) of the patient Anaesthesia machine or Ventilator (If not the type screw thread interface tube, please connect the requirement type tube) the sampling tube's port can also been fixed on the naris of patient with adhesive plaster.



Fig. 5-7-3 Connected with dehydration flask

- Select < CO₂ Setup> button in Main Screen, then select the <Start> and press this button to start sampling pump, begin measuring EtCO₂.
- Pay attention to the water level of dehydration flask. If the highest water level reaches, Please replace the dehydration flask in time to prevent the module from soaking by water.
- When air is getting across the sampling tube, a period of time will cost. So, a delay time will appear from starting measure to showing waveform in the screen and measuring result.
- Please keep the sampling tube clean, and prevent the tube from clogging by dust.

C Note : Dehydration flasks and sampling tubes are disposable, please use products provided or designated by manufacturer.

5.7.2 Setup of CO₂ parameters



Scan Speed Select the scanning speed of RESP waveforms, and options are 6.25mm/s, 12.5mm/s and 25mm/s.

Resp Source Select Resp source. And options are AUTO, ECG and EtCO₂.

Unit Select the unit for CO₂, and options are **mmHg**, % and **kPa**.

Resp Gain Select the gain of RESP waveform, and options are 1x, 2x and 4x.

Alarm Setup Click and open the dialog of CO₂ alarm.

Start Press this button to start the sampling pump to initiate the measurement of CO_2 (only valid on sidestream CO_2).

Stop Press this button to switch off the sampling pump and terminate the measurement of CO_2 (only valid on sidestream CO_2).

Offset Cal Select the mode of drift calibration. Options are Cancel, Automatic and Manual. During the common measurements, please remain the default configuration as Automatic. Only when it is necessary to conduct gain calibration should this option be configured as Manual (only valid on sidestream CO_2 which sampling pump has been started).

Wave Type Select display type of CO₂ waveform. The options are Draw and Fill.Back to Main Return to main screen.
Alarm Setup



Adjust alarm Select this option to enter the configuration of alarm limits; conduct the configurations by turning the trim knob to select high or low limits and exit by selecting $\langle EXIT \rangle$. The upper part is the high limit and the lower one is the low limit.

EtCO₂ alarm high limit, its configuration range is $0.0 \sim 13.1\%$ ($0 \sim 99.6$ mmHg) continuously adjustable, no lower than the low limit; the configuration range of EtCO₂ alarm low limit is $0.0 \sim 13.1\%$ ($0 \sim 99.6$ mmHg) continuously adjustable, no higher than the high limit.

FiCO₂ alarm high limit, its configuration range is $0.0 \sim 13.1\%$ ($0 \sim 99.6$ mmHg) continuously adjustable, no lower than the low limit; the configuration range of FiCO₂ alarm low limit is $0.0 \sim 13.1\%$ ($0 \sim 99.6$ mmHg) continuously adjustable, no higher than the high limit.

Resp alarm high limit, its configuration range is $0 \sim 150$ rpm continuously adjustable, no lower than the low limit; the configuration range of Resp alarm low limit is $0 \sim 150$ rpm continuously adjustable, no higher than the high limit.

Apnea alarm when the time of zero Resp rate has reached this time scale, the alarm will be set off. Options are Off, 5s, 10s, 20s, 40s, 60s, 80s, 100s and 120s.

Alarm Source Select the alarm source for CO₂ parameters. When the selected parameter or one of the parameter exceed alarm limit, the monitor will give out alarm signal. The options are as follow:

 $\langle Et \rangle$: Only EtCO₂ exceeds the alarm limit will trigger the alarm system.

 $\langle \mathbf{RR} \rangle$: Only RR exceeds the alarm limit will trigger the alarm system.

 $\langle Et/RR \rangle$: EtCO₂ or RR exceeds the alarm limit will trigger the alarm system.

Alarm Switch Select the switch of CO2 alarm.

Caution: When the monitor is powered on, the pump in the CO₂ module is set off as default configuration. Since long-time running of sampling pump could shorten the life of CO₂ module, please start sampling pump manually, and stop the sampling pump after monitoring has been finished.

5.7.3 Gain Calibration

Please carry out gain calibration and manual offset calibration, when the following conditions happened:

- 1. The module has been used for between half a year and one year.
- 2. The precision of EtCO₂ reading has been doubted by clinical physician.
- 3. After the latest calibration, atmospheric pressure or height above sea level varies evidently.

The apparatus has already been calibrated before leaving factory. User can directly apply it to measuring in normal conditions, to the exclusion of the previous conditions.

Gain calibration and manual offset calibration must be carried out if the previous conditions happened. The following procedures must be observed:

1. Select <**Menu>→**<**Monitor Setup>→** <**System Setup>→**<**Machine>**, please set <**CO**₂ **Cal Mode> ON**;

2. When the monitor has been run for 30 minutes, one end of the sampling tube has been connected with the module, the other end has been exposed in the undefiled atmosphere, please adjust **CFFSET CAL>** to **Manual** in the CO₂ setup dialog, and press the button of **CFFSET CAL>** to start offset calibration.

3. Please connect the adjusting device according to Fig.5-7-4. While the standard gas of pressure CO_2 5.0% (38.0mmHg) getting across the sampling tube, observe pressure measuring apparatus carefully to ensure that the pressure of standard gas is one standard atmosphere (the range of error is ±5%). Then select <Menu>→<Monitor Setup> → <System Setup>→<Machine>, press the <CO₂ GAIN CAL>,and input 5% in the popped up window. Select<OK>to start gain calibration. About 5 seconds later, the reading having calibrated will be shown in the screen.



This end Connects with pressure measurement apparatus.



Warning: The standard gas of which the pressure of CO₂ is 5.0% (38.0mmHg) must be used during gain calibration. Otherwise, measurement values will not be accurate.

Note: User may only calibrate the device under the instruction of the technical personnel authorized by company. Moreover, wrong calibrating procedure may result in false reading.

5.8 Measurement of CO2 (Microstream, LoFlo)

Use the CO_2 measurement to monitor the patient's respiratory status and to control patient ventilation.

5.8.1 Preparing to Measure CO2

1. Attaching the LoFlo Module Cable

To attach the LoFlo module cable, plug the cable into the CO_2 socket on the left panel of monitor by matching the key on the cable to the key on the connector.

Caution: To remove the module cable from the monitor, grasp the collar surrounding the cable and pull up.

2. Attaching the Sample Cell

Follow these steps:

1) Insert the LoFlo sample cell into the LoFlo sample cell receptacle .A "click" will be heard when the sample cell is properly inserted. (Fig.5-8-1, Fig.5-8-2)



Fig.5-8-2

P Note:

- Inserting the sample cell into the receptacle automatically starts the sampling pump. Removal of the sample cell turns the sample pump off.
- To remove the sample cell from the sample cell receptacle, press down on the locking tab and pull the sample cell from the sample cell receptacle.

2) If the sampling pump fails to turn on, or runs intermittently, perform a "Zero" procedure. (Refer to the *chapter 5.8.3*)

3) Ensure that the LoFlo module exhaust tube vents gases away from the module environment.

4) Wait for the CO2 module to warm up.

The monitor will display the **Sensor Warm Up** message for approximately one minute while the module warms up to operating temperature. The message disappears when the module is ready for use.

Note: Warm up time varies with ambient temperature of the module.

5.8.2 Setup of CO2 parameters



Scan speed Select the scanning speed of RESP waveforms, and options are 6.25mm/s, 12.5mm/s and 25mm/s.

RESP source	Select RESP source.	And options ar	e AUTO,	ECG and EtCO	2.
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Unit Select the unit for CO₂, and options are **mmHg**, % and **kPa**.

Resp Gain Select the gain of RESP waveform from ECG, and options are 1x, 2x and 4x.

Alarm setup Click and open the dialog of CO₂ alarm.

CO₂ setup Click and open the dialog of CO₂ setup.

Wave Type Select display type of CO₂ waveform. The options are Draw and Fill.

Back to Main Return to main screen.

	Constantion (Constantion)
	CU2 Setup
Gas Temp	35
Barometric	760
EtCO2 Period	1 breath
Zero Gas	Air
Compensation	16
Balance gas	Air
Anesthetic	
ZERO	
Previous	
Set temper	ature of the gas
mixture. D	efault is 35 °C.

Gas Temp Select the temperature of gas. (Turn the trim knob with an increment or decrement of 1° C)

Barometric Select the Atmospheric pressure. (Turn the trim knob with an increment or decrement of 1mmHg)

EtCO2 PeriodSelect the response time of EtCO2, the options are 1 breath, 10s and 20s.Zero GasSelect the gas type of zeroing, the options are Air and N2.

Compensation Select the concentration of oxygen. (Turn the trim knob with an increment or decrement of 1%)

Balance gas Select the balance gas type, the options are Air, N_20 and Helium.

AnestheticSelect the concentration of balance gas. (Turn the trim knob with anincrement or decrement of 0.1%)

Zero Press the button to start zeroing. It is only valid when the system detects that the module can be zeroed.

5.8.3 Zero

Zeroing allows the LoFlo module or CAPNOSTAT 5 sensor to adjust to the optical characteristics, in order to obtain accurate readings. While zeroing is recommended the first time a LoFlo module or CAPNOSTAT 5 sensor is connected to the unit, it is only absolutely necessary when the message **Zero Required** is displayed.



* Warning:

Always ensure that the sample cell is properly connected to the LoFlo module before zeroing.

Always ensure that the CAPNOSTAT5 sensor is properly connected to the airway adapter before zeroing.

Follow these steps:

1) Ensure that the nasal cannula or airway adapter is not connected to the patient or close to any source of CO2 (including the patient's, your own, exhaled breath and ventilator exhaust valves).

1) Press the $\langle Zero \rangle$ option in $\langle CO_2 Setup \rangle$ menu. The unit zeroes the module and displays the **Zero In Progress** message for approximately 15-20 seconds. The message disappears upon completion of the zeroing.

P Note:

- Do not attempt zeroing for 20 seconds after removing the adapter or cannula from the patient's airway. This time allows any CO2 remaining in the adapter or cannula to dissipate before zeroing.
- Do not attempt to zero the module while the adapter or cannula is in the patient's airway.
- **Do not attempt zeroing if the temperature is not stable.**
- Zeroing with CO2 in the adapter or cannula can lead to inaccurate measurements or other error conditions. If you attempt zeroing while CO2 remains in the adapter or cannula, the time required to zero the module may be increased.

5.8.4 Applying Microstream airway adapter or cannula

For intubated patients requiring an airway adapter: Install the airway adapter at the proximal end of the circuit between the elbow and the ventilator Y section. (Fig.5-8-3)



Fig.5-8-3

For intubated patients with an integrated airway adapter in the breathing circuit: Connect the male connector on the straight sample line to the female port on the airway adapter. (Fig.5-8-4)



Fig.5-8-4

For non-intubated patients: Place the nasal cannula onto the patient. (Fig.5-8-5)





For patients prone to mouth breathing use an oral-nasal cannula. Trim the oral sampling tip if necessary to fit the patient. It should extend down past the teeth and be positioned in the mouth opening. Remove the cannula from the patient if the tip needs to be trimmed. (Fig.5-8-6)



Fig.5-8-6

For nasal or oral-nasal cannulas with oxygen delivery, place the cannula on the patient as shown then attach the oxygen supply tubing to the oxygen delivery system and set the prescribed oxygen flow.

Warning: Always connect the airway adapter to the sensor before inserting the airway adapter into the breathing circuit. In reverse, always remove the airway adapter from the breathing circuit before removing the sensor.

Caution: Always disconnect the cannula, airway adapter or sample line from the sensor when not in use.

5.8.5 Removing Exhaust Gases from the System

Warning: When using the microstream CO2 measurement on patients who are receiving or have recently received anesthetics, connect the outlet to a scavenging system, or to the anesthesia machine/ventilator, to avoid exposing medical staff to anesthetics.

Use an exhaust tube to remove the sample gas to a scavenging system. Attach it to the microstream sensor at the outlet connector.

5.8.6 Safety considerations



- Do not use in the presence of flammable anesthetics or other flammable gasses. Use of the LoFlo Module in such environment may present an explosion hazard.
- Electrical Shock Hazard: Always disconnect the LoFlo Module before cleaning. Do not use if it appears to have been damaged. Refer servicing to qualified service personnel.
- Do not position the sensor cables or tubing in any manner that may cause entanglement or strangulation.
- Reuse, disassembly, cleaning, disinfecting or sterilizing the single patient use cannula kits and on-airway adapters may compromise functionality and system performance leading to a user or patient hazard. Performance is not guaranteed if an item labeled as single patient use is reused.
- Inspect the microstream on- airway adapters, microstream sampling kits and CO2 airway adapters for damage prior to use. Do not use the microstream onairway adapters, microstream sampling kits and CO2 airway adapters if they appear to be damaged or broken.
- Replace the microstream on- airway adapters, microstream sampling kits and CO2 airway adapters if excessive secretions are observed.
- Monitor the CO₂ waveform (Capnogram). If you see changes or abnormal appearance check the airway adapters and the sampling line. Replace it if needed.
- **Do not operate the LoFlo Module when it is wet or has exterior condensation.**
- **Do not apply excessive tension to any cable.**
- Do not use device on patients that can not tolerate the withdrawal of 50 ml/min±10 ml/min from the airway or patients that can not tolerate the added dead space to the airway.
- **Do not connect the exhaust tube to the ventilator circuit.**

Caution:

- Use only accessories provided by manufacturer.
- **Do not sterilize or immerse the LoFlo Module in liquids.**
- **Do not clean the LoFlo Module and accessories except as directed in this manual.**
- **Remove the LoFlo sampling kit sample cell from the receptacle when not in use.**
- **Do not stick appendage into sample receptacle.**
- Always insert sample cell before inserting the on-airway adapter into the ventilated circuit
- Always remove the on-airway adapter from the ventilated circuit before removing the sample cell.

P Note:

- This product and its accessories are latex free.
- After the life cycle of the LoFlo Module and its accessories have been met, disposal should be accomplished following national and local requirements.
- Nitrous oxide, elevated levels of oxygen and helium can influence the CO2 measurement. Please setup gas compensation according to actual state.
- Barometric pressure compensation is required to meet the stated accuracy of the LoFlo Module.

5.9 Measurement of CO2 (Mainstream, CAPNOSTAT5)

5.9.1 Preparing to Measure CO2

1. Attaching the CAPNOSTAT 5 sensor cable

To attach the CAPNOSTAT 5 sensor cable, plug the cable into CO_2 socket on the left panel of monitor by matching the key on the cable to the key on the connector.

Q Caution: To remove the sensor cable from the monitor, grasp the collar surrounding the cable and pull up.

2. Selecting a mainstream airway adapter

Select an airway adapter based on the patient's size, ET tube diameter and monitoring situation. For more information refer to the following table or contact manufacturer.

Airway Adapter Type	ET Tube Diameter	
SPU* Pediatric/Adult	> 4.0 mm	
Adult Reusable	> 4.0 mm	
SPU* Neonatal/Pediatric	≤ 4.0 mm	
Neonatal Reusable	≤ 4.0 mm	

*SPU = Single Patient Use

3. Attaching the airway adapter to the CAPNOSTAT 5 sensor

Before attaching the airway adapter to the CAPNOSTAT 5 sensor, verify that the airway adapter windows are clean and dry. Clean or replace the adapter if necessary. Follow these steps:

1) Align the arrow on the bottom of the airway adapter with the arrow on the bottom of the sensor.

2) Press the sensor and airway adapter together until they click.

3) Wait for the airway adapter and sensor to warm up.

The monitor will display the **Sensor Warm Up** message for approximately one minute while the sensor and adapter warm to operating temperature. The message disappears when the sensor is ready for use.

Note: Warm up time varies with ambient temperature of the module.

4. Zero

Please refer to chapter 5.8.3

5. Attaching the airway adapter to the airway circuit

After zeroing, attach the airway adapter to the airway circuit as follow.(Fig.5-9-1)





6. Ensure the airway air-proof and ready to measure

5.9.2 Setup of CO2 parameter

Please refer to chapter 5.8.2

5.9.3 Zero

Please refer to chapter 5.8.3

5.9.4 Safety considerations



Warning:

- **Do not use in the presence of flammable anesthetics or other flammable gasses.** Use of the CAPNOSTAT5 sensor in such environment may present an explosion hazard.
- Electrical Shock Hazard: Always disconnect the CAPNOSTAT5 sensor before cleaning. Do not use if it appears to have been damaged. Refer servicing to qualified service personnel.
- Do not position the sensor cables or tubing in any manner that may cause entanglement or strangulation.
- Reuse, disassembly, cleaning, disinfecting or sterilizing the single patient use CO2 airway adapters may compromise functionality and system performance leading to a user or patient hazard. Performance is not guaranteed if an item labeled as single patient use is reused.
- Inspect the CO2 airway adapters for damage prior to use. Do not use the CO2 airway adapters if they appear to be damaged or broken.
- Replace the CO2 airway adapters if excessive secretions are observed.
- If the CO2 waveform (Capnogram) appears abnormal, inspect the CO2 airway adapters and replace if needed.
- Monitor the CO2 waveform (Capnogram) for elevated baseline. Elevated baseline can be caused by sensor or patient problems.
- Periodically check the CAPNOSTAT5 sensor and tubing for excessive moisture or secretion buildup.
- Do not operate the CAPNOSTAT5 sensor when it is wet or has exterior condensation.

Caution:

- Use only accessories provided by manufacturer.
- Do not sterilize or immerse the CAPNOSTAT5 sensor in liquids.
- Do not clean the CAPNOSTAT5 sensor and accessories except as directed in this manual.

- It is recommended that the CO₂ sensor be removed from the circuit whenever an aerosolized medication is delivered. This is due to the increased viscosity of the medications which may contaminate the sensor windows, causing the sensor to fail prematurely.
- Do not apply excessive tension to the CAPNOSTAT5 sensor cable.

P Note:

- This product and its accessories are latex free.
- After the life cycle of the CAPNOSTAT5 sensor and its accessories have been met, disposal should be accomplished following national and local requirements.
- Nitrous oxide, elevated levels of oxygen and helium can influence the CO2 measurement. Please setup gas compensation according to actual state.
- Barometric pressure compensation is required to meet the stated accuracy of the CAPNOSTAT5 sensor.
- Do not place the combined CO2 sensor between the ET tube and the elbow (pediatric or adult circuit), as this may allow patient secretions to block the adapter windows.
- Position the combined CO2 sensor with its windows in a vertical and not a horizontal position: this helps keep patient secretions from pooling on the windows.

5.9.5 Maintenance and cleaning

For CAPNOSTAT 5 Sensor and LoFlo Module

The outside of the module or sensor may be cleaned and disinfected by wiping with 70% isopropyl alcohol, a 10% bleach solution, or mild soap. After cleaning, wipe with a clean, water-dampened cloth to rinse. Dry before use.

For Reusable Airway Adapters

Reusable airway adapters may be cleaned by rinsing in a warm soapy solution, followed by soaking in a liquid disinfectant such as 70% isopropyl alcohol, a 10% bleach solution, Cidex® or System 1® (refer to the disinfectant manufacturer's instructions for use). Adapters should then be rinsed with sterile water and dried.

Reusable airway adapters may also be pasteurized or autoclaved. Autoclave at 121° C (250°F) for 20 minutes, unwrapped.

Before reusing the adapter, ensure the windows are dry and residue-free, and that the adapter has not been damaged during handling or by the cleaning process.

5.10 Measurement of AG

AG module is used to measure respiratory and anesthetic gases of a patient during anesthesia, including CO₂, N₂O, O₂, Halothane, Isoflurane, Enflurane, Sevoflurane and Desflurane. It is applicable for adult, pediatric and infant patients.

There are two methods for measuring anesthetic gas in the patient's airway:

■ Mainstream: Uses a AG sensor attached to an airway adapter directly inserted into the patient's breathing system.

■ Sidestream: Takes a sample of the respiratory gas with a constant sample flow from the patient's airway and analyzes it with the AG module.

Provide and Series an

5.10.1 Preparatory Steps for Measurement of AG

♦ Mainstream AG sensor

1. Plug the AG sensor connector into the AG connector on the monitor.

2. Attach AG sensor on the AG airway adapter. Shown as follows:



3. A green LED indicates that the AG sensor is ready for use. A blue LED indicates that may measurement of anesthetic gases.



4. Connect the 15 mm male connector of AG airway adapter to the breathing circuit Y-piece, and connect the 15mm female connector of AG airway adapter to the patient's endotracheal tube.



Alternatively, connect an HME (Heat Moisture Exchanger) between the patient's endotracheal tube and the AG sensor. Placing an HME in front of the AG sensor protects the airway adapter from secretions and effects of water vapor and eliminates the need of changing the adapter. It allows free positioning of the AG sensor as well.



5. Unless the AG sensor is protected with an HME always position the AG sensor with the indicating LED pointing upwards



Pre-use Check

- Check the reading on the monitor is correct or not before connecting the airway adapter to the breathing circuit.
- Always verify gas readings and waveforms on the monitor before connecting the airway adapter to the patient circuit.
- Perform the tightness check of the patient circuit with the AG sensor snapped on the AG airway adapter.

The state of the L	ED on the AC	sensor:
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Indication	Status
Steady green light	System OK
Blinking green light	Zeroing in progress
Steady blue light 1)	Anesthetic agent present
Steady red light	Sensor error
Blinking red light	Check airway adapter

Note 1: Valid for IRMA multi-gas sensors only.

• Sidestream AG module

- 1) Connect the AG module cable into the CO₂/AG socket on the left panel of the monitor..
- 2) Connect a Nomoline sampling line to the inlet port of the AG module.



3) Connect the gas sample exhaust port to a scavenging system or return the gas to the patient circuit.

- 4) Power up the patient monitor.
- 5) A green LED indicates that the AG module is ready for use.
- 6) Perform a pre-use check:
- ① Connect the sampling line to the inlet port of the AG module.
- ② Check that the AG module shows a steady green light (indicating that the system is OK)

(3) For AG module with O_2 option fitted: Check that the O_2 reading on the monitor is correct (21%).

④ Breathe into the sampling line and check that valid CO₂ waveforms and values are displayed on the monitor.

- ⑤ Occlude the sampling line with a fingertip and wait for 10 seconds.
- ⁽⁶⁾ Check that an occlusion alarm is displayed and that the AG module shows a flashing red light.

 \bigcirc If applicable: Perform a tightness check of the patient circuit with the sampling line attached.

Indication	Status
Steady green light	System OK
Blinking green light	Zeroing in progress
Steady blue light	Anesthetic agent present
Steady red light	Sensor error
Blinking red light	Check sampling line

The state of the LED on the AG module:

5.10.2 Setup of AG parameter



AA type Select the types of anesthetic gas, and options are **AA**, **HAL**, **ENF**, **ISO**, **SEV** and **DES**. After the monitor is turn on, if no AA types are configured, there will be a technical alarm prompting the configuration of AA and need to designate a kind of anesthetic gas. Considering safety, the configuration will not be saved after the monitor is switched off.

Label	Meanings	Label	Meanings
AA	Anesthetic agent	ISO	Isoflurane
HAL	Halothane	SEV	Sevoflurane
ENF	Enflurane	DES	Desflurane

Gas zero Conduct zero calibration on anesthesia gas module. Press this button, the following dialog will pop up. Select $\langle OK \rangle$ to conduct zero-calibration operation. If $\langle Cancel \rangle$ is selected, the zero-calibration will not be implemented.



Note: The zero-calibration of Gas is only valid on the AG module of PHASEIN Company.

 O_2 Module Turn on O_2 module . When choose ON, parameters of O_2 appear on the screen. When choose OFF, parameters of O_2 disappear.

Auto Color Select On or Off to enable or disable the auto color function for the gas.

Operate Mode Select the operate mode of the AG module, the options are **Sleep Mode**, **Work Mode** and **Self Test**.

Alarm Setup Click and open the dialog of anesthetic gas.



• **AA alarm** Click and open the dialog of AA alarm.



Adjust alarm Select this option to enter the configuration of alarm limits; conduct the configurations by turning the trim knob to select high and low limits and exit by selecting $\langle EXIT \rangle$. The upper part is the high limit and the lower one is the low limit.

EtAA alarm high limit, its configuration range is $0.0 \sim 30.0\%$ continuously adjustable, no lower than the low limit; the configuration range of EtAA alarm low limit is $0.0 \sim 30.0\%$ continuously adjustable, no higher than the high limit.

FiAA alarm high limit, its configuration range is $0.0 \sim 30.0\%$ continuously adjustable, no lower than the low limit; the configuration range of FiAA alarm low limit is $0.0 \sim 30.0\%$ continuously adjustable, no higher than the high limit.

Alarm Source Select the alarm source for AA parameters. When the selected parameter or one of the parameter exceed alarm limit, the monitor will give out alarm signal. The options are as follow:

 $\langle EtAA \rangle$: Only EtAA exceeds the alarm limit will trigger the alarm system.

(FiAA): Only FiAA exceeds the alarm limit will trigger the alarm system.

 $\langle Et/Fi \rangle$: EtAA or FiAA exceeds the alarm limit will trigger the alarm system. Alarm Switch Select the switch of AA alarm.

O₂ alarm Click and open the dialog of O_2 alarm.



Adjust alarm Select this option to enter the configuration of alarm limits. conduct the configurations by turning the trim knob to select high and low limits and exit by selecting $\langle EXIT \rangle$. The upper part is the high limit and the low one is the low limit.

FiO₂ alarm high limit, its configuration range is $18 \sim 100\%$ continuously adjustable, no lower than the low limit; the configuration range of FiO₂ alarm low limit is $18 \sim 100\%$ continuously adjustable, no higher than the high limit. FiO₂ alarm cannot be switched off, and when lower than 18% it will trigger high alarm.

EtO₂ alarm high limit, its configuration range is $10 \sim 100\%$ continuously adjustable, no lower than the low limit; the configuration range of EtO₂ alarm low limit is $10 \sim 100\%$ continuously adjustable, no higher than the high limit.

FiO₂ alarm Select \langle **ON** \rangle to enable FiO₂ over limit alarm; select \langle **OFF** \rangle to disable FiO₂ over limit alarm.

EtO₂ alarm Select \langle **ON** \rangle to enable EtO₂ over limit alarm; select \langle **OFF** \rangle to disable EtO₂ over limit alarm.



N₂O alarm Click and open the dialog of N_2O alarm.



Adjust alarm Select this option to enter the configuration of alarm limits; conduct the configurations by turning the trim knob to select high and low limits and exit by selecting $\langle EXIT \rangle$. The upper part is the high limit and the lower one is the low limit.

FiN₂O alarm high limit, its configuration range is $0 \sim 82\%$ continuously adjustable, no lower than the low limit; the configuration range of FiN₂O alarm low limit is $0 \sim 82\%$ continuously adjustable, no higher than the high limit.

EtN₂O alarm high limit, its configuration range is $0 \sim 100\%$ continuously adjustable, no lower than the low limit; the configuration range of EtN₂O alarm low limit is $0 \sim 100\%$ continuously adjustable, no higher than the high limit.

Alarm Source Select the alarm source for N₂O parameters. When the selected parameter or one of the parameter exceed alarm limit, the monitor will give out alarm signal. The options are as follow:

 $\langle EtN_2O \rangle$: Only EtN₂O exceeds the alarm limit will trigger the alarm system.

 $\langle FiN_2O \rangle$: Only FiN₂O exceeds the alarm limit will trigger the alarm system.

 $\langle Et/Fi \rangle$: EtN₂O or FiN₂O exceeds the alarm limit will trigger the alarm system.

Alarm Switch Select the switch of N₂O alarm.

5.10.3 Safety Information

◆ Mainstream AG sensor

Warning:

■ Disposable IRMA airway adapters shall not be reused. Reuse of the single use adapter can cause cross infection.

■ Used airway adapters shall be disposed of in accordance with local regulations for medical waste.

■ Use only PHASEIN manufactured oxygen sensor cells. Depleted oxygen sensors shall be disposed of in accordance with local regulations for batteries.

■ Do not use the IRMA Adult/Pediatric airway adapter with infants as the adapter adds 6 ml dead space to the patient circuit.

■ Do not use the IRMA Infant airway adapter with adults as this may cause excessive flow resistance.

■ Measurements can be affected by mobile and RF communications equipment. It should be assured that the IRMA probe is used in the electromagnetic environment specified in this manual.

■ Do not place the IRMA airway adapter between the endotracheal tube and an elbow as this may allow patient secretions to block the adapter windows and result in incorrect operation. Shown as follows:



■ To keep secretions and moisture from pooling on the windows or oxygen sensor port, always position the IRMA probe in a vertical position with the LED pointing upwards. Shown as follows:



■ Do not use the IRMA airway adapter with metered dose inhalers or nebulized medications as this may affect the light transmission of the airway adapter windows.

■ Do not try to open the oxygen sensor assembly. The oxygen sensor is a disposable product and contains a caustic electrolyte and lead.

■ The IRMA sensor is intended only as an adjunct in patient assessment. It must be used in conjunction with other assessments of clinical signs and symptoms.

- Replace the adapter if rainout/condensation occurs inside the airway adapter.
- Use only PHASEIN manufactured IRMA airway adapters.
- The IRMA sensor is not intended to be in patient contact.

Caution:

Do not apply tension to the sensor cable.

■ Do not operate the IRMA sensor outside the specified operating temperature environment.

■ Do not leave depleted oxygen sensors mounted in the IRMA sensor, even if the probe is not in use

■ Always disconnect the IRMA sensor from the monitor when not in use to prolong the lifetime of IRMA sensor.

■ The materials of patient breath tubing which is connected to the gas adapter, can't be anti-static and electric ones. Or it will be more dangerous when using HF electrosurgical equipments.

■ If error occurs in IRMA sensor, the indicating light will keep in red, and blink in red means the sensor is check the airway adapter.

■ The lifetime of the oxygen sensor is up to six months since its leaving factory. If it cannot work normally or the parameter cannot be accurate measured due to exceeding time limit, please replace the oxygen sensor timely.

■ If the AG airway adapter is detached from the sensor, or low voltage of oxygen sensor cell, or there is something wrong with the sensor, the prompting message may pop up on one of above conditions

• Sidestream AG module:

🂕 Warning:

■ Use only Nomoline sampling lines manufactured by PHASEIN.

■ The sidestream AG module must not be used with flammable anesthetic agents.

■ Carefully route the sampling line to reduce the risk of patient entanglement or strangulation.

■ Do not re-use disposable sampling lines.

■ Do not lift the monitor by the sampling line as it could disconnect from the monitor, casing the monitor to fall on the patient.

■ Used disposable sampling lines shall be disposed of in accordance with local regulations for medical waste.

■ Do not use adult/pediatric type sampling line configurations with infants, as this may add dead space to the patient circuit.

■ Do not use infant type sampling line configurations with adults, as this may cause excessive flow resistance.

Do not use the sidestream AG module with metered-dose inhalers or nebulized

medications as this may clog the bacteria filter.

• Check that the gas sample flow is not too high for the present patient category.

■ Measurements can be affected by mobile and RF communications equipment. Make sure that the sidestream AG module is used in the electromagnetic environment specified in this manual.

■ The sidestream AG module is intended only as an adjunct in patient assessment. It must be used in conjunction with other assessments of clinical signs and symptoms.

Replace the sampling line if the sampling line input connector starts flashing red, or a Nomoline occlusion message is displayed on the host monitor.

■ No modification of this equipment is allowed without authorization of the manufacturer. If this equipment is modified, appropriate inspection and testing must be conducted to ensure continued safe operation.

- The sidestream AG module is not designed for MRI environments.
- During MRI scanning, the monitor must be placed outside the MRI suite.

■ Operating high frequency electrosurgical equipment in the vicinity of the monitor may produce interference and cause incorrect measurements.

■ Too strong positive or negative pressure in the patient circuit might cause incorrect readings and internal damage.

Strong scavenging suction pressure might cause incorrect readings and internal damage.

• Exhaust gases should be returned to the patient circuit or a scavenging system.

Do not place the sidestream AG module in any position that might cause it to fall on the patient.

Caution:

Do not apply tension to the sidestream AG module cable.

Do not operate the sidestream AG module outside the specified operating temperature environment.

5.10.4 Zeroing

• Mainstream AG sensor:

In order to ensure the accuracy of gas measurement, zero reference calibration should be performed at regular intervals. Under the following conditions, it is necessary to perform zero reference calibration:

——The measured reading occurs error;

——A"GAS CONC. OUT OF RANGE" alarm message is displayed (When using CO₂ module, a "CO₂ CONC. OUT OF RANGE" alarm message is displayed);

——Airway adapter is replaced.

As following procedures:

- Snap a new AG airway adapter onto the AG sensor. Ensure that the airway adapter is not connected to the breath circuit of patient. The presence of ambient air (21% O₂ and 0% CO₂) in the AG airway adapter is very important.
- 2. Select 〈 **Gas Zero** 〉 in the AG parameter setting menu, this will start zeroing. The monitor zeroes the module and displays the "Zero In Progress" message for about 5s.
- 3. There will be a blinking green light on the AG sensor LED during 5s while Zero Reference calibration is in progress. Wait until the AG indicating lamp light on green and the reading of oxygen on the screen is "21". (When using CO₂ module, the reading of CO₂ on the screen is "0".) The message disappears upon completion of the zeroing.

Warning: Incorrect zero reference calibration will result in false gas readings.

Caution:

- User may only perform zero reference calibration under the instruction of the technical personnel authorized by manufacturer.
- Ensure the AG sensor is running more than 15min before replacing a new airway adapter. Please perform zero more than 2min after replacement.

• Sidestream AG module:

The sidestream AG module needs to establish a zero reference level for the CO₂, N₂O and anesthetic agent gas measurement. This zero calibration is here referred to as "zeroing".

The sidestream AG module performs zeroing automatically by switching the gas sampling from the respiratory circuit to ambient air. The automatic zeroing is performed every 24 hours, and takes less than 3 seconds for CO₂ module and less than 10 seconds for AG module.

If the sidestream AG module is fitted with an oxygen sensor, the automatic zeroing

will also include room air calibration of the oxygen sensor.

Warning: Since a successful zeroing requires the presence of ambient air (21% O₂ and 0% CO2) in the AG module, ensure that the AG module is placed in a well ventilated place. Avoid breathing near the sidestream AG module before or during the zeroing procedure.

5.10.5 MAC Calculation

Minimum alveolar concentration (MAC) is a standard for comparing the potency of inhalation anesthetics. The MAC value represents the alveolar concentration of an anesthetic (at one atmosphere) that, in 50 percent of a tested population, prevents gross muscular movement in response to a painful, standardized stimulus.

The MAC value may be calculated and displayed by using end-tidal (Et) gas concentrations according to the following formula:

$$MAC = \frac{\% Et(AA1)}{X(AA1)} + \frac{\% Et(AA2)}{X(AA2)} + \frac{\% Et(N2O)}{100}$$

X(AA): HAL=0.75%, ENF=1.7%, ISO=1.15%, SEV=2.05%, DES=6.0%

P Note:

■ Altitude, patient age and other individual factors are not considered in the formula above.

■ ET gas concentrations for secondary agent (AA2) is only available for IRMA AX+ sensors.

5.10.6 Maintenance and disinfecting

• Mainstream AG sensor:

The IRMA sensor can be cleaned using a cloth moistened with maximum 70% ethanol or maximum 70% isopropyl alcohol.

Remove the disposable IRMA Airway Adapter prior to cleaning the IRMA sensor.

Caution:

■ The IRMA airway adapters are non-sterile devices. Do not autoclave the devices as this will damage them.

• Never sterilize or immerse the IRMA sensor in liquid.

Sidestream AG module:

The ISA sidestream module should be cleaned on a regular basis.

Use a cloth moistened with max 70% ethanol or isopropyl alcohol to clean the module.

To prevent cleaning liquids and dust from entering the ISA sidestream module through

its inlet port, keep the Nomoline sampling line connected while cleaning the module.

Warning:

■ The Nomoline sampling lines are non-sterile devices. To avoid damage, do not autoclave any part of the sampling line.

• Never sterilize or immerse the ISA sidestream module in liquid.

Replacement of sampling line

The Nomoline sampling line is non-reusable.

Every two weeks or whenever "Sampling line clogged" appears, whichever comes first, the Nomoline sampling line should be replaced.

5.10.7 Maintenance

The ISA sidestream module is permanently factory calibrated. The module's stable design, results in no routine calibrations. If you want to conduct a maintenance task for the module, please send it to manufacturer.

5.10.8 PHASEIN Information

5.10.8.1 Patients

PHASEIN AB holds the following patents regarding products: SE519766; SE519779; SE523461; SE524086. Other patents pending.

5.10.8.2 Trademarks

PHASEIN IRMATM, PHASEIN ISATM, PHASEIN XTPTM, Sigma Multigas TechnologyTM, LEGITM, NomolineTM, IRMA EZ IntegratorTM, PHASEIN GasMasterTM and ISA MaintenanceMasterTM are trademarks of PHASEIN AB.

5.11 Measurement of ICG

5.11.1 General

Impedance cardiography (ICG) is a safe, non-invasive method to measure a patient's hemodynamic status. The ICG waveform is generated by thoracic electrical bioimpedance (TEB) technology, which measures the level of change in impedance in the thoracic fluid. Four small sensors send and receive a low amplitude electrical current through the thorax to detect the level of change in resistance in the thoracic fluid. With each cardiac cycle,

fluid levels change, which affects the impedance to the electrical signal transmitted by the sensors.

ICG uses an advanced algorithm to determine the level of change in impedance, to generate ICG waveform, and to calculate or derive hemodynamic parameters.

5.11.2 Precautions during ICG monitoring

Warning:

- Do not allow patients to wear ICG sensors when undergoing Magnetic Resonance Imaging (MRI) due to risk of severe burns.
- Ensure that the sensor gel, sensors or lead wires do not contact any other conductive materials (including earth-grounded materials) during patient monitoring.
- The ICG measurement is designed for assessment of most adult patients. (Height 122-229 cm and weight 30-159 kg)
- Do not conduct ICG monitoring on the patient with minute ventilation sensor function pacemaker (MV pacemaker), and the MV pacemaker function is enabled.
- The ICG sensor can only be applied to one patient at one time.

5.11.3 Effect factor

When patients present with the following conditions or anomalies may demonstrate reduced accuracy:

- Septic shock
- Aortic valve regurgitation
- Severe hypertension (Mean > 130 mmHg)
- Patient's height and weight exceed the specified range
- Connection to an intra-aortic balloon pump or chest tubes
- Patient movement, includes tremor
- Signal interference from patient cable or power wire connection
- The chest surgery that may change the blood and current in chest

5.11.4 Monitoring Procedure

Skin preparation

Good sensor-to-skin contact is important for a good ICG signal, as the skin is a poor

conductor of electricity. It is necessary to clean the patient's skin for sensors placement, as the following step:

1. Select sites with intact skin, without impairment of any kind.

2. Clip or shave hair from sites as necessary.

3. Gently abrade the skin and remove dead skin cells to improve the conductivity of the sensor site.

4. Wash sites thoroughly with soap and water, leaving no soap residue.

We do not recommend using ether or pure alcohol, because this dries the skin and increases the resistance.

5. Dry skin thoroughly.

Connect ICG cable

Plug the ICG cable into the ICG socket on the left panel of monitor (refer to Fig.1-5-1).

Place sensors on patient

In order to get good signals and accurate data, it is important to place ICG sensors on appropriate position, as follow:



- 1. A pair of sensors placed below the earlobe on each side of the neck.
- 2. Another pair of sensors placed on the midaxillary line at the xiphoid process level.
- 3. The two sensors must be placed right on the opposite position (180°) .

Input patient information

- 1. Select $\langle ICG \rangle$, open the $\langle ICG Setup \rangle$ menu;
- 2. Select (Input Info);

3. Set the parameters, including height, weight, sex, age, SYS, DIA, MEAN, CVP and PaoP.

The parameters (height, weight, sex, age) are associated with the parameters in the

patient info of system. If the parameters have been changed here, the parameters in the patient info of system alter at the same time, and vice versa.

There are default value of SYS, DIA, MEAN, CVP and PaoP, user may change them according to actual value.

5.11.5 ICG parameter Setup

	ICG Setup
ICG PARAM	
Input Info	
Alarm Setup	
Param Select	<u></u>
Scan Speed	25mm/s
UpdateRate	20
BeatAverage	20
Back to Main	
Show ICG Pa	aram.

ICG Param Click and open the window to show the current hemodynamic parameters of patient.

Input Info Click and open the dialog of patient's info input.

Alarm Setup Click and open the dialog of ICG alarm setup.

Param Select Click and open the dialog of parameter selection.

Scan Speed Select the scanning speed of ICG waveform, and the options are 6.25 mm/s, 12.5mm/s, 25mm/s.

Update Rate Select the interval of parameter updating, and the options are 5, 10, 20, 30, 40, 50, 60.

Beat Average Selects how many beats are averaged to determined displayed data, and the options are 5, 10, 20, 30, 40, 50, 60.

■ ICG Parameter



Note on the hemodynamic parameters

Abbreviation	Unit	Full name
C.O.	L/min	Cardiac output
C.I.	L/min/m ²	Cardiac index
SV	mL	Stroke volume
SI	mL/m ²	Stroke index
SVR	DS/cm ⁵	Systemic vascular resistance
SVRI	$DS \cdot m^2/cm^5$	Systemic vascular resistance index
TFI	Ω	Thoracic fluid index
TFC	/kΩ	Thoracic fluid content
HR	bpm	Heart rate

The HR reading is not obtained form ECG module, and it is obtained from ICG module.

Input Patient Info



Sex Gender of patient, female or male.

Age Age of patient. Turn the trim knob with an increment or decrement of 1 year.

Height Patient's height. Turn the trim knob with an increment or decrement of 1cm.

Weight Patient's weight. Turn the trim knob with an increment or decrement of 1kg.

SYS Systolic pressure of patient. Turn the trim knob with an increment or decrement of 1mmHg.

DIA Diastolic pressure of patient. Turn the trim knob with an increment or decrement of 1mmHg.

MEAN Mean pressure of patient. Turn the trim knob with an increment or decrement of 1mmHg.

CVP Cardiac venous pressure of patient. Turn the trim knob with an increment or decrement of 1mmHg.

PaoP Pulmonary artery occluded pressure of patient. Turn the trim knob with an increment or decrement of 1mmHg.

Alarm Setup



Adjust Alarm Select this option to enter the configuration of alarm limits, conduct the configuration by turning the trim knob to select high or low limit and exit by selecting **EXIT**. The upper part is the high limit and the lower one is the low limit.

Alarm Source Select the alarm source for ICG parameters. When the selected parameter or one of the parameter exceed alarm limit, the monitor will give out alarm signal. The options are as follow:

 $\langle C.I. \rangle$: Only C.I. exceeds the alarm limit will trigger the alarm system.

 $\langle TFC \rangle$: Only TFC exceeds the alarm limit will trigger the alarm system.

 $\langle C.I./TFC \rangle$: C.I. or TFC exceeds the alarm limit will trigger the alarm system.

Alarm Switch Select the switch of ICG alarm.

Param Select

Param Select
SelectParamI SVRI
SelectParamII TFC
Previous
Select Param I.

Param1 Select the parameter to display on the area1, and the options are C.O., C.I., SV, SI, SVR, SVRI, TFI, TFC, HR.

Param2 Select the parameter to display on the area1, and the options are C.O., C.I., SV, SI, SVR, SVRI, TFI, TFC, HR.

5.11.6 Maintenance and cleaning

The maintenance and cleaning of ICG cable is the same as ECG cable.

Chapter 6 Alarm

This chapter gives general information about the alarm and corresponding remedies.

Note: The equipment generates all the auditory and visual alarms through speaker, LED and screen.

6.1 Alarm Priority

There are two kinds of alarms, defined as physiological alarm and technical alarm. Physiological alarms refer to those alarms triggered by patient's physiological situation which could be considered dangerous to his or her life, such as SpO₂ exceeding alarm limit (parameter alarms). Technical alarms refer to system failure, which can make certain monitoring process technically impossible or make monitoring result unbelievable. General alarm belongs to those situations that cannot be categorized into these two cases but still need to pay some attention. Each alarm, either technical or physiological, has it's own priority.

Alarms in the monitor are divided into three priorities, that is: high priority, medium priority and low priority.

- High priority alarm indicates the patient's life is in danger. It is the most serious alarm.
- Medium priority alarm means serious warning.
- Low priority alarm is a general warning.

Only alarm priority of parameters exceeding limits alarm can be modified by the user, the other alarm priorities of physiological and technical alarms are preset by the system and they can not be changed by the user.

6.2 Alarm Modes

When alarm occurs, the monitor may raise the user's attention in two ways, which are auditory prompt, visual prompt and description. Visual prompt is given by alarm indicating lamp and screen of the monitor, auditory prompt is given by speaker in the device. Physiological alarm information is displayed in the Physiological Alarm area. Most of technical alarm information is displayed in the Technical Alarm area. Technical alarms related to NIBP measurement are displayed in the NIBP parameter area. The Physiological Alarm area is on the upmost right part of the screen. The Technical Alarm area is to the left side of the Physiological Alarm area.

The alarm sound and visual display comply with clause 201.3.2 of the standard IEC 60601-1-8.

Note: The concrete presentation of each alarm prompt is related to the alarm priority.

Alarm Sound

The high/medium/low-level alarms are indicated by the system in following different audio ways:

Alarm level	Audio prompt		
High	Mode is "DO-DO-DO-DO-DO, DO-DO-DO-DO", which is triggered once every 10 seconds.		
Medium	Mode is "DO-DO-DO", which is triggered once every 25 seconds.		
Low	Mode is "DO-", which is triggered once every 25 seconds.		

🖃 Alarm Lamp

For M8000A&M9000A patient monitor:

When physiological alarm occurs, the physiologic alarm lamp lights according to the alarm level. Shown as follows:

Alarm level	Visual prompt
High	Alarm indicating lamp flashes in red with 2 Hz.
Medium	Alarm indicating lamp flashes in yellow with 0.5 Hz.
Low	Alarm indicating lamp lights on in yellow.

When technical alarm occurs, the technical alarm lamp lights on in blue.

For M8000&M9000 patient monitor:

The high/medium/low-level alarms are indicated by the system in following different visual ways:

Alarm level	Visual prompt
High	Alarm indicating lamp flashes in red with 2 Hz.
Medium	Alarm indicating lamp flashes in yellow with 0.5 Hz.
Low	Alarm indicating lamp lights on in yellow.

Screen Display

Physiological alarm: The parameter, which triggers the alarm, splashes in the frequency of 2Hz on the screen. The physiological alarm area on the screen displays alarm message, and red indicates high priority alarm, yellow indicates medium or low priority alarm.

When Technical alarm or General alarm occurs, the Technical alarm area displays alarm message, red indicates high priority alarm, yellow indicates medium or low priority alarm, cyan indicates general message.

Note: When alarms of different priorities occur at the same time, the monitor prompts the one of the highest priority.

6.3 Alarm Setup

📼 Set Alarm volume

Step 1: Select <Alarm Volume> item in Menu: <MENU> \rightarrow <Alarm Setup> \rightarrow <Alarm Volume>.

Step 2: Set < Alarm Volume > item to <Off>, <1>, <2>, <3>.

■ Set alarm limits of physiological parameters

The alarm limit of each physiological parameter can be set in its menu, and they are continuous in alarm range. For example:

ECG alarm setup:

Step 1: Select Menu < ECG>

Step 2: Configure the following parameters related to ECG alarm, <HR LO> and <HR HI>.

Please refer to above operation for Methods of Alarm setup of the other parameters

It is important to set physiological alarm limits properly. The monitor can't give medicinal alarm prompt in clinical application with improper setting of physiological alarm limit.

The physiological alarm occurs when the measurement exceeds the set parameter limits. Please refer to above operation for Methods of alarm setup of the other parameters.
ECG Alarm configuration



Alarm levels configuration



Alarm recording configuration

Alarm Record	Alarm R	ecord		
Alarm Record	EXIT			
All OFF	ECG	Sp02	NIBP	Temp
Default Setup	OFF	OFF	OFF	OFF
Previous	IBP	C02	Ges	ARR
	OFF	OFF	OFF	UFF
Set record on alarm event.				

Alarm indication of physiological parameters

Audio: when alarm occurs, the system generates alarm sound to raise the user's attention (audio alarm can be disabled).

Visual: The parameter flashes on the display area of the screen and alarm LED lights.

Warning: The lower limit and the upper limit of parameter must be set based on clinical practices and general clinical experiences.

Proof Note: When parameter alarm level is off, alarm will be disabled, even if the measurement results exceed the limits. Alarm indicating lamp in the front of the monitor will alarm at the highest level, if different levels alarms coexist.

6.4 Alarm Cause

Alarm of the monitor includes:

- 1. Physiological Alarm
- 2. Technical Alarm
- 3. General Prompt
- Physiological Alarm

When the measuring value has exceeded the set parameter limit and its <ALM LEV> is not <OFF>, the monitor alarms. The monitor wouldn't alarm with absence of either of the two conditions.

Technical Alarm

Once system fault occurs, the monitor will alarm immediately and trigger corresponding operations, such as stop displaying values and waveforms, erase the last screen to avoid misleading. The screen displays more than one fault message by alterative.

General Prompt

Sometimes there are alarms similar to Technical Alarms but can be considered as normally. The condition, which triggers this kind of alarm wouldn't bring danger to the patient.

6.5 Silence/Suspension

SILENCE

Press the < SUSPEND/SILENCE > button on the front panel for more than 2 seconds can shut off all sounds until the < SUSPEND/SILENCE> button is pressed again. When the system is in SILENCE status, any newly generated alarm will cancel the SILENCE status and make the system back to normal status.

When in the SILENCE status, the icon 💥 will be displayed in the left upper of the screen.

SUSPENSION

Press the < SUSPEND/SILENCE> button on the front panel for less than 2 seconds can close all audio and visual prompt and description about all the physiological alarms and to make the system enter ALARM PAUSE status. The rest seconds for alarm pause is displayed in the Physiological Alarm area. And the symbol is displayed in the System Prompt area.

The time for Alarm Suspension is 2 minutes.

When in the PAUSE status, press the < SUSPEND/ SILENCE> button again to restore the normal alarm status. Besides, during PAUSE status, newly occurring technical alarm will cancel the PAUSE status and the system will come back to the normal alarm status. The symbol is disappears, too.

Solution Note: Whether an alarm will be reset depends on the status of the alarm cause. But by pressing <SUSPEND/SILENCE> button can permanently shut off audio sound of Lead Off or Sensor Off alarms.

6.6 Parameter Alarm

You can check and set the parameters' alarm limit in their menus or in the alarm setup window in main setup menu.

If you want to set the alarm state of parameters, enter the machine maintenance state please.

For the parameters whose alarm switch is set to ON, the alarm will be triggered when at least one of them exceeds alarm limit. The following actions take place:

- 1. Alarm message displays on the screen as described in alarm mode;
- 2. The monitor beeps in its corresponding alarm class and volume;
- 3. If alarm recording is on, the recorder starts alarm recording at set interval.

6.7 When an Alarm Occurs

Note: When an alarm occurs, you should always check the patient's condition first.

Check the alarm message appeared on the screen. It is needed to identify the alarm and act appropriately, according to the cause of the alarm.

- 1. Check the patient's condition.
- 2. Identify which parameter is alarming or which kind of alarm it is.
- 3. Identify the cause of the alarm.
- 4. Silence the alarm, if necessary.
- 5. When cause of alarm has been over, check that the alarm is working properly.

6.8 Alarm Description and Prompt

6.8.1 ECG Alarm

Message	Cause	Alarm Level
HR too high	HR measuring value is above the upper	User-selectable
	alarm limit	
HR too low	HR measuring value is below the lower	User-selectable
	alarm limit	

Message	Cause	Alarm Level
ECG RA LA LL V- LEAD	ECG electrode fall off the skin or ECG	Low
OFF	cables fall off the monitor	LOW
ECG electrode polarized	ECG electrode polarized	Low
ECC communication owner	ECG measurement failure or communication	Low
ECG communication error	failure	LOW
HR alarm error	Alarm failure	Low

Technical Alarm:

6.8.2 RESP Alarm

Physiological Alarm:

Message	Cause	Alarm Level
RR too high	RR measuring value is above the upper alarm limit	User-Selectable
RR too low	RR measuring value is below the lower alarm limit	User-Selectable
RESP Apnea	No signal for breath in specific interval	User-Selectable

Technical Alarm:

Message	Cause	Alarm Level
RR alarm error	Alarm failure	Low

6.8.3 SpO2 Alarm

Physiological Alarm:

Message	Cause	Alarm Level
Sec. to a high	SpO ₂ measuring value is above the upper	Medium ,High
SpO ₂ too nign	alarm limit	User-Selectable
Sec. to a large	SpO ₂ measuring value is below the lower	Medium ,High
SpO ₂ too low	alarm limit	User-Selectable
PR too high	PR measuring value is above the upper alarm limit	User-Selectable
PR too low	PR measuring value is below the lower alarm limit	User-Selectable

Technical Alarm:

Message	Cause	Alarm Level
SpO ₂ sensor off	SpO ₂ sensor may be disconnected from the patient or the monitor	Low
SpO ₂ communication error	SpO ₂ measurement failure or communication error	Low

SpO ₂ alarm error	Alarm failure	Low
PR alarm error	Alarm failure	Low
SpO ₂ sensor failure	SpO ₂ sensor failure	Low
SpO2 pulse timeout	Search pulse too long	High
SpO2 Interference Detected	SpO2 module detects the interference.	Low
SpO ₂ No sensor Connected	No SpO2 sensor connected	Low
SpO2 No Cable	No cable connected	Low
SpO2 Sensor Calibrate	SpO2 sensor calibrate	Low
SpO2 has R wave	SpO2 wave has R wave	Low
SpO2 Defective Sensor	SpO2 sensor is defective	Low
SpO2 Too Much Ambient	There are too much ambient light.	Low
Light		
SpO2 Low Signal IQ	SpO2 signal IQ is low	High
SpO2 Unrecognized Sensor	SpO2 sensor is unrecognized.	Low
SpO2 Processing Active	SpO2 processing is active.	Low
SpO2 No Tape	SpO2 no tape	Low
SpO2 Demo	The monitor is at demo mode	Low
SpO2 Sensor Fault	SpO2 sensor has some fault	Low
SpO2 Board Fault	SpO2 board has some fault	Low
SpO2 Low Perfusion	SpO2 low perfusion	Low
SpO2 Diagnostic Failure	SpO2 diagnostic failure	Low

Prompt:

Message	Cause	Alarm Level
Search pulse	SpO ₂ module is searching for pulse	No alarm
Motion interference	Patient movement too much.	No alarm
Disconnected	SpO2 sensor may be disconnected form the monitor.	No alarm

6.8.4 TEMP Alarm

Message	Cause	Alarm Level
TEMP1 too high	TEMP1 measuring value is above upper alarm limit	User-Selectable
TEMP1 too low	TEMP1 measuring value is below lower alarm limit	User-Selectable

TEMP2 too high	TEMP2 measuring value is above upper alarm limit	User-Selectable
TEMP2 too low	TEMP2 measuring value is below lower alarm limit	User-Selectable

Technical Alarm:

Message	Cause	Alarm Level
TEMP1 sensor off	TEMP1 sensor may be disconnected from user or monitor	Low
TEMP2 sensor off	TEMP2 sensor may be disconnected from user or monitor	Low
TMEP communication error	TEMP measurement error or communication error	Low
TMEP1 alarm error	Alarm failure	Low
TEMP2 alarm error	Alarm failure	Low
T1 over measuring range	TEMP1 over measuring range	Low
T1 below measuring range	TEMP1 below measuring range	Low
T2 over measuring range	TEMP2 over measuring range	Low
T2 below measuring range	TEMP2 below measuring range	Low
TEMP Self checking error	TEMP calibration failure	Low

6.8.5 IBP Alarm

Message	Cause	Alarm Level
IBP SYS1 too high	SYS measuring value of channel 1 is above upper alarm limit	User-Selectable
IBP SYS1 too low	SYS measuring value of channel 1 is below lower alarm limit	User-Selectable
IBP DIA1 too high	DIA measuring value of channel 1 is above upper alarm limit	User-Selectable
IBP DIA1 too low	DIA measuring value of channel 1 is below lower alarm limit	User-Selectable
IBP MAP1 too high	MAP measuring value of channel 1 is above upper alarm limit	User-Selectable
IBP MAP1 too low	MAP measuring value of channel 1 is below lower alarm limit	User-Selectable
IBP SYS2 too high	SYS measuring value of channel 2 is above upper alarm limit	User-Selectable
IBP SYS2 too low	SYS measuring value of channel 2 is below lower alarm limit	User-Selectable

IBP DIA2 too high	DIA measuring value of channel 2 is above upper alarm limit	User-Selectable
IBP DIA2 too low	DIA measuring value of channel 2 is below lower alarm limit	User-Selectable
IBP MAP2 too high	MAP measuring value of channel 2 is above upper alarm limit	User-Selectable
IBP MAP2 too low	MAP measuring value of channel 2 is below lower alarm limit	User-Selectable

Technical Alarm

Message	Cause	Alarm Level
IBP1 sensor off	IBP cable of channel 1 falls off from monitor	Low
IBP2 sensor off	IBP cable of channel 2 falls off from monitor	Low
IBP communication error	IBP communication error	Low
IBP1 alarm error	Alarm failure	Low
IBP2 alarm error	Alarm failure	Low

Prompt:

Message	Cause	Alarm Level
IBP1 Checking	IBP1 zero calibration is in progress.	No alarm
IBP1 Errlose	IBP1 zero calibration failed for IBP1 cable falls off.	
IBP1 Errtimeout	IBP1 zero calibration failed for time is out.	
IBP1 Check OK	IBP1 zero calibration succeed.	
IBP2 Checking	IBP2 zero calibration is in progress.	No alarm
IBP2 Errlose	IBP2 zero calibration failed for IBP2 cable falls off.	
IBP2 Errtimeout	IBP2 zero calibration failed for time is out.	
IBP2 Check OK	IBP2 zero calibration succeed.	

6.8.6 NIBP Alarm

Message	Cause	Alarm Level
NIBP SYS too high	NIBP SYS measuring value is above upper alarm limit	User-Selectable
NIBP SYS too low	NIBP SYS measuring value is below lower alarm limit	User-Selectable

NIBP DIA too high	NIBP DIA measuring value is above upper alarm limit	User-Selectable
NIBP DIA too low	NIBP DIA measuring value is below lower alarm limit	User-Selectable
NIBP MAP too high	NIBP MAP measuring value is above upper alarm limit	User-Selectable
NIBP MAP too low	NIBP MAP measuring value is below lower alarm limit	User-Selectable

Technical Alarm 1(display in description area):

Message	Cause	Alarm Level
NIBP communication error	NIBP measurement failure or communication failure	Low
NIBP SYS alarm error	Alarm failure	Low
NIBP DIA alarm error	Alarm failure	Low
NIBP MAP alarm error	Alarm failure	Low

Technical Alarm 2(display in description area below NIBP mean arterial pressure value):

Message	Cause	Alarm Level
SELF-TEST FAILED	Transducer or other hardware failure.	Low
LOOSE CUFF	a. Cuff is completely unwrapped.b. The cuff is not connected.c. Adult cuff used in neonate mode.	Low
AIR LEAK	Air leak in pneumatics, hose, or cuff.	Low
AIR PRESSURE ERROR	Unable to maintain stable cuff pressure, e.g. kinked hose.	Low
WEAK SIGNAL	a. Very weak patient signal due to a loosely wrapped cuff.b. The pulse of patient is too weak.	Low
RANGE EXCEEDED	Measurement range exceeds module specification.	Low
EXCESSIVE MOTION	 a. Too many retries due to interference of motion artifact. b. Signal is too noisy during measurement, e.g. patient has severe tremor. c. Irregular pulse rate, e.g. arrhythmia. 	Low
OVERPRESSURE SENSED	Cuff pressure exceeds the specified upper safety limit. Could be due to rapid squeezing or bumping of cuff.	Low
SIGNAL SATURATED	Large motion artifact that saturates the BP amplifier's amplitude handing capability.	Low
AIR SYSTEM LEAK	Module reports Air Leakage failure while in the Pneumatic Test mode.	Low
SYSTEM FAILURE	Module occurs abnormal processor event.	Low

TIME OUT	Measurement took more than 120 seconds in adult, 90 seconds in neonate mode.	Low
CUFF TYPE ERR	Neonate cuff used in adult mode.	Low

Prompt (display in description area below NIBP mean arterial pressure value):

Message	Cause	Alarm Level
NIBP Resetting	NIBP measurement module is resetting	
Over Press Testing	NIBP is testing Over-Pressure	Na alarma
Manometer Testing	NIBP is testing Manometer	ino alarini
Pneumatic Testing	NIBP is testing Pneumatic	

6.8.7 System Alarm and Prompt

Technical Alarm

Message	Cause	Alarm Level
Battery failure	Battery failure or no battery	Low
Battery low	Voltage of battery is too low	Medium
Key error	Keyboard error	Low
Recorder error	No paper in the recorder when recording or the recorder door is open or recorder is absent	Low

Prompt

Message	Cause	Alarm Level
Recording	Recorder is in printing operation	No alarm

6.8.8 CO2Alarm (CPT module)

Physiological Alarm:

Message	Cause	Alarm Level
EtCO ₂ too high	EtCO ₂ measuring value is above upper alarm limit	User-Selectable
EtCO ₂ too low	EtCO ₂ measuring value is below lower alarm limit	User-Selectable

Technical Alarm:

Message	Cause	Alarm Level
CO ₂ sensor off	CO ₂ sensor off patient or off the monitor	Low
CO ₂ communication error	CO ₂ module failure or communication failure	Low
CO2 alarm error	Co2 alarm function failure	Low

CO ₂ measurement Over range	CO ₂ measurement Over range, need verify zero	Medium
CO ₂ sensor error	CO ₂ sensor error	Medium

6.8.9 CO2Alarm (LoFlo module, CAPNOSTAT5 module)

Physiological Alarm:

Message	Cause	Alarm Level
EtCO ₂ Hi	EtCO ₂ measuring value is above upper alarm	User-Selectable
EtCO ₂ Lo	EtCO ₂ measuring value is below lower alarm	User-Selectable
FiCO ₂ Hi	FiCO ₂ measuring value is above upper alarm	User-Selectable
FiCO ₂ Lo	FiCO ₂ measuring value is below lower alarm	User-Selectable
Apnea	No breath detected in the set period	User-Selectable

Technical Alarm:

Message	Cause	Alarm Level
Sensor Over Temp	Sensor over temperature.	High
Sensor Faulty	Sensor error	High
Check Sampling Line	Sampling line blockage or damage;	Low
	Sampling line is kinked or pinched;	
	Exhaust tube is blocked.	
Zero Required	Negative CO_2 detected; the module needs to be zeroed.	High
CO ₂ Out of Range	The calculated CO ₂ value is out of range.	Low
Check adapter	The adapter is removed from the module.	Low
Sensor no initialized	Sensor or module is not initialized	Low

Prompt:

Message	Cause	Alarm Level
Zero in Progress	Zeroing is in progress.	No Alarm
Sensor Warm Up	Module is warming up.	No Alarm

6.8.10 AG alarm and promotion

Message	Cause	Alarm Level
EtAA too high	EtAA is above upper alarm limit	User Selectable
EtAA too low	EtAA is below lower alarm limit	User-selectable
FiAA too high	FiAA is above upper alarm limit	User-Selectable

FiAA too low	FiAA is below lower alarm limit	
EtN ₂ O too high	EtN ₂ O is above upper alarm limit	
EtN ₂ O too low	EtN ₂ O is below lower alarm limit	User-Selectable
FiN ₂ O too high	FiN ₂ O is above upper alarm limit	User-Selectable
FiN ₂ O too low	FiN ₂ O is below lower alarm limit	
EtO ₂ too high	EtO ₂ is above upper alarm limit	User-Selectable
EtO ₂ too low	EtO ₂ is below lower alarm limit	User-Selectable
FiO ₂ too high	FiO ₂ is above upper alarm limit	User-Selectable
FiO ₂ too low	FiO ₂ is below lower alarm limit	

Technical Alarm:

Message	Cause	Alarm Level
GAS communication error	GAS module failure or communication error	Medium
Check Airway Adapter	Airway adaptor of GAS module disconnected with sensor	Medium
Replace O ₂ sensor	Oxygen sensor disconnected with module	Medium
O ₂ sensor low	Weak oxygen sensor signal	Medium
GAS sensor error	GAS sensor error	Low
GAS CONC. Out of Range	Measurement of GAS module over range	Medium
Room Air Calibration Required	Measurement of oxygen density is not correct.	High
Sampling line Clogged	Sampling line is occluded or damaged; Sampling tube is kinked or pinched.	Medium
No sampling line	Sampling line disconnected with the AG module.	Low
O ₂ Sensor Err	O ₂ Sensor error	Low
Last Span Cal. command fail	The command of the last span calibration failed.	Low

Prompt messages:

Messages	Cause	Level
Span Cal. in progress	Span calibration is in progress.	No alarm
O2 Span Calibration Required	The O2 span needs to be calibrated.	No alarm
Mixed Agents	More than one type of agent exist	No alarm

6.8.11 ICG alarm and promotion

Physiological Alarm:

Message	Cause	Alarm Level
C.I. High	C.I. measuring value is above the high alarm limit	
C.I. Low	C.I. measuring value is below the low alarm limit	
TFC High	TFC measuring value is above the high alarm limit	User-selectable
TFC Low	TFC measuring value is below the low alarm limit	

Technical Alarm:

Message	Cause	Alarm Level
ICG sensor off	ICG sensor may be detached from the patient or monitor	Low

Chapter 7 Recording

Monitor carries out the recording function by the built-in recorder.

Alarm recording

Monitor provides the function of alarm trigger recording. To make alarm recording available, Please keep <**Alarm Record** >of <**Recorder setup**> of <**System setup**> in <**Monitor setup**> menu is **ON**, and adjust alarm level of alarm parameter to non-close. If any monitoring parameter exceeds the limit and <**Alarm Record**> is **ON**, recorder will print all monitoring parameter values in the alarm time. Moreover, if monitor alarms continuously, recorder will print every two minutes.

■ Auto recording

Monitor has the function of Auto recording. To make Auto recording available, user can adjust **<Record Interval**> of **<Recorder Setup**> of **<System Setup**> in **<Monitor Setup**> to a necessary interval time. All monitoring parameter values and waveforms will be recorded automatically according to the determined period.

Real-Time recording

Monitor has the function of real time recording. If **<FREEZE/RECORD>** key in the front panel has been pressed over 2 seconds, the waveform and data of cardiac electro and SpO₂ can be recorded in real time. If **<FREEZE/RECORD>** pressed again, real time recording will end. The lead ECG waveform (determined by **<Record Wave>** in **<Recorder Setup>**) will be monitoring in emphasis, when ECG waveforms are being recorded.

Note: During real time recording, three waveforms can be recorded at the same time. Users can configure the waveforms according to need. Please refer to chapter 4.2.1. Measurement parameter values of individual module have been recorded on the top of waveforms.

Chapter 8 The Maintenance and Cleaning

8.1 System Check

An effective maintenance schedule should be established for your monitoring equipment and reusable supplies. This should include inspection as well as general clearing on a regular basis. The maintenance schedule must comply with the policies of your institution's infection control unit and/or biomedical department.

Check with your biomedical department to be sure preventive maintenance and calibration has been done. The User Maintenance Instruction contains detailed information.

Before using the monitor, check the equipment following these guidelines:

- Check the equipment for obvious mechanical damage.
- Check all the outer cables, inserted modules and accessories for fraying or other damage. Qualified service personnel should repair or replace damaged or deteriorated cables.
- Check all the functions relevant to patient monitoring, make sure that the monitor is in good condition.

If you find any damage on the monitor, stop using the monitor on patient, and contact the biomedical engineer of the hospital or Manufacturer's Customer Service immediately.

Note: Refer to the User Maintenance Instruction for more comprehensive checkout procedures.

The overall check of the monitor, including the safety check, should be performed only by qualified personnel once every 6 to 12 month, and whenever the monitor is fixed up.

- Inspect the safety relevant labels for legibility.
- Verify that the device functions properly as described in the instructions for use.
- Test the protection earth resistance according IEC 60601-1:1995, Limit 0.10hm.
- Test the earth leakage current according IEC 60601-1:1995, Limit: NC 500uA, SFC 1000uA.
- Test the patient leakage current according IEC 60601-1:1995, Limit: 100uA(BF), 10uA(CF).
- > Test the patient leakage current under single fault condition with mains voltage on

the applied part according IEC 60601-1:1995, Limit: 5mA(BF), 50uA(CF).

The leakage current should never exceed the limit. The data should be recorded in an equipment log. If the device is not functioning properly or fails any of the above tests, the device has to be repaired.

The synchronism of the defibrillator should be checked by in the frequency described in the hospital regulations. At least every 3 months, it should be checked by the biomedical engineer of the hospital or qualified service technician.

All the checks that need to open the monitor should be performed by qualified service technician. The safety and maintenance check can be conducted by persons from the manufacturer. You can obtain the material about the customer service contract from the local office.

The circuit diagrams, parts lists and calibration instructions of the patient monitor can be provided by the manufacturer.

Warning: If the hospital or agency that is responding to using the monitor does not follow a satisfactory maintenance schedule, the monitor may become invalid, and the human health may be endangered.

Solution Note: To ensure maximum battery life, please ensure that the battery is fully charged when you are keeping the device in storage for an extended period of time, and then take out the battery.

Warning: Refer the battery replacement only to manufacturer's service technician.

8.2 Battery Maintenance

A rechargeable and maintenance-free battery is designed for Patient Monitor, which enables continuous working when AC power off. Special maintenance is not necessary in the normal situation. Please pay attention to the followings in using for more durable usage and a better capability.

■ Operate the patient monitor in the environment according to the instruction.

 \blacksquare Use AC power for the patient monitor when available.

■ Recharge the battery sooner when it is off. The volume of battery will not be charged to what it should be, when the battery has not been charged for a long time.

■ If the monitor is not used for long time, the AC power should be plugged in until the battery is fully recharged, then take out the battery, so that the service life of the battery will not be shortened.

 \blacksquare Avoid exposed and sun shine.

■ Avoid infrared and ultraviolet radiation.

■ Avoid moist, dust and erosion from acid gas.

For Lithium ion battery:

A lithium ion battery needs at least two conditioning cycles when it is put into use for the first time. A battery conditioning cycle is one complete, uninterrupted charge of the battery, followed by a complete, uninterrupted discharge of the battery. A lithium ion battery should be conditioned regularly to maintain its useful life. Condition a battery once when it is used or stored for two months, or when its run time becomes noticeably shorter.

To condition a lithium ion battery, follow this procedure:

1. Disconnect the monitor from the patient and stop all monitoring and measuring procedures.

2. Place the lithium ion battery in need of conditioning into battery compartment of the monitor.

3. Connect the monitor to the AC mains. Allow the battery to be charged uninterruptedly for above 6 hours.

4. Remove the AC mains and allow the monitor to run from the battery until it shuts off.

5. Reconnect the monitor to the AC mains. Allow the battery to be charged uninterruptedly for above 6 hours.

Now the battery is conditioned and the monitor can be returned to service.

8.3 General Cleaning

Warning: Before cleaning the monitor or the sensors, make sure that the equipment is switched off and disconnected from the power line.

The Patient Monitor must be kept dust-free.

Regular cleaning of the monitor shell and the screen is strongly recommended. Use only non-caustic detergents such as soap and water to clean the monitor shell.

Please pay special attention to the following items:

- 1. Avoid using ammonia-based or acetone-based cleaners such as acetone.
- 2. Most cleaning agents must be diluted before use. Follow the manufacturer's

directions carefully to avoid damaging the monitor.

- 3. Don't use the grinding material, such as steel wool etc.
- 4. Don't let the cleaning agent enter into the chassis of the system.
- 5. Don't leave the cleaning agents at any part of the equipment.

8.4 Cleaning Agents

Examples of disinfectants that can be used on the instrument casing are listed below:

- Diluted soap solution
- Diluted Ammonia Water
- Diluted Sodium Hypochlorite (Bleaching agent).

Note: The diluted sodium hypochlorite from 500ppm (1:100 diluted bleaching agent) to 5000ppm (1:10 bleaching agents) is very effective. The concentration of the diluted sodium hypochlorite depends on how many organisms (blood, mucus) on the surface of the chassis to be cleaned.

- Hydrogen Peroxide 3%
- Alcohol 70%
- Isopropyl alcohol 70%

The surface of patient monitor can be cleaned with hospital-grade ethanol and dried in air or with crisp and clean cloth.

The manufacturer has no responsibility for the effectiveness of controlling infectious disease using these chemical agents. Please contact infectious disease experts in you hospital for details.

8.5 Disinfection

To avoid extended damage to the equipment, disinfection is only recommended when stipulated as necessary in the Hospital Maintenance Schedule. Disinfection facilities should be cleaned first.

Appropriate disinfection materials for ECG leads, SpO₂ sensor, blood pressure cuff, TEMP probe, CO₂ sensor, AG sensor and ICG cable are introduced in the corresponding chapters respectively.

Warning: Do not use EtO gas or formaldehyde to disinfect the monitor.

Chapter 9 Accessories

This chapter lists the recommendation accessories used in this device.

Warning: The accessories listed below are specified to be used in this device. The device will be possibly damaged or lead some harm if any other accessories are used.

1. ECG

ECG Electrode

Туре	Patient category	PN
Disposable	Adult	15-100-0008
	Pediatric/ Neonatal	15-100-0009

ECG Cable

Туре	Description	Standard	PN
Snap	5-lead	IEC	15-100-0002
Snap	3-lead	IEC	15-100-0004
Snap	5-lead	AHA	15-100-0001
Snap	3-lead	AHA	15-100-0003
Snap	Chest lead	AHA	15-100-0005

2. SpO2

Nellcor SpO₂ Sensor

Туре	Model	Patient category	
MAX-A		Adult finger (patient size>30kg)	
	MAX-P	Pediatric foot/hand (patient size 10-50kg)	
Disposable	MAX-I	Infant foot/hand (patient size 3-20kg)	
	MAX-N	Adult finger or neonatal foot/hand (patient size >40 kg or <3 kg)	
	DS-100A	Adult	
Reusable	OXI-A/N	Adult / neonatal	
	OXI-P/I	Pediatric / infant	

Nellcor-SpO₂ Extension cable

Accessories	PN
Extension cable	15-100-0016

BLT SpO₂ Sensor

Туре	Patient category	PN
	Adult	15-100-0010
Reusable	Pediatric	15-100-0011
	Neonatal	15-100-0012

Masimo SpO₂ Sensor

Туре	Model / PN	Patient category
Daugahla	DCI / 2501	Adult finger
Reusable	DCIP / 2502	Pediatric finger
Disposable	Neo / 2514	Infant foot/hand

Masimo SpO₂ Extension cable

Accessories	Model / PN
Extension cable	M-LNC1 / 2523

3. Temp

Temp probe

Туре	Applied site	PN
Reusable	Surface	15-100-0027
	Coelom	15-100-0028

4. NIBP

Disposable cuffs

Model	Patient category	Limb circumference (cm)	Bladder width (cm)
M1866A	Neonatal	3.1-5.7	2.5
M1868A		4.3-8.0	3.2
M1870A		5.8-10.9	4.3
M1872A		7.1-13.1	5.1

Disposable cuffs (SunTech)

Patient category	Limb circumference (cm)	PN
	7-13	15-100-0173
Neonatal	6-11	15-100-0174
	3-5.5	15-100-0175

Patient category	Limb circumference (cm)	Bladder width (cm)	PN
Large adult			15-100-0021
Adult	25-35	14.4	15-100-0019
Small adult	20-28	11	15-100-0023
Pediatric	13-20	8	15-100-0022
Infant	10-18	5	15-100-0024
Neonatal	6-11		15-100-0025
Adult (Thigh)			15-100-0020

Reusable cuffs

5. IBP

Accessories	Material	PN
IBP kits	DPT-248 pressure transducer	15-100-0031
(UTAH)	Extension cable	15-100-0029
IBP kits (BD)	TNF-R pressure transducer	15-100-0034
	Pressure tubing	15-100-0033
	Extension cable	15-100-0032

6. CO₂

Sidestream CO₂ (CPT)

Accessories	PN
CO ₂ sampling tube	15-100-0035
CO ₂ dehydration flash	15-100-0036
CO ₂ 3-way stopcock	15-100-0037

Mainstream CO₂ (C5)

Accessories	PN
CAPNOSTAT5 CO ₂ sensor	16-100-0015
Airway adapter (adult)	15-100-0042
Airway adapter(neonatal)	15-100-0043

Microstream CO₂ (LoFlo)

Accessories	PN
LoFlo CO ₂ sensor	16-100-0016
Airway adapter (adult)	15-100-0045
CO ₂ nasal cannula (adult)	15-100-0044
CO ₂ /O ₂ nasal cannule (adult)	15-100-0046

CO ₂ nasal cannule (pediatric)	15-100-0048
CO ₂ nasal cannule (infant)	15-100-0049

7. AG

Mainstream AG (IRMA)

Accessories	PN
IRMA CO ₂ sensor	16-100-0017
IRMA AX sensor	16-100-0019
Airway adapter (adult/pediatric)	15-100-0039
Airway adapter (infant)	15-100-0040
Extension cable	15-024-0001

Sidestream AG (ISA)

Accessories	PN
ISA OR+ module	16-100-0037
ISA module Modura holder	15-100-0090
ISA module Clamp Adapter	15-100-0091
Nomoline Sampling line	15-100-0089

8. ICG

Accessories	Model	PN
ICG cable	BoiZ DX 5551	15-100-0050
ICG lead	BoiZ DX 5562	15-100-0051
ICG electrode	BZ-1550-50	15-100-0052

Appendix A Specifications

A.1 Classifications

Refer to chapter 2.3.

A.2 Specifications

Environment

Ambient Temperature	Operating temperature: $5^{\circ}C \rightarrow 40^{\circ}C$ Transportation and storage temperature: $-20^{\circ}C$ $\rightarrow +55^{\circ}C$
Relative humidity	Working $\leq 85\%$ Transportation and storage $\leq 93\%$
Atmospheric pressure	Working 700 hPa~1060 hPa Transportation and storage 500 hPa~1060 hPa

Size and Weight

Size	318mm×264mm×152mm
Weight	4.5kg

Power supply

Power Voltage	AC 100V-240V
Frequency	50Hz/60Hz
Input current	0.6A -0.3A
Fuse	T 2A/250V, integrated in the power module
Earth leakage current	<0.3 mA
Safety class	Category I

Display

LCD	
Size	M9000, M9000A: 12.1" M8000, M8000A: 10.4"
Туре	Color TFT-LCD

Resolution	800×600 pixels or higher	
Indicators		
Physiological alarm LED	1 (Yellow/Red)	Only for
Technical alarm LED	1 (Blue)	M8000A&M9000A
Alarm LED	1 (Yellow/Red) (Only for M8000&M9000)	
AC Power LED	1 (Green/Orange)	
Battery Charge LED	1 (Yellow)	

Interface

Power	1 AC power inlet
Wired network	1, standard RJ45 socket
Wireless network	1, 433MHz, 10mW (optional)
Equipotential grounding	1
terminal	
Defibrillator synchronization	1 RJ11 socket
output	

Signal output

Defibrillator synchronization signal output		
Output impedance	50Ω	
Delay	\leq 35 ms (from R wave crest to pulse raise)	
Amplitude	High level : 3.5 V to 5 V, the maximal output current	
	1mA.	
	Low level: < 0.5 V, the maximal input current 5mA.	
Pulse width	100ms ±10%	
Rise and drop time	< 1ms	

Battery

Tumo	Rechargeable Lithium ion battery	
Туре	11.1V/4.0AH	
Model	Li1104C	
Charge time	\leq 6 hours (2 batteries for 12 hours)	
Operating time under the	\geq 240 minutes (2 batteries for 480 minutes)	

normal use and full charge	New and fully charged battery at 25 °C ambient temperature and NIBP work on AUTO mode for 20 minutes interval.
Operating time after the first alarm if low battery	≥ 10 minutes

Battery

Туре	Rechargeable Lead acid cell, 12V/2.0AH		Rechargeable Lead acid cell, 12V/2.0AH	
Model	FP1223C			
Charge time	≤ 10 hours (2 batteries for 20 hours)			
Operating time under the normal use and full charge	≥60 minutes (2 batteries for 120 minutes) New and fully charged battery at 25 °C ambient temperature and NIBP work on AUTO mode for 20 minutes interval.			
Operating time after the first alarm if low battery	\geq 5 minutes			

ECG

Lead Mode	1. 5-leads ECG input		
	3 12-leads ECG input (option)		
Lead selection	1. I, II, III, aVR, aVL, aVF, V- 2. I, II, III 3. I, II, III, aVR, aVL, aVF, V1~V6 (option)		
Gain	AUTO, 0.25x, 0.5x, 1.0x, 2.0x, 4.0x		
Input impedance	≥5.0 MΩ		
CMRR	$ \begin{array}{c} MON \geq 105 dB \\ OPS \geq 105 dB \end{array} $		
Frequency response	MON 0.5~40Hz OPS 1~25Hz		
Electrode offset potential	±500mV d.c.		
Leakage Current	<10 uA		
ECG signal range	±6.0 mV		
Baseline recovery	<5s after Defibrillation. (MON or OPS mode)		
Pacemaker pulses	No rejection of pulses with amplitudes of $\pm 2mV \sim \pm 700 \text{ mV}$ and durations of $0.5 \sim 2.0 \text{ ms}$.		
Insulation	Breakdown Voltage 4000VAC 50/60Hz		

Indication of electrode separation	Every electrode (exclusive of RL)	
Sweep speed	12.5mm/s, 25mm/s, 50mm/s	

HR

Measurement range	10~350 bpm	
Refreshing time	Per 4 pulses	
Resolution	1 bpm	
Accuracy	$\pm 1\%$ or ± 1 bpm, whichever is greater	
Sensitivity	≥0.2mVpp	
Alarm range	0~350 bpm, continuously adjustable between upper limit and lower limit	
Alarm indication	Sound and light alarming	
Time to Alarm for Tachycardia	Average 4s	
Tall T-Wave Rejection Capability	0-1 mV T-Wave amplitude	
Response Time of Heart Rate Meter to Change in Heart Rate	HR change from 80 to 120 bpm: Range: 6 to 10s HR change from 80 to 40 bpm: Range: 6 to 10s	

ST segment

Measurement range	-2.0mV~2.0mV
Accuracy	-0.8mV~0.8mV : ±0.02mV or ±10% whichever is greater Over ±0.8mV: unspecified
Resolution	0.01mV
Refreshing time	10s
Alarm range	-2.00mV \sim 2.00mV, continuously adjustable between upper limit and lower limit.

NIBP

Way of measurement	Automatic oscillometry		
Range of measurement	Adult	SYS	30~270 mmHg
		DIA	10~220 mmHg
		MEAN	20~235 mmHg

		SYS	30~235 mmHg	
	Child	DIA	10~220 mmHg	
		MEAN	20~225 mmHg	
		SYS	30~135 mmHg	
	Neonate	DIA	10~110 mmHg	
		MEAN	20~125 mmHg	
Range of HYPER measurement SYS DIA MEAN	(Only for a $40 \sim 300 \text{m}$ $10 \sim 250 \text{ m}$ $20 \sim 270 \text{ m}$	(Only for adult) $40 \sim 300$ mmHg $10 \sim 250$ mmHg $20 \sim 270$ mmHg		
Cuff pressure range	0~280 mm	$0 \sim 280 \text{ mmHg} (0 \sim 300 \text{ mmHg at HYPER mode})$		
Resolution	1 mmHg	1 mmHg		
Pressure Accuracy Static Clinical	±2% or ±3 ±5 mmHg 8 mmHg s	±2% or ±3 mmHg, whichever is greater ±5 mmHg average error 8 mmHg standard deviation		
Unit	mmHg, kP	mmHg, kPa		
Pulse rate range	40 ~ 240 b	40 ~ 240 bpm		
Inflation time for cuff	Less than 4	Less than 40s. (standard adult cuff)		
Total cycle time	20 to 45s motion art	20 to 45s typical (dependent on heart rate and motion artifact)		
Intervals for AUTO measurement time	1,2,3,4,5,1 2,4,8 hours	1,2,3,4,5,10,15,20,30,45,60,90 minutes 2,4,8 hours		
Initial inflation pressure	Adult : 170 Pediatric : Neonatal : (The value	Adult : 170 mmHg Pediatric : 130 mmHg Neonatal : 100 mmHg (The value can be adjusted manually)		
Overpressure Protection	Hardware	Hardware and software double protections		
Adult	297±3 mm	297±3 mmHg		
Child	252±3 mm	252±3 mmHg		
Neonatai	14/±3 mm	нд		
Range of alarm	SYS	0~300 mmHg between uppe	, continuously adjustable r limit and lower limit	
	DIA	0~300 mmHg between uppe	, continuously adjustable r limit and lower limit	

	MEAN	0~300 mmHg, continuously adjustable between upper limit and lower limit	
Alarm indication	Sound and light alarming		
Measurement Mode	Adult	Manual, Auto and STAT	
	Child	Manual, Auto and STAT	
	Neonatal	Manual, Auto	
	HYPER	Manual, Auto and STAT	

SunTech NIBP (option)

Way of measurement	Oscillometry		
Range of measurement	Adult	SYS	40~260 mmHg
		DIA	20~200 mmHg
		MEAN	26~220 mmHg
	Child	SYS	40~160 mmHg
		DIA	20~120 mmHg
		MEAN	26~133 mmHg
		SYS	40~130 mmHg
Range of measurement	Neonate	DIA	20~100 mmHg
		MEAN	26~110mmHg
Pressure Accuracy			
Static	±3 mmHg		
Clinical	±5 mmHg average error		
	8 mmHg standard deviation		
Unit	mmHg, kPa		
Pulse rate range	30 ~ 220 bpm		
Pulse veracity	2bpm 或 3%, whichever is greater		
Inflation time for cuff	<75s		
Measurement protection	Adult: <180s		
	Child: <180s		
	Neonate: <90s		
	Adult: 120~280mmHg, default 160mmHg		
Initial inflation pressure	Child: 80~170mmHg, default 120mmHg		
	Neonate: 60~140mmHg, default 90mmHg		
Intervals for AUTO	1,2,3,4,5,10,15, 30, 60,90 minutes		
measurement time	2,4,6 hours		

Overpressure Protection	Hardware and software double protections	
Adult	<300 mmHg	
Child <3 Neonatal <1	<300 mmF <150 mmF	Ig Ig
Alarm indication	Sound and light alarming	
Measurement Mode	Adult	Manual, Auto and STAT
	Child	Manual, Auto and STAT
	Neonatal	Manual, Auto

SpO2

BLT-SpO ₂	
Measurement Range	0~100%
Resolution	1%
Accuracy	At 70~100%, ±2% At 0~69%, unspecified
Data update period	<13s
Alarm range	50~100%, continuously adjustable between upper limit and lower limit.
PR	
Measurement Range	25~250 bpm
Resolution	1 bpm
Accuracy	$\pm 1\%$ or ± 1 bpm, whichever is greater
Data update period	<13s
Alarm range	0~300 bpm, continuously adjustable between upper limit and lower limit.

Nellcor-SpO ₂ (option)	
Measurement Range	0~100%
Resolution	1%
Accuracy	At 70~100%, ±2 digits (Adult)
	At 70~100%, ±3 digits (Neonate)
	At 70~100%, ±2 digits (Low Perfusion)
	At 0~69%, unspecified
Perfusion Range	$0.03\% \sim 20\%$
Data update period	Average 7s

Alarm range	50~100%, continuously adjustable between upper limit and lower limit.
PR	
Measurement Range	20~250 bpm
Resolution	1 bpm
Accuracy	±3 digits
Data update period	Average 7s
Alarm range	0~300 bpm, continuously adjustable between upper limit and lower limit.

Masimo-SpO ₂ (option)	
SpO2	
Measurement Range	0~100%
Resolution	1%
Accuracy	At 70~100%: ±2% (adult/pediatric, non-motion conditions)At 70 ~ 100%: ±3% (neonate, non-motion conditions)At 70 ~100%: ±3% (motion conditions)At 70 ~100%: ±3% (motion conditions)At 0~69%, unspecified
Average time	2-4s, 4-6s, 8s, 10s, 12s, 14s, 16s
Alarm range	0~100%, continuously adjustable between upper limit and lower limit.
PR	
Measurement Range	25~240 bpm
Resolution	1 bpm
Accuracy	 ±3 bpm (non-motion conditions) ±5 bpm (motion conditions)
Alarm range	0~300 bpm, continuously adjustable between upper limit and lower limit.
PI	
Measurement Range	0.02 ~ 20.0% (for disposable sensor) 0.05~ 20.0% (for reusable sensor)
Resolution	0.1%
PVI	· ·
Measurement Range	0~100%

	-
Resolution	1%

TEMP

Measurement Range	0.0~50.0°C
Accuracy	±0.1°C
Resolution	0.1°C
Unit	Celsius (°C), Fahrenheit (°F)
Refreshing time	1s
Self check	Every 10 minutes
Accuracy	At 45.1~50.0°C, ± 0.2 °C (exclusive of probe) At 25.0~45.0°C, ± 0.1 °C (exclusive of probe) At 0.0~24.9°C, ± 0.2 °C (exclusive of probe)
Connecting cable	Compatible with YSI-400
Alarm range	$0.0{\sim}50.0~{}^\circ\!\mathrm{C}$, continuously adjustable between upper limit and lower limit
Alarm indication	Sound and light alarming

RESP

Method	Impedance variation between RA-LL (R-F)
Measuring impedance range	0.2 ~3 Ω
Excitation frequency	64.8 kHz
Excitation current	≤300 µ A at 64.8 kHz
Base line impedance range	500~4000 Ω (50~120 kHz exciting frequency)
Measurement Range	0~150 rpm
Resolution	1 rpm
Accuracy	±2 rpm
Gain	x1, x2, x4
Sweep speed	6.25mm/s, 12.5mm/s, 25mm/s
Delay of Apnea Alarm	Off, 10s, 20s, 40s, 60s
Alarm range	0~150 rpm, continuously adjustable between upper limit and lower limit
Alarm indication	Sound and light indication

Measurement Range	-50 ~ +300 mmHg		
Resolution	1 mmHg		
Unit	mmHg, kPa		
Accuracy Static Dynamic	 ± 2mmHg or 2% of the reading, whichever is greater (exclusive of transducer) ± 4mmHg or 4% of the reading, whichever is greater (inclusion of transducer) ± 4mmHg or 4% of the reading, whichever is greater 		
Sensitivity of transducer	5uV/V/m	5uV/V/mmHg, 2%	
Impedance of transducer	300~3000) Ω	
Bandwidth	d.c. ~ 15H	łz	
Transducer sites	Arterial Pressure (ART) Pulmonary Artery Pressure (PA) Left Atrium Pressure (LAP) Right Atrium Pressure (RAP) Central Venous Pressure (CVP) Intracranial Pressure (ICP)ART PA CVP LAP0~200mmHg 0~300 mmHg -10~20 mmHg		
Selection of measurement range	RAP		-50~300 mmHg AUTO
	(Among them, the AUTO switches automatically at an interval of 10 mmHg so as to ensure the waveform is at the state most suitable for observation)		
Alarm range	SYS	SYS -50~300 mmHg, continuously adjustate between upper limit and lower limit	
	DIA -50~300 m between u		mHg, continuously adjustable pper limit and lower limit
	MEAN	-50~300 m between u	mHg, continuously adjustable pper limit and lower limit
Alarm indication	Sound and light indication		

IBP

CO2 (Sidestream, CPT)

Measure method	Infrared spectrum
Measure mode	Sidestream

Measurement Range	0.0~13.1% (0~99.6 mmHg)
Resolution	1 mmHg
Unit	%, mmHg, kPa
Accuracy	At <5 % CO ₂ , $\pm 0.3\%$ (± 2.0 mmHg) At ≥ 5 % CO ₂ , $\leq \pm 10$ % of reading
Range of respiration rate measurement	3~150 rpm
Warm up time	Reach 97% of full specifications within 45s and full specifications within 10 minutes.
Rise time	About 100ms when using 1.5m sampling tube and adult dehydration flask with 120ml/min flow rate.
CO ₂ Response Time	<4s when using 1.5m sampling tube and adult dehydration flask with 120ml/min flow rate.
Sample Flow Rate	User adjustable, 50ml/min, 100ml/min, 150ml/min, 200ml/min, 250ml/min
Calibration	Offset calibration: auto, manual Gain calibration
Alarm range	0.0~13.1 % (0~99.6mmHg), continuously adjustable between upper limit and lower limit
Alarm indication	Sound and light indication

CO2 (Microstream, LoFlo)

Measure method	Infrared spectrum		
Measure mode	Microstream		
Warm up time	Capnogram displayed in less than 20 s, At an		
	ambient temperature of 25°C, full specifications		
	within 2 minutes.		
CO ₂ Measurement Range	0~19.7%(0~150 mmHg)		
CO ₂ Resolution	1mmHg		
CO ₂ Stability	Short-Term Drift: Drift over four hours≤0.8mmHg.		
	Long-Term Drift: Accuracy specification will be		
	maintained over a 120 hours period.		
unit	%, mmHg, kPa		
CO ₂ Accuracy	$0 \sim 40 \text{ mmHg}, \pm 2 \text{ mmHg}$		
(at 760 mmHg, ambient	$41 \sim 70$ mmHg, $\pm 5\%$ of reading		
temperature of 25°C)	$71 \sim 100 \text{ mmHg}, \pm 8\% \text{ of reading}$		
	$101 \sim 150 \text{ mmHg}, \pm 10\% \text{ of reading}$		
	Above 80 breath per minute \pm 12% of reading		
	Gas temperature at 25°C.		
CO ₂ response time	<3s (includes transport time and rise time)		

Respiration Rate Range	2~150 rpm
Respiration Rate Accuracy	±1 rpm
Sample Flow Rate	50 ml/min ±10 ml/min
Alarm range	0.0~13.1 % (0~99.6mmHg), continuously adjustable between upper limit and lower limit
Alarm indication	Sound and light indication

CO2 (Mainstream, CAPNOSTAT5)

Measure method	Infrared spectrum	
Measure mode	Mainstream	
Warm up time	Capnogram displayed in less than 15 s, At an ambient temperature of 25°C, full specifications within 2 minutes.	
CO ₂ Measurement Range	0~19.7%(0~150 mmHg)	
CO ₂ Resolution	1mmHg	
CO ₂ Accuracy	0 ~ 40 mmHg, ±2 mmHg 41 ~ 70 mmHg, ±5% of reading 71 ~100 mmHg, ±8% of reading 101 ~ 150 mmHg, ±10% of reading Temperature at 35°C.	
CO ₂ Stability	Short-Term Drift: Drift over four hours≤0.8 mmHg. Long-Term Drift: Accuracy specification will be maintained over a 120 hours period.	
Rise time	<60ms	
unit	%, mmHg, kPa	
Respiration Rate Range	0~150 rpm	
Respiration Rate Accuracy	±1 rpm	
Alarm range	0.0~13.1 % (0~99.6mmHg), continuously adjustable between upper limit and lower limit	
Alarm indication	Sound and light indication	

CO2 (Mainstream, IRMA)

Measure method	Infrared spectrum		
Measure mode	Mainstream		
Fi and Et values	CO ₂		
Resolution	1%		

Calibration	Zeroing recommended when changing Airway adapter. No span calibration required for the IR bench. Room air calibration of oxygen sensor performed automatically when changing airway adapter (<5s)
Warm-up time	Concentrations are reported and the automatic agent identification is running within 10s. Full accuracy within 10s.
Rise time(at 10 L/min)	\leq 90 ms
Total system response time	< 1 s
CO ₂ Measurement Range	0~15%
CO2 accuracy	$\pm (0.3\%_{ABS} + 4\%_{REL})$

AG (Mainstream, IRMA)

Measure method		Infrared spectrum			
Measure mode		Mainstream			
Fi and Et values		CO ₂ , N ₂ O, agent (HAL, ISO, ENF, SEV, DES)			
Resolution		1%			
Calibration		Zeroing recommended when changing Airway adapter. No span calibration required for the IR bench. Room air calibration of oxygen sensor performed automatically when changing airway adapter (<5s)			
Warm-up time		Concentrations are reported and the automatic agent identification is running within 10s. Full accuracy within 20s for IRMA AX+.			
Rise time (at 10 L/min)		$\begin{array}{l} CO_2 \leqslant 90 ms \\ N_2O \ \leqslant \ 300 \ ms \\ HAL, ISO, ENF, SEV, DES \ \leqslant \ 300 \ ms \end{array}$			
Total system respo	em response time < 1 s				
Primary agent threshold		0.15%. When an agent is identified, concentrations will reported even below 0.15% as long as apnea is not detected			
Second agent three	shold	0.2% + 10% of total agent concentration			
Agent identification time		< 20s. (Typically < 10s)			
Measurement range and accurac		acy of gas:			
Gas	Range (%)	Accuracy ¹⁾			

CO ₂	0-15	$\pm (0.3\%_{ABS} + 4\%_{REL})$
N ₂ O	0-100	$\pm (2\%_{ABS}+5\%_{REL})$
HAL, ISO, ENF	0-8	$\pm (0.2\%_{ABS} + 10\%_{REL})$
SEV	0-10	$\pm (0.2\%_{ABS} + 10\%_{REL})$
DES	0-22	$\pm (0.2\%_{ABS} + 10\%_{REL})$
Respiration rate range	0~150 rpm	
Respiration rate accuracy	±1 rpm	
Alarm indication	Sound and light indication	

Note1: The accuracy specification is valid for the operating temperature and humidity conditions specified, except for interference specified in the table "Interfering gas and vapor effects" below. The accuracy specification for IRMA AX+ is not valid if more than two agents are present in the gas mixture.

Interfering gas and vapor effects

If the concentration of the various gases is below the following level, there will be negligible interference for measurement under the specification above.

Gas or vapor	Gas level	Effect (CO ₂)	Effect (Agents)	Effect (N ₂ O)
N ₂ O	60%		— —	— —
HAL	4%		—	
ENF, ISO, SEV	5%	+8% of reading ¹)		
DES	15%	+12% of reading ¹)		
Xe (Xenon)	80%	-10% of reading ¹⁾		
He (Helium)	50%	-6% of reading ¹⁾		
Metered dose inhaler Propellants	Not for use with metered dose inhaler propellants			
C ₂ H ₅ OH (Ethanol)	0.3%		—	
C ₃ H ₇ OH (Isopropanol)	0.5%	—	—	
CH ₃ COCH ₃ (Acetone)	1%	—		—
CH ₄ (Methane)	3%	—	—	—
CO (Carbon monoxide)	1%			
NO (Nitrogen monoxide)	0.02%			
O ₂	100%	—	—	—

♦ IRMA CO₂ sensor:
Gas or vapor	Gas level	Effect (CO ₂)	Effect (Agents)	Effect
N ₂ O	60%			—
HAL	4%		—	
ENF, ISO, SEV	5%		—	
DES	15%			
Xe (Xenon)	80%	-10% of reading ¹⁾		
He (Helium)	50%	-6% of reading ¹⁾	—	
Metered dose inhaler	Not for use with m	netered dose inhaler p	ropellants	
Propellants				
C ₂ H ₅ OH (Ethanol)	0.3%	—		—
C ₃ H ₇ OH (Isopropanol)	0.5%	—	—	—
CH ₃ COCH ₃ (Acetone)	1%		—	
CH ₄ (Methane)	3%		—	
CO (Carbon monoxide)	1%		—	
NO (Nitrogen	0.02%	—	—	—
monoxide)				
O ₂	100%			

♦ IRMA AX+ sensor:

Note 1: Interference at indicated gas level. For example, 50 % Helium typically decreases the CO2 readings by 6%. This means that if measuring on a mixture containing 5.0 % CO2 and 50 % Helium, the actual measured CO2 concentration will typically be (1-0.06) * 5.0 % = 4.7 % CO2.

Measurement mode	Sidestream
Sampling flow rate	50ml/min ± 10ml/min
Measurement parameters	CO ₂ , N ₂ O, O ₂ , agent (ISO, ENF, SEV, HAL, DES)
Compensations	Automatic compensation for pressure, temperature and broadening effects on CO ₂ .
Calibration	No span calibration is required for the IR bench. An automatic zero reference calibration is performed at startup and then every 24 hours.
Warm-up time	< 20 s (Concentrations reported, automatic agent identification enabled and full accuracy)
Typical rise time at 50 ml/min sample flow	$\begin{array}{l} \text{CO}_2 \leq 200 \text{ ms} \ (\leq 250 \text{ ms for ISA OR} + / \text{AX} +) \\ \text{N}_2\text{O} \leq 350 \text{ ms} \\ \text{O}_2 \leq 450 \text{ ms} \\ \text{ISO, ENF, SEV, HAL, DES} \leq 350 \text{ ms} \end{array}$

AG (Sidestream, ISA)

Total system response time		< 3 s (with 2 m sampling line)	
Primary agent threshold		0.15%. When an agent is identified, concentrations will be reported even below 0.15%.	
Secondary agent threshold		0.2 % + 10% of total agent concentration	
Agent identification time		< 20 s (typically < 10 s)	
Measurement range and accura		acy of gas:	
Gas	Range		Accuracy ¹⁾
CO ₂	0-15%		\pm (0.3 kPa + 4% of reading)
N ₂ O	0-100%		\pm (2 kPa + 5% of reading)
HAL, ENF, ISO	0-8%		\pm (0.2 kPa + 10% of reading)
SEV	0-10%		\pm (0.2 kPa + 10% of reading)
DES	0-22%		\pm (0.2 kPa + 10% of reading)
O ₂	0-100%		\pm (2 kPa + 2% of reading)

Note 1: The accuracy specification is valid for the operating temperature and humidity conditions specified, except for interference specified in the table "Interfering gas and vapor effects" below. The accuracy specification is not valid if more than two agents are present in the gas mixture. If more than two agents are present, an alarm will be set.

Interfering gas and vapor effects

If the concentration of the various gases is below the following level, there will be negligible interference for measurement under the specification above.

♦ ISA AX+ sensor:

Cas or vapor	Casloval	Effort (CO.)	Effect	Effect
Gas of vapor	Gas level		(Agents)	(N ₂ O)
N ₂ O	60%		—	—
HAL	4%			
ENF, ISO, SEV	5%			
DES	15%			
Xe (Xenon)	80%	-10% of reading ¹⁾		
He (Helium)	50%	-6% of reading ¹⁾		
Metered dose inhaler	Not for use with metered dose inhaler propellants			
Propellants				
C ₂ H ₅ OH (Ethanol)	0.3%			
C ₃ H ₇ OH (Isopropanol)	0.5%			
CH ₃ COCH ₃ (Acetone)	1%			—
CH ₄ (Methane)	3%		—	—
CO (Carbon monoxide)	1%			
NO (Nitrogen	0.02%			

monoxide)			
O ₂	100%	 	

Note 1: Interference at indicated gas level. For example, 50 % Helium typically decreases the CO2 readings by 6%. This means that if measuring on a mixture containing 5.0 % CO2 and 50 % Helium, the actual measured CO2 concentration will typically be (1-0.06) * 5.0 % = 4.7 % CO2.

ICG

Measuring Method	Measurement of thoracic electrical bioimpedance
	HR: 40~250 bpm
	SV: 5~250 mL
Measuring Range	SI: 5~125 mL/m ²
	C.O.: 1.4~15 L/min
	TFC: $15 \sim 143/k$ Ω
	HR: ±2 rpm
Measuring accuracy	SV: unspecified
	C.O.: unspecified
	C.I.: 0.0~15.0 L/min/m ² ,
Alarm range	TFC: 10~150 /k Ω
	continuously adjustable between high limit and low
	limit
Alarm indication	Sound and light indication

Recorder

Method	Thermal dot array
Paper width	50 mm
Record width	40 mm
Paper Speed	12.5 mm/s ,25 mm/s ,50 mm/s
Traces	Maximum 3 tracks

Alarm

Level	Low, medium and high
Indication	Auditory and visual
Setup	Default and custom
Silence	All alarms can be silenced
Volume	45~85 dB measured at 1 meter

Appendix B Default System Setup

There are three options of default system setup: ADULT, CHILD, NEONATAL. The followings are the detail:

B.1 System

1. Standard Configuration

1) Trend Graph Configuration

Region	Parameter	
Region 1	HR	
Region 2	SpO ₂	
Region 3	PR	
Region 4	NIBP	
Region 5	Resp	
Region 6	T1+T2	

2) Trend Table Configuration

Page 1

Region	Parameter
Region 1	HR
Region 2	SpO ₂
Region 3	PR
Region 4	NIBP(S/D)
Region 5	NIBP(M)
Region 6	Resp

Page 2

Region	Parameter
Region 1	HR
Region 2	T1
Region 3	T2

2. Standard Configuration + dual IBP

1) Trend Graph Configuration

Page 1

Region	Parameter
Region 1	HR
Region 2	SpO ₂
Region 3	P1
Region 4	P2
Region 5	Resp

Page 2

Region	Parameter
Region 1	PR
Region 2	NIBP
Region 3	T1+T2
Region 4	NIBP

2) Trend Table Configuration

Page 1

Region	Parameter
Region 1	HR
Region 2	SpO ₂
Region 3	P1
Region 4	P2
Region 5	Resp

Page 2

Region	Parameter
Region 1	PR
Region 2	NIBP(S/D)
Region 3	NIBP(M)
Region 4	T1
Region 5	T2

3. Standard Configuration + dual IBP + EtCO₂

1) Trend Graph Configuration

Page 1

Region	Parameter
Region 1	HR
Region 2	SpO ₂
Region 3	P1
Region 4	P2
Region 5	CO ₂

Page 2

Region	Parameter
Region 1	PR
Region 2	NIBP
Region 3	Resp
Region 4	T1+T2

2) Trend Table Configuration

Page 1

Region	Parameter
Region 1	HR
Region 2	SpO ₂
Region 3	P1(S/D)
Region 4	P2(M)
Region 5	CO ₂

Page 2

Region	Parameter
Region 1	PR
Region 2	NIBP(S/D)
Region 3	NIBP(M)
Region 4	Resp
Region 5	T1
Region 6	T2

4. Standard Configuration + dual IBP + EtCO₂+GAS

1) Trend Graph Configuration

Page 1

Region	Parameter
Region 1	HR
Region 2	SpO ₂
Region 3	P1
Region 4	P2
Region 5	CO ₂

Page 2

Region	Parameter
Region 1	PR
Region 2	NIBP
Region 3	Resp
Region 4	O ₂ +N ₂ O
Region 5	AA
Region 6	T1+T2

2) Trend Table Configuration

Page 1

Region	Parameter
Region 1	HR
Region 2	SpO ₂
Region 3	P1(S/D)
Region 4	P2(M)
Region 5	CO ₂

Page 2

Region	Parameter
Region 1	PR
Region 2	NIBP(S/D)
Region 3	NIBP(M)
Region 4	Resp
Region 5	T1
Region 6	T2

Page 3

Region	Parameter
Region 1	CO ₂
Region 2	N ₂ O
Region 3	AA
Region 4	O ₂

B.2 Alarm Limit

1. Setup of parameters alarm limit for adult

Parameter	Low limit	High limit
HR (bpm)	50	120
SpO ₂ (%)	90	100
PR (bpm)	50	120
RR (rpm)	8	30
T1 (°C)	36.0	39.0
T2 (°C)	36.0	39.0
NIBP SYS(mmHg)	90	160
NIBP DIA (mmHg)	50	90
NIBP MEAN (mmHg)	60	110
IBP1 SYS (mmHg)	90	160
IBP1 DIA (mmHg)	50	90
IBP1 MEAN (mmHg)	60	110
IBP2 SYS (mmHg)	6	14
IBP2 DIA (mmHg)	-4	6
IBP2 MEAN (mmHg)	0	10
EtCO ₂ (mmHg)	20	50
FiCO ₂ (mmHg)	0	20
EtAA (%)	0.0	3.0
FiAA (%)	0.0	5.0
EtN ₂ 0 (%)	0	82

FiN ₂ 0 (%)	0	82
EtO ₂ (%)	10	100
FiO ₂ (%)	18	100
C.I.(L/min/m2)	1.5	5.0
$TFC(/k\Omega)$	15.0	60.0

2. Setup of parameters alarm limit for child

Parameter	Low limit	High limit
HR (bpm)	75	160
SpO ₂ (%)	90	100
PR (bpm)	75	160
RR (rpm)	8	30
T1 (°C)	36.0	39.0
T2 (°C)	36.0	39.0
NIBP SYS (mmHg)	70	120
NIBP DIA (mmHg)	40	70
NIBP MEAN (mmHg)	50	90
IBP1 SYS (mmHg)	70	120
IBP1 DIA (mmHg)	40	70
IBP1 MEAN (mmHg)	50	90
IBP2 SYS (mmHg)	2	10
IBP2 DIA (mmHg)	-4	2
IBP2 MEAN (mmHg)	0	4
EtCO ₂ (mmHg)	20	50
FiCO ₂ (mmHg)	0	20
EtAA (%)	0.0	3.0
FiAA (%)	0.0	5.0
EtN ₂ 0 (%)	0	82
FiN ₂ 0 (%)	0	82
EtO ₂ (%)	10	100
FiO ₂ (%)	18	100
C.I.(L/min/m2)	1.5	5.0
TFC(/kΩ)	15.0	60.0

3.	Setup of	f parameters	alarm	limit	for neonate
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Parameter	Low limit	High limit
HR (bpm)	90	200
SpO ₂ (%)	85	95
PR (bpm)	90	200
RR (rpm)	30	100
T1 (°C)	36.0	39.0
T2 (°C)	36.0	39.0
NIBP SYS(mmHg)	40	90
NIBP DIA (mmHg)	20	60
NIBP MEAN (mmHg)	25	70
IBP1 SYS (mmHg)	40	90
IBP1 DIA (mmHg)	20	60
IBP1 MEAN (mmHg)	25	70
IBP2 SYS (mmHg)	2	10
IBP2 DIA (mmHg)	-4	2
IBP2 MEAN (mmHg)	0	4
EtCO ₂ (mmHg)	20	45
FiCO ₂ (mmHg)	0	20
EtAA (%)	0.0	3.0
FiAA (%)	0.0	5.0
EtN ₂ 0 (%)	0	82
FiN ₂ 0 (%)	0	82
EtO ₂ (%)	10	100
FiO ₂ (%)	18	100
C.I.(L/min/m2)	1.5	5.0
TFC(/kΩ)	15.0	60.0

Appendix C Guidance and Manufacture's Declaration of EMC

Guidance and manufacture's declaration – electromagnetic emissionsfor all EQUIPMENT and SYSTEMS

Guidance and manufacture's declaration – electromagnetic emission				
The Monitor is intended for use in the electromagnetic environment specified below. The customer of the user of the Multi-parameter Monitor should assure that it is used in such and environment				
Emission test Compliance Electromagnetic environment – guidance				
RF emissions CISPR 11	Group 1	The Monitor uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.		
RF emission CISPR 11	Class A	The Monitor is suitable for use in all establishments other than domestic and those directly connected to the public		
Harmonic emissions IEC 61000-3-2	Class A	low-voltage power supply network that supplies buildings used for domestic purposes.		
Voltage fluctuations/ flicker emissions IEC 61000-3-3	Complies			

Guidance and manufacture's declaration – electromagnetic immunity – for all EQUIPMENT and SYSTEMS

Guidance and manufacture's declaration – electromagnetic immunity					
The Monitor is intended for use in the electromagnetic environment specified below. The customer or the user of Low Frequency Therapeutic Device should assure that it is used in such an environment.					
Immunity test	IEC 60601 test level Compliance level Electromagnetic environn guidance				
Electrostatic discharge (ESD) IEC 61000-4-2	±6 kV contact ±8 kV air	±6 kV contact ±8 kV air	Floors should be wood, concrete or ceramic tile. If floor are covered with synthetic material, the relative humidity should be at least 30%.		
Electrical fast transient/burst IEC 61000-4-4	±2 kV for power supply lines ±1 kV for input/output lines	±2 k V for power supply lines ±1 kV for input/output lines	Mains power quality should be that of a typical commercial or hospital environment.		
Surge IEC 61000-4-5	±1 kV differential mode ±2 kV common mode	±1 kV differential mode ±2 kV common mode	Mains power quality should be that of a typical commercial or hospital environment.		
Voltage dips, short interruptions and voltage variations on power supply input lines IEC 61000-4-11	<5% UT (>95% dip in UT) for 0.5 cycle 40% UT (60% dip in UT) for 5 cycles 70% UT (30% dip in UT) for 25 cycles <5% UT (>95% dip in UT) for 5 sec	<5% UT (>95% dip in UT) for 0.5 cycle 40% UT (60% dip in UT) for 5 cycles 70% UT (30% dip in UT) for 25 cycles <5% UT (>95% dip in UT) for 5 sec	Mains power quality should be that of a typical commercial or hospital environment.		
Power frequency (50Hz) magnetic field IEC 61000-4-8	3A/m	3A/m	Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.		

Guidance and manufacture's declaration – electromagnetic immunity – for EQUIPMENT and SYSTEMS that are not LIFE-SUPPORTING

Guidance and manufacture's declaration – electromagnetic immunity				
The Monitor is intended for use in the electromagnetic environment specified below. The customer or the user of Low				
Immunity test IEC 60601 test level Compliance level Electromagnetic environment - guidance				
Conducted RF IEC 61000-4-6 Radiated RF IEC 61000-4-3	3 Vrms 150 kHz to 80 MHz 3 V/m 80 MHz to 2.5 GHz	1 Vrms 3 V/m	Portable and mobile RF communications equipment should be used no closer to any part of the patient monitor, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter. Recommended separation distance $d = \left[\frac{3.5}{V_1}\right]\sqrt{P}$ $d = \left[\frac{3.5}{E_1}\right]\sqrt{P}$ 80 MHz to 800 MHz $d = \left[\frac{7}{E_1}\right]\sqrt{P}$ 800 MHz to 2.5 GHz 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 Interference may occur in the vicinity of equipment marked with the following symbol:	

NOTE 1 At 80 MHz and 800 MHz, the higher frequency range applies.

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

- a 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 Low Frequency Therapeutic Device is used exceeds the applicable RF compliance level above, the Low Frequency Therapeutic Device should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as reorienting or relocating the Low Frequency Therapeutic Device.
- b Over the frequency range 150 kHz to 80 MHz, field strengths should be less than 3 V/m.

Recommended separation distances between portable and mobile RF communications equipment and the EQUIPMENT or SYSTEM – for EQUIPMENT or SYSTEM that are not LIFE-SUPPORTING

Recommended separation distances between

portable and mobile RF communications equipment and the Low Frequency Therapeutic Device The Monitor is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the Low Frequency Therapeutic Device can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the Low Frequency Therapeutic Device as recommended below, according to the maximum output power of the communications equipment.

Rated maximum	Separation distance according to frequency of transmitter (m)				
output power of	150 kHz to 80 MHz	80 MHz to 800 MHz	800 MHz to 2.5 GHz		
transmitter (W)	$d = \left[\frac{3.5}{V_1}\right]\sqrt{P}$	$d = \left[\frac{3.5}{E_1}\right]\sqrt{P}$	$d = \left[\frac{7}{E_1}\right]\sqrt{P}$		
0.01	0.35	0.12	0.23		
0.1	1.1	0.38	0.73		
1	3.5	1.2	2.3		
10	11	3.8	7.3		
100	35	12	23		

For transmitters rated at a maximum output power not listed above, the recommended separation 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 for the higher frequency range applies. NOTE 2 These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people. Product name: Patient monitor

Product type: M8000A

Manufacture for BIOLIGHT MEDITE USA

Address: 533 Broadhollow Road Ste B-24, Melville, New York 11747

Fax: 888-785-8948

Toll-free consultation hot line: 888-82-BLTUSA (888-822-5887)

Post code: 519085

PN: 22-024-0002