Product Information

Product Model: P12/P15/P18/P22
 Product Name: Patient monitor

Manufacturer Name: Guangdong Biolight Meditech Co., Ltd.

After Service Contact Information:

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Revision History

This manual has a revision number. This revision number changes whenever the manual is updated due to software or technical specification change. Contents of this manual are subject to change without prior notice.

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CE mark

C€₀₁₂₃

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Statement

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

Manufacturer's Responsibility

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.
- All the replaced components, assorted accessories and consumables involved in maintenance are original or approved by Biolight.
- The storage condition, operation condition and electrical status of the instrument conform to the product specification. The electrical installation of the relevant room complies with the applicable national and local requirements;
- The instrument is used in accordance with the user's manual.

Signs in this manual:



WARNING: Indicates a potential hazard or unsafe practice that, if not avoided, will result in death or serious injury.



▲ CAUTION: Indicates a potential hazard or unsafe practice that, if not avoided, could result in minor personal injury or product/property damage.



NOTE: Provides application tips or other useful information to ensure that you get the most from your product.

Warranty and maintenance service

The warranty period for the purchased products shall be subject to the sales contract. The consumables means disposable consumables material needed to be replaced after using every time or vulnerable material needed to be replaced regularly. The consumables don't have warranty service.

The warranty period starts from the "installation date" filled in on the attached < Equipment Warranty Card>, which is the only proof to calculate the warranty period. In order to protect your rights, please fill in the warranty card after the installation of the equipment, and give the second warranty card ("Biolight company retention") to the installer or mail it back to the after-service department of Biolight.

Please note that the following conditions will not be covered by the warranty:

- The customer does not fill in or return the equipment warranty card within 30 days after the installation acceptance;
- The equipment series number provided by the customer is not correct (our company confirms the equipment series number is warranty or not).

Within the warranty period, all products can enjoy free after-sales service; however, please note that, even within the warranty period, due to the following reasons, the products need to be repaired, the company will carry out maintenance service, you need to pay maintenance fees and accessories fees:

- Man-made damage;
- Improper use;
- The voltage of the power network exceeds the product's specified range;
- Irresistible natural disasters;
- Replace or use parts, accessories, consumables that are not approved by Biolight or maintained by non-authorized personnel of Biolight;
- Other faults not caused by the product itself.

After the expiration of the warranty, Biolight can continue to provide charged maintenance services. If you do not pay or delay in paying the maintenance fee, Biolight will suspend the maintenance service until you pay for it.

After service

Manufacturer: Guangdong Biolight Meditech Co., Ltd.

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About this manual

This manual contains the instructions necessary to operate the product safely and in accordance with its function and intended use. Observance of this manual is a prerequisite for proper product performance and correct operation and ensures patient and operator safety.

This manual is based on the maximum configuration and therefore some contents may not apply to your product. If you have any question, please contact us.

This manual is an integral part of the product. It should always be kept close to the equipment so that it can be obtained conveniently when needed.

The manual is geared for clinical professionals who are expected to have a working knowledge of medical procedures, practiced and terminology as required for monitoring patients.

All illustrations in this manual serve as examples only. They may not necessarily reflect the setup or data displayed on your product.

Conventions:

- ♦ **Bold Italic** text is used in this manual to quote the referenced chapter or sections.
- ◆ 【】 is used to enclose screen texts.
- → is used to indicate operational procedures.

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Chapter 1 General Introduction

1.1. Intended Use

The P series patient monitors (P12/P15/P18/P22), hereafter called the monitor, are intended to be used for monitoring, displaying, reviewing, storing and alarming of multiple physiological parameters of patients, including ECG, Heart Rate(HR), Respiration Rate(RR), Temperature(TEMP), Pulse Oxygen Saturation(SpO₂), Pulse Rate(PR), Non-invasive Blood Pressure(NIBP), Carbon dioxide(CO₂), Invasive Blood Pressure(IBP), Impedance Cardiograph(ICG), Bispectral Index(BIS), Electroencephalograph(EEG), Anesthetic Gas(AG), Respiratory Mechanics(RM), Regional oxygen saturation(rSO₂), Neuromuscular Transmission(NMT), Total Hemoglobin (SpHb), Carboxyhemoglobin (SpCO) and Methemoglobin (SpMet).

Impedance Cardiograph(ICG) monitoring is only intended for adult patients.

The following functions are not applicable to neonates: Total Hemoglobin (SpHb), Carboxyhemoglobin (SpCO), Methemoglobin (SpMet), Neuromuscular Transmission(NMT) and Bispectral Index(BIS).

The monitors are to be used in medical facilities by clinical professionals or under their guidance.



WARNING:

■ The monitor is intended for use only by clinical professionals or under their guidance. It must only be used by persons who have received adequate training in its use. Anyone unauthorized or untrained must not perform any operations on it.

1.2. Composition

The monitor is composed of main unit, main control software, battery, auxiliary module slots, plug-in modules and corresponding accessories (including CO₂ measuring module and accessories, IBP measuring module, ICG measuring module and accessories, BIS measuring module and accessories, AG measuring module and accessories, rSO₂ measuring module and accessories, NMT measuring module and accessories, EEG cables, RM sensors, ECG cables, NIBP cuffs, SpO₂ sensors and temperature sensor).

1.3. Contraindications

- Do not measure NIBP on patients with sickle-cell disease or any condition where skin damage has occurred or is expected.
- Use clinical judgments to decide whether to perform frequent Auto NIBP measurements on patients with severe thromboembolism disease because of the risk of hematoma in the limb fitted with the cuff.
- Use clinical judgments to decide whether to perform Auto BP measurement on the patients of thrombasthemia.
- NIBP measurements are impossible with heart rate extremes of less than 40bpm or greater than 240bpm, or if the patient is on a heart-lung machine.

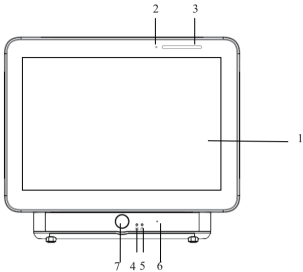
 The measurement may be inaccurate or impossible:
 - —with excessive and continuous patient movement such as shivering or convulsions;
 - ——if a regular arterial pressure pulse is hard to detect;
 - ----with cardiac arrhythmias;
 - ----with rapid blood pressure changes;
 - —with severe shock or hypothermia that reduces blood flow to the peripheries;
 - ----on a edematous extremity.
- RESP monitoring and apnea alarm based on chest impedance method are not suitable for patients with obstructive sleep apnea.
- Patients with chronic septicemia or hypercoagulable state cannot consider using this equipment. Because this equipment may cause suppurative or non-irritating thrombus;
- Patients with Parkinson's disease and tricuspid valve prolapse may be at risk of arrhythmia.
- Do not allow patients to wear ICG sensors when undergoing Magnetic Resonance Imaging (MRI) due to risk of severe burns.
- Before monitoring patients with pacemakers, ensure that the function of the pacemaker cannot be influenced by the measuring current used for impedance cardiography. In the case of minute ventilation pacemakers the use of the ICG

module is not allowed if the minute ventilation function of the pacemaker is activated.

Factors which can affect the accuracy of ICG measurement, as following:
——Septic shock;
——Aortic valve regurgitation and defect of septum;
——Severe aortic sclerosis, aortic prosthesis;
——Severe hypertension (Mean>130 mmHg);
——Cardiac arrhythmia;
——Tachycardia with a heart rate higher than 200 bpm;
——The patient's weight and height are out of range: Patient heights below
120 cm (48") or above 230 cm (90"); Patient weights less than 30 kg (67
lbs.) or greater than 155 kg (341 lbs.);
Aortic balloon or aortic balloon pump;
——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.
——Simultaneous use of electrical cautery systems during surgical procedures;

1.4. Main Unit

1.4.1. Front View



- 1. Display screen
- 2. Light induction: For example, when the environment is dark, the display brightness can be automatically adjusted.

3. Alarm lamp

Alarm lamp with different color and flashing frequency indicates the level of technical alarm and physiological alarm:

- ♦ High level alarm: the lamp quickly flashes red.
- Medium level alarm: the lamp slowly flashes yellow.
- ◆ Low level alarm: the lamp lights cyan without flashing.

4. Power indicating lamp

It is a LED that lights green and orange, the status of the LED is specified as follows:

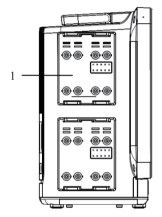
- Green: When the AC mains is connected.
- Orange: When the AC mains is not connected and monitor is powered by battery.
- Off: When the mains is not connected.
- 5. Charging indicating lamp

- ◆ Light up: When the battery is being charged.
- Off: When the battery is fully charged or no battery in monitor.
- 6. Phonetic holes.
- 7. Power switch

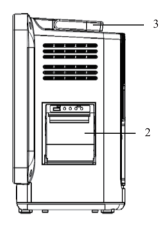
1.4.2. Side View

> P12

Left View

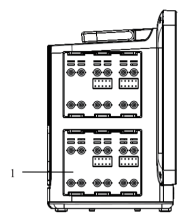


Right View



> P15&P18&P22

Left View



Right View

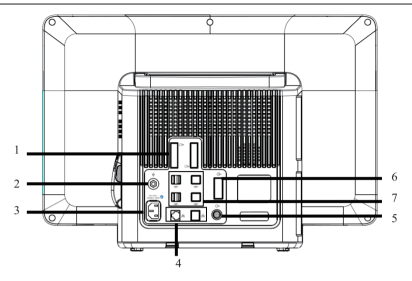
(1) External parameter module slots (2) Recorder (3) Handle



CAUTION: In order to prevent poor contact due to dust accumulated, please regularly clean the contact point according to actual application condition. Before cleaning, the monitor must be powered off. When cleaning, please wipe the point with medical cotton dipped into medicinal alcohol by use of a nipper.

1.4.3. Rear View

The rear views of P series monitors are different because of the size, but the interfaces are the same. Here is the rear view of P22 monitor.



1. DVI display connector

Connect to standard DVI display for secondary displaying.

2. Equipotential grounding terminal

When other devices and monitors are unified used, the wires should be used to connect the equipotential grounding terminal of other devices and monitors to eliminate the potential difference between different devices and ensure safety.

3. AC power socket

4. Network connector

The standard RJ45 interface, enabling networking with the central monitoring system, other bed communications and system upgrades via.

5. Nurse call connector

6. Auxiliary Module Slots connector

Connect with Auxiliary Module Slots.

7. USB connector

Standard USB2.0 connectors, which can be connected to USB devices such as U disk and barcode scanner.

1.5. Measurement Modules

1.5.1. Available Plug-in Modules

The plug-in modules are used to monitor the patient's physiological parameters and connect external devices. The monitor provides the following modules:

Modules	Functions	
P1 patient monitor	Can be used as a parameter module with LCD in P series monitor,	
	and it can simultaneously monitor ECG, Resp, CO2, SpO2, PR,	
	Temp, NIBP and IBP.	
Multi-parameter	It is a Multi-parameters measurement module without LCD, and it	
module (MPS-P)	can simultaneously monitor ECG, Resp, SpO ₂ , PR, Temp, NIBP	
	and IBP.	
IBP module	Support IBP monitoring.	
Temp module	Support temperature monitoring	
SpO ₂ module	Support SpO ₂ monitoring (Nellcor, Masimo)	
CO ₂ module	Support Mainstream CO ₂ , Sidestream CO ₂ and Microflow CO ₂	
	monitoring	
AG module	Support Mainstream AG and Sidestream AG monitoring.	
ICG module	Support ICG monitoring.	
BIS module	Support BIS monitoring.	
NIBP module	Support NIBP module (Suntech)	
EEG module	Support EEG monitoring.	
DM module	Support DM monitoring.	
RM module	Support RM monitoring.	
rSO ₂ module	Support rSO ₂ monitoring. (Masimo, Nonin)	
NMT module	Support NMT monitoring.	
BIOLINK module	Connects external device.	
	(Before use, please contact manufacturer to confirm if the external	
	device can be connected)	

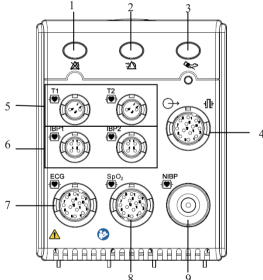
The multi-parameter modules (MPS-P) have the following configuration combinations:

Models	Configuration combinations
MPS-P01	Standard configuration: 3/5/6-lead ECG, RESP, BLT NIBP ,TEMP,
	BLT SpO ₂
MPS-P02	Standard configuration+12-lead
MPS-P03	Standard configuration+IBP
MPS-P04	Standard configuration+12-lead+ Multifunctional connector
MPS-P05	Standard configuration+IBP+12-lead+Multifunctional connector

Because different modules occupy different amount of slots, hence the amount of plug-in modules on the monitor may vary.

1.5.2. Example parameter Module

The plug-in modules have similar structures, and we take the MPS-P module as an example:



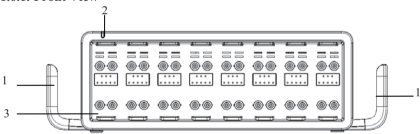
- 1. Alarm pause button: Pause all alarms.
- 2. Alarm reset button: To reset the alarm system.
- 3. NIBP button: To start or stop NIBP measurement.

- 4. Multifunctional connector: output analog signals and defibrillation synchronization signal.
- 5. Temp connector
- 6. IBP connector
- 7. ECG connector
- 8. SpO_2 connector (BLT SpO_2)
- 9. NIBP connector

1.6. Auxiliary Module Slots

The plug-in modules can be connected through Auxiliary Module Slots. The Auxiliary Module Slots has eight module slots. It is connected to the monitor (P15/P18/P22) through the Auxiliary Module Slots connector.

1.6.1. Front View



- 1. Handle
- 2. Status indicator: Lights green when the Auxiliary Module Slots is powered on.
- 3. Module slots

1.6.2. Rear View



1. Auxiliary Module Slots connector: Connect the P series monitor.

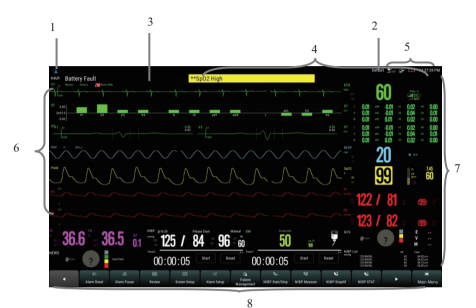
1.7. Applied Parts

The applied parts of the monitor are:

- ECG electrode and leadwire
- SpO₂ sensor
- Temp probe
- NIBP cuff
- IBP transducer
- ICG sensor
- CO₂ sampline tube
- AG sampline tube
- RM sensor
- EEG electrode
- BIS sensor
- NMT sensor and electrode
- rSO₂ sensor

1.8. Screen Display

The monitor adopts a display screen of high-resolution TFT LCD. Measurement parameters, waveforms, patient information, alarm area and menu can be displayed on the screen. Standard screen is shown as follows:



1. Patient info area

Shows the room number, bed number, patient name, patient category and so on. Select this area to enter the **[Patient Management]** menu and detailed description please to refer *Chapter 4 Patient Management*.

2. Current configuration

3. Technical alarm area

Display technical alarm information and prompt information. Cyclic displaying when there are multiple messages. Select this area to open the 【Alarm Informations】 menu to view the current technical alarm.

4. Physiological alarm area

Shows the physiological alarm messages, medium-level and low-level alarm messages display on the left, while the high-level alarm messages display on the right.

Cyclic displaying when there are multiple messages. Select this area to open the **[Alarm Informations]** menu to view the current physiological alarm.

5. System status information area

Display alarm volume, network and storage devices connection status, battery and system time. For the battery status icon please to refer *chapter 28 Battery*.

6. Waveform area

Show the waveforms of physiological parameter. Label displays on the top left corner of each waveform area. Select the waveform area of a parameter and enter the corresponding parameter setting menu.

7. Parameter area

It consists of various parameter areas, and shows parameter value, unit, alarm limit and alarm status, etc. Label displays on the top left corner of each parameter area. Select the parameter area of a parameter to enter the corresponding parameter setting menu.

8. Area of touch quick keys

Shows quick keys, these quick keys are used to conduct some common operations.

1.8.1. Screen Symbols

The following table shows the symbols and meanings displayed in the system information area:

Symbol	Note	Symbol	Note
?	Wireless network connected. The physical part represents the network signal strength.	⊕ X	Wireless network not connected.
	Wired network connected	-	Wired network not connected.
緻	All the alarms are paused.	\bigotimes	The parameter alarm is turned off or the alarm system of the monitor is turned off.
%	The alarm has been confirmed and the alarm system has been reset.	\bowtie	Audible alarm tones are turned off

Symbol	Note	Symbol	Note
	Indicates that the battery is fully charged.		Indicates that the battery is half charged.
	Indicates that the battery is empty and needs to be charged.		Indicates that the battery is almost depleted and need to be charged immediately, otherwise the monitor will automatically turn off.
<u>-</u> \d\d_+	Indicates that the battery is being charged.	*	Indicates that the monitor is being powered by AC power.
(X)	No battery is installed.		

1.8.2. Quick keys

Quick keys are displayed at the bottom of the monitor's main screen. Through the quick keys, you can easily and quickly access some functions or perform operations.

1.8.2.1. Quick keys list

The symbols on the quick keys are shown as follows:

Symbol	Quick key Note	Function
▼	Previous page	Previous page
	Next page	Next page
	Main Menu	Enter the main menu
\bigcirc	Standby	Enter the Standby mode
\triangle	Alarm Setup	Enter the 【Alarm Setup】 menu
***	Review	Enter the 【Review】 interface

Symbol	Quick key Note	Function
Q.	NIBP Measure	Enter the 【NIBP Measure】 menu
E S	NIBP Start/ Stop	Start/Stop NIBP measurement
€	NIBP Stop All	Stop all NIBP measurement
6 9	NIBP STAT	NIBP STAT measurement mode
P	Venipuncture	Start/Stop Assistant venipuncture
→ ()←	Zero	Start IBP, CO ₂ Zero
lacktriangle	Freeze	Freezes Waveforms
:	Alarm Reset	Alarm reset
緻	Alarm Pause	Pause the current alarms
	Screen Setup	Enter the 【Screen Setup】 menu
	Patient Management	Enter the [Patient Management] menu
2→	Discharge Patient	Discharge the current patient
Q» HIII	Sound	Enter the 【Sound】 setting menu
äill	Brightness	Enter the 【Brightness】 menu
<u>+</u>	Lock Screen/Unlock Screen	Disable/activate touch screen
<u> </u>	Wireless Setup	Enter the 【Wireless Setup】 menu
4	Intubation Status	Enter / exit Intubation status

Symbol	Quick key Note	Function
	Record Setup	Enter the 【Record Setup】 menu
\$	Realtime Record	Manually start/stop real-time recording
	Calculations	Enter the 【Calculations】 menu
ψψ	Remote View	Enter the 【Remote View】interface
	Night Mode / Exit Night Mode	Enter / Exit night mode
Q	Voice Assistant/Close Voice Assistant	Turn voice assistant on/off
目	Manual Event	Manually trigger and save events
2	Loops	Open the 【Loops】 window
<u>.</u>	Rescue Mode/Exit Rescue Mode	Enter/Exit the Rescue Mode
Ü	CCHD	Entering the CCHD Screening Page



■ The monitor provides the intubation status function during RESP, CO₂, AG and RM monitoring. See 7.12 Intubation status for details.

1.8.2.2. Setting the quick keys

You can set quick keys which need to be displayed on the interface, as follows:

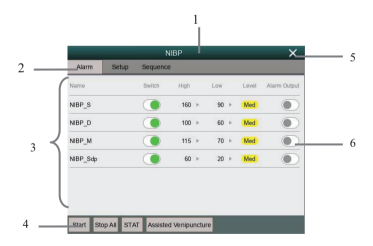
- 1. Enter the **[Quick Keys]** setting menu in one of the following ways:
 - > Select **[Screen Setup]** quick key → select **[Quick Keys]** submenu.
 - Select the 【Main Menu 】 quick key → select 【Quick Keys】 from
 【Display】 column.

2. Set the required quick keys:

- Add quick key: Select desired quick key from the 【Choices】 column on the left, and then select 【Add】.
- > Delete parameters: Select desired quick key from the **[Selected]** column on the right, and then select **[Delete]**.
- Move the display position of quick key: From the [Selected] quick key column on the right to select quick keys needed to be moved. And select [Move To Up], [Move To Down], [Move To Top] or [Move To Bottom] as needed.
- > Select **[Default Setting]** and the quick key settings will restore the factory default settings.

1.8.3. Menu

The styles of the various menus are basically similar, see the picture below:



- 1. Menu title: Summary of the current menu.
- 2. Submenu button: Press this button to enter the corresponding submenu.
- 3. Main display area of menu: Display menu options.
- 4. Operation button: Click to start an operation.

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- 5. Exit button: Exit the current menu.
- 6. Function switch:
 - Green: The function switch is on;
 - > Gray: The function switch is off.

Chapter 2 Safety

2.1. Safety Information



- The physiological data and alarm messages displayed on the monitor are for reference only and cannot be directly used for diagnostic interpretation.
- The monitor can only be used for a single patient at the same time
- This monitor can only be connected to a power outlet that has a protective ground. Do not use a removable multi-hole socket. If the power outlet is not connected to a grounding conductor, do not use the outlet and use a rechargeable battery to power the monitor.
- Before use, you must check the equipment, cables and accessories to ensure they work properly and safely.
- To avoid explosion hazard, do not use the monitor in the presence of flammable anesthetics or other flammable substance in combination with air, oxygen-enriched environments, or nitrous oxide.
- Do not open the shell of the device; otherwise the electric shock hazard may result. All maintenance and upgrades must be operated by the personnel trained and authorized by manufacturer only.
- Do not use the monitor in nuclear magnetic resonance (MR) environments.
- Do not touch the patient and live parts simultaneously. Otherwise patient injury may result.
- Do not come into contact with the patient during defibrillation. Otherwise serious injury or death could result.
- Do not rely exclusively on the audible alarm system for patient monitoring. Adjustment of alarm volume to a low level or off may result in a hazard to the patient. Remember that alarm settings should be customized according to different patient situations and always keeping the patient under close surveillance is the most reliable way for safe patient monitoring.
- Carefully place the power cord and various accessory cables to avoid risk of

entanglement or suffocated, the cables entangled, or subject to electrical interference.

- To avoid danger or environment pollution, the packaging materials must be handled in accordance with local regulations or the hospital's waste disposal system. Packaging materials must be placed out of reach of children.
- Ensure that the equipment is supplied with continuous electric power during work. Sudden power failure may cause data loss.
- The user should periodically check and move the sensor on the skin to avoid adverse skin or tissue effects.



∆ CAUTION:

- To ensure patient safety, use only parts and accessories specified in this manual.
- Magnetic and electrical fields are capable of interfering with the proper performance of the monitor. For this reason make sure that all external devices operated in the vicinity of the monitor comply with the relevant EMC requirements. Mobile phone, X-ray equipment or MRI devices are a possible source of interference as they may emit higher levels of electromagnetic radiation.
- To avoid contamination or infection of personnel, environment or other equipments, the equipment and its accessories that meet the service life must be disposed in accordance with relevant local regulations or hospital systems.
- The lifetime of the patient monitor is 5 years.
- Before connecting the equipment to the power line, check that the voltage and frequency ratings of the power line are the same as those indicated on the equipment's label or in this manual.
- Always install or carry the equipment properly to avoid damage caused by drop, impact, strong vibration or other mechanical force.
- The monitor must be wiped dry immediately after exposure to rain or splashes.
- Please do not mix different types and brands of electrodes. Mixing the electrodes may result in a large baseline drift or a long baseline recovery time after defibrillation. It is forbidden to use dissimilar metal electrodes, which

may cause high polarization voltages.



NOTE:

- Put the monitor in a location where you can easily see the screen and access the operating controls.
- The equipment use a mains plug as isolation means to the mains power. Do not locate the equipment in a place difficult to operate the mains plug.
- In normal use, the operator should stand in front of the device
- Keep this manual near the device so that it can be easily and timely obtained when needed.
- This manual describes the product in the most complete configuration. This manual describes all features and options. Your monitor may not have all of them.

2.2. Equipment Symbols

Your device may not have all of the symbols below.

Symbol	Note	Symbol	Note
1	Defibrillation-proof Type BF applied part	ECG	Abbreviation of "Electrocardiogram".
•	Defibrillation-proof Type CF applied part	SpO ₂	Abbreviation of Pulse "Oxygen Saturation".
\triangle	Attention: Consult accompanying documents (this manual).	ТЕМР	Abbreviation of "Temperature".
((•))	Non-ionizing radiation	CO ₂	Abbreviation of "Carbon dioxide".
4	Dangerous voltage	NIBP	Abbreviation of "Non-invasive Blood Pressure".
$\frac{1}{\sqrt{1}}$	Equipotential grounding	IBP	Abbreviation of "Invasive Blood Pressure".
\rightarrow	Auxiliary output	DM	Abbreviation of "Drip Monitor".

_	Alternating current (AC)	ICG	Abbreviation of "Impedance
	Alternating current (AC)	ICG	Cardiograph".
4Пь	Defibrillator synchronization	BIS	Abbreviation of "Bispectral
716	output connector	DIS	Index".
AAA	Manufacturer	EEG	Abbreviation of
		220	"Electroencephalograph".
IP21	Degree of protection against	AG	Abbreviation of "Anesthetic
	ingress of liquid		Gas".
晃	Network	RM	Abbreviation of "Respiratory
			Mechanics".
\hookrightarrow	DVI display connector	rSO ₂	Abbreviation of "Regional
	1 7		oxygen saturation".
_			Abbreviation of
•	USB	NMT	"Neuromuscular
			Transmission".
\sim	Manufacture date	RESP	Abbreviation of "Respiration".
(€ ₀₁₂₃	CE mark	EC REP	Authorized representative in
0123	OL mark	LO INLI	the European Community.
SN	Serial number	LOT	Batch code
↔	Input/output	(2)	Refer to this user's manual.
\wedge	Warning: the protection against	the effects of the	e discharge of a cardiac
	defibrillator is dependent upon the	e appropriated c	able
7	Symbol for the marking of elect	rical and electro	onics devices according to Waste
13	Electrical and Electronic Equipme	ent Directive.	

2.3. Packaging Symbols

Symbol	Symbol Note
	Fragile. Handle with care.

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Symbol	Symbol Note
	This Side Up.
	Keep dry.
	Stacking layer limit, where 'n' represents the maximum permissible number of layers. $(N = 6)$.

Chapter 3 Basic Operations

3.1. Installation



WARNING:

- The equipment should be installed by personnel designated by the manufacturer.
- The copyright of the software of this device belongs to the manufacturer. No organization or individual can tamper with, copy or exchange any infringement by any means or form without permission.
- Equipment connected to the analog or digital interfaces must comply with the respective IEC standards (e.g. IEC 60950 for data processing equipment and IEC 60601-1 for medical equipment). Furthermore all configurations shall comply with the current version of the standard for SYSTEMS IEC 60601-1. 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 current version of the requirements of the system standard IEC 60601-1. If in doubt, consult the technical service department or your local representative.
- When the equipment is connected to other electrical equipment into a combination with specific functions, if it is impossible to determine whether the combination is dangerous (for example, the electric shock hazard caused by the accumulation of leakage current), please contact the expert of the company or the hospital to ensure that the necessary safety of all equipment will not be damaged in the combination.

3.1.1. Unpacking and Checking

1. Unpacking

Before unpacking, examine the packing case carefully for signs of damage. If any damage is detected, contact the carrier. If the packing case is intact, open the package.

2. Remove the monitor and accessories carefully.

- 3. Keep all the packaging materials for future use in transportation or storage.
- 4. Check the monitor and accessories

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



- Keep the packing materials out of children's reach. Disposal of the packing materials should comply with local regulations or the hospital's waste disposal system.
- The monitor might be microbial contaminated during storage and transport. Before use, please verify whether the packages, especially the package of disposable accessories are intact. In case of any damage, do not apply it to the patient.

3.1.2. Environmental equipments

The operating environment of the equipment must meet the specifications in this manual.

The operating environment of the equipment should also reasonably free from the noises, vibration, dust, corrosive or flammable, explosive substances. If it is installed in the cabinet, make sure that there is enough space in front of the cabinet for operation, maintenance and repair; In order to maintain ventilation, the equipment should be at least 2 inches (5cm) away from around the cabinet.

When the equipment is transferred from one place to another, condensation may occur due to differences in temperature or humidity. At this point, you must wait for the condensation to disappear before you can use the equipment.



Please ensure the monitor is working under specified conditions; otherwise, the technical specifications mentioned in this manual will not be met, thus possibly leading to damage of equipment and other unexpected results.

3.2. Getting Started

3.2.1. Connecting the Power

Connecting the AC power

When the monitor need to be supplied by AC power, you can plug one end of the AC power cord into the AC power connector on the back of the monitor and the other end plug into the AC power outlet.



- Always use the accompanying power cord delivered with the monitor.
- Use the battery if the integrity of the protective earth conductor or the protective earthing system in the installation is in doubt.

Using battery

The monitor can be powered by a lithium battery. After the battery is installed, if the external power supply is suddenly interrupted, the monitor can automatically use the lithium battery to supply the power. For the use of the battery, please refer to *Chapter 28 Battery*.

3.2.2. Connecting the Input Device

Connect the keyboard, mouse and barcode scanner if necessary.

3.2.3. Connecting the Auxiliary Module Slots

Use the Auxiliary Module Slots cable to connect the monitor and Auxiliary Module Slots.

3.2.4. Installing the Plug-in Module

Insert the modules into the host monitor's module slots. A click is heard when the module is pushed into place.

3.2.5. Removing the Plug-in Module

Lift the latch at the bottom of the module and pull it out.



■ To prevent the P1 from falling, catch it with another hand while pulling it out.

3.3. Starting the Monitor

After installing the monitor, you can monitor the patient.

- 1. Before powering on the monitor, please check whether there has mechanical damaged, external cables and accessories connect correctly.
- 2. Plug the power cord into an AC outlet. If using battery power, make sure there is enough power in the battery.
- 3. When pressing the power switch, the alarm light will display red, yellow and cyan in turn. After the alarm light is turned off, the screen will display the startup interface. After the system emits a beep, the startup screen disappears and enters the main monitoring interface.



■ If the monitor is damaged or does not work properly, do not use it for any monitoring procedure on a patient. And then please contact the maintenance personnel or the manufacturer immediately.

3.4. Starting Monitoring

- 1. Decide what parameters should be monitored or measured.
- Connecting required cables and sensors.
- 3. Check whether the connection of cables and sensors is correct.
- 4. Check whether all kinds of settings are correct, for example: **[Patient Type]** and **[Paced]**. For detailed information on the measurement or monitoring of each

parameter, please refer to the corresponding chapters.

3.5. Operation and Browse

Almost every element on the screen is interactive. Screen elements include parameter values, waveforms, quick keys, information area, alarms area, menus, etc. Often you can access the same element in different ways. For example, you can access a parameter setup menu by selecting corresponding parameter area or waveform area.

3.5.1. Using the Touch screen

Click on the touch screen to complete some operations quickly and easily.

In order to prevent misoperation of the touch screen, you can operate the **【Lock Screen】** quick key to temporarily lock the touch screen. After the touch screen is locked, the **【Lock Screen】** quick key become **【Unlock Screen】**, and its background color is blue, which indicating that the touch screen operation is disabled.

The lock time of the touch screen can be customized, the steps are as follows:

- 1. Enter the **[Other]** interface in the one of the following ways:
 - ◆ Select **[Screen Setup]** quick key → select **[Other]** submenu;
 - ◆ Select [Main Menu] quick key → from [Display] column to select [Other] submenu.
- 2. Set the **[Screen Lock Duration]**. The touch screen is unlocked under the following conditions:
 - When the set lock screen duration is reached, the touch screen is automatically unlocked.
 - Select the **[Unlock Screen]** quick key to unlock the touch screen.



- Check the touch screen whether it is damaged or breakage before use. If it is found to be damaged or broken, please stop using the monitor immediately and contact the service personnel.
- If you find that the touch screen is loose, stop using the monitor immediately and contact your service personnel.

3.5.2. Using the Mouse

The monitor supports the mouse with USB connector. You can use the mouse to select a screen element by moving the cursor on the element and then click on it.

3.5.3. Using the Barcode scanner

The monitor supports the barcode scanner to input patient's medical record number or registration number, and connects to the monitor through USB interface

3.5.4. Using the On-Screen Keyboard

The on-screen keyboard enables you to enter information:

- Use the key to delete the previous character.
- > \text{\text{\$\text{\$\Delta}\$}} Use the key to toggle between uppercase and lowercase letters.}
- Use the key to confirm what you have entered and close the onscreen keyboard.
- > Use the key to clear the entered character
- > **@#%** Use the key to access the symbol keyboard.
- > **ABC** Use the key to return to alphabetic keyboard.

3.6. General Settings

This chapter only introduces the general settings. For the setting of parameters and other functions, please refer to the corresponding chapter.

3.6.1. Language Settings

- Select 【Main Menu】 → from 【System】 column to select 【Maintenance】
 →enter the maintenance password.
- 2. Select **Other** submenu.
- 3. Select **[Language]** and then select the desired language.
- 4. Restart the patient monitor.

3.6.2. Adjusting the Screen Brightness

The steps to set the screen brightness are as follows:

- 1. Set the brightness in one of the following ways:
 - ♦ Select **Brightness** quick key.
 - ◆ Select **Screen Setup** quick key→select **Other** submenu.
 - ◆ Select [Main Menu] quick key → from [Display] column to select
 [Other] submenu.
- 2. If the monitor operates on AC power, please set [Brightness]; If the patient monitor operates on battery power, please set [Brightness On Battery].
- 3. When the **[Brightness]** set as **[Auto]**, the screen will change to the brightness automatically according to the environment light intensity.



NOTE:

- When the patient monitor enters standby mode, the screen brightness will be automatically adjusted to the lowest.
- When the AC power is interrupted and the battery is powered, the screen brightness is automatically set to the corresponding brightness when the battery is powered. You can still manually adjust the brightness as needed.

3.6.3. Setting the Date and Time

- 1. Enter the **[System Time]** page in one of the following ways:
 - ➤ Select [Main Menu] → from [System] column to select [Time], enter the [System Time].
 - Click the system time in the system status information area on the monitor to enter the 【System Time】 page.
- 2. Select [Date] and [Time] to set the current date and time.
- 3. Select [Date Format].
- 4. If you need to use 12 hours format, turn off **[24-Hour]**.

If the monitor is connected to the central monitoring system, the system time of the monitor will be automatically adjusted according to the time of the central monitoring system.



■ When starting to use the monitor, please modify the date and time of the device according to the local time. Incorrect date and time setting may lead to misjudgment of patient trend data.

3.6.4. Adjusting Volume

Select [Sound] quick key or the sound icon in the system status information area to enter the page, set [Alarm Volume], [High Alarm Volume], [Reminder Volume], [QRS Volume], [Touch Tone] and [NIBP End Tone] switch respectively.

3.7. Measurement Settings

3.7.1. Setting Parameters

You can manually turn the parameter switch on or off. The steps to set the parameter switch are as follows:

- 1. Enter **[Parameter Switch]** interface in one of the following ways:
 - ◆ Select [Screen Setup] quick key→select [Param Switch] submenu.
 - ◆ Select [Main Menu] quick key, from [Parameter] column to select [Parameter Switch].
- 2. Turn on or off the corresponding parameters as needed

When a parameter is off, the monitor interface will not display the parameter value and waveform.

3.7.2. Setting Display Screen

You can set the parameter waveform and its order displayed in the normal interface

as required. The steps are as follows:

- 1. Enter **[Screen Layout]** interface in one of the following ways:
 - ➤ Select 【Screen Setup 】 quick key→select 【Screen Layout 】 submenu.
 - ➤ Select 【Main Menu】 quick key→from 【Display】 column to select 【Screen Layout】.
- 2. Select a parameter area or waveform. From the pop-up parameter list to select the elements needed to display in the area. The selected parameters and waveforms are displayed according to the set position. The parameters and waveforms that are not selected will not be displayed on the interface.



NOTE: The first line of the parameter area and the waveform area always

displays the ECG parameters and the first ECG waveform.



■ The parameters of the 【Screen Layout】 are not assigned to the display area, which will not be displayed on the monitor interface and relevant alarms for this parameter will still be provided

3.7.3. Setting the Parameter

Each parameter has an independent setting menu, through which alarm and parameter setting can be modified. You can enter the Parameter Setting Menu in the following ways:

- > Select the parameter area or waveform area of a parameter
- Select [Main Menu] quick key, from [Parameter] column to select
 [Setup], and then select corresponding parameter.

3.8. Operation Mode

3.8.1. Monitor Mode

Monitor mode is the most common clinical working mode used to monitor patient. When the monitor is turned on, it automatically enters the monitor mode.

3.8.2. Standby Mode

You can temperately stops patient monitoring without switching off the monitor by entering the standby mode. You can enter the Standby Mode in the following ways:

3.8.2.1. Entering the Standby Mode

Select either way showed in the following to enter the Standby Mode:

- Press [Standby] quick key, or
- Select 【Main Menu】 quick key→from the 【Patient】 column to select
 【Standby】 button, or
- ➤ Select 【Patient Management】 quick key→discharging patient and the enter Standby Mode.

The monitor behaves as follows after entering the standby mode:

- > Stops all parameter measurements.
- Disables all the alarms and prompt messages, except for the battery low alarm
- > Turns screen brightness to the dimmest after entering the standby mode for 30 seconds.



Pay attention to the potential risk of placing the monitor to standby. In the standby mode, the monitor stops all parameters' measurements and disables all the alarm indications, except for the battery low alarm.

3.8.2.2. Exit the Standby Mode

To exit the standby mode, choose any of the following ways:

- Select Resume Monitor to exit the standby mode and resume monitoring the current patient.
- > Select [Discharge Patient] to discharge the current patient.

If the monitor automatically enters the standby mode after a patient is discharged, choose any of the following ways to exit the standby mode:

- > Select [Patient Management] to exit the standby mode and admit a new patient.
- Select [Monitor] to enter the patient information for preparing to admit a new patient.

3.8.3. Demo Mode

In Demo mode, the monitor can demonstrate its major functions when patient or patient simulator is not connected. The Demo mode is password protected

Choose the following way to enter the Demo Mode:

- 1. Select 【Main Menu】→from 【System】 column to select 【Demo】.
- 2. Input the password→select 【OK】.

Shut down and restart to exit the Demo Mode.



■ The demonstration function is mainly used to display machine performance and to train users. In the actual clinical use, it is forbidden to use the demonstration function, so as to prevent the medical staff from mistakenly thinking that the monitor displays the waveform and parameters of the monitored patient, thus affecting the patient's monitoring and delaying the diagnosis and treatment.

3.8.4. Night Mode

Night mode is a special clinical monitoring mode. Under the night mode, the alarm volume, QRS volume and screen brightness turn to be lowest automatically. To avoid disturbing the patient, night mode may be used.

3.8.4.1. Entering Night Mode

The steps of entering [Night Mode] as following:

- Select 【Night Mode】 quick key or select 【Main Menu】 quick key→from
 【Display】 column to select 【Night Mode】 button.
- 2. In the pop-up menu, set the night mode.
- 3. Select [Enter Night Mode].

The night mode settings are as follows by default:

- Brightness: 1
- Alarm Volume: 2
- ORS Volume: 1
- Touch Tone: Off
- NIBP End Tone: Off



■ Verify the settings of brightness, alarm volume, QRS volume and key tone before entering the night mode. Pay attention to the potential risk if the setting value is low.

3.8.4.2. Exit the Night Mode

To exit the night mode, follow this step:

- Press [Exit Night Mode] quick key or select [Main Menu] quick key→from [Display] column to select [Exit Night Mode] button.
- 2. In the pop-up box to select **(OK)**

F

NOTE: The monitor resumes the previous settings after exiting the night mode.

3.8.5. Privacy Mode

The privacy mode is a special monitoring mode. In the privacy mode, the monitor does not display patient information and monitoring data to protect patient information from non-clinicians such as visitors.

The privacy mode is only available when the patient admitted by the monitor is also monitored by the Central monitoring system. The monitor continues monitoring the patient, but patient data is only visible at the Central monitoring system.

3.8.5.1. Entering the Privacy Mode

To enter the privacy mode, choose either of the following ways:

◆ Select 【Main Menu】 quick key→from the 【Display】 column to select 【Privacy Mode】 button→Select 【OK】.

The monitor has the following features after entering the privacy mode:

- The screen turns blank, and prompt **[Being monitoring]** at same time.
- All parameters and waveforms display are shield.
- Except for the low battery alarm, the monitor inactivates alarm tone and alarm light of all other alarms.
- The monitor suppresses all system prompt tone, including heart beat tone, pulse tone, etc.



■ In Privacy mode, all audible alarms are suppressed and the alarm light is deactivated at the monitor. Alarms are presented only at the Central monitoring system. Pay attention to potential risk.



■ Cannot enter the privacy mode if a low battery alarm occurs.

3.8.5.2. Exit the Privacy Mode

The monitor automatically exits the privacy mode in any of the following situations:

- ◆ The monitor disconnects from the Central Monitoring System.
- ◆ The low battery alarm occurs.

You can also press **[Exit Privacy Mode]** on the screen to manually exit the privacy mode.

3.9. Using the Timer

The monitor provides timer function, which can display up to four timers at the same time. You can set each timer separately, which prompts when the set time arrives.

3.9.1. Displaying Timer

To display a timer, follow this procedure:

- 1. Access **(Screen Layout)** in either of the following ways:
 - ◆ Select 【Screen Setup 】 quick key→Select 【Screen Layout 】 submenu.
 - ◆ Select [Main Menu] quick key→from the [Display] column to select [Screen Layout].
- 2. Click the parameter area where you want to display the timer, select **[Timer]**
 - → select [Timer1], [Timer2], [Timer3], [Timer4].

3.9.2. Controlling the Timer

The timer provides the following controls:

[Start]: Start the timer.

[Pause]: Pause the timer.

Resume: Resume the timer.

Reset: Clear the current timing results and reset the timer.



■ Do not use the timers to schedule critical patient-related tasks.

3.9.3. Setting the Timer

You can set each timer independently. To set the timer, follow this procedure:

- 1. Select the timer area to enter the **[Timer]** menu.
- 2. Set 【Timer Type】:
 - ♦ 【Normal】: The timer is timed according to the present 【Run Time】.

 Stop timing when running time is reached.
 - ◆ 【Cycled】: The timer cycles, that is, the timer counts according to the preset 【Run Time】, and starts timing again after reaching the run time. The timer area displays the number of timer cycles.
 - ◆ 【Unlimited】: The timer displays the time elapsed since the timer was started.
- 3. Set [Direction]
- 4. Set [Run Time].
- 5. Set [Reminder Volume] When the remaining time is 10 seconds, the monitor issues a reminder tone and the timer flashes in red, prompting you that the run time is to expire.



- Cannot change timer settings when a timer is running.
- Set [Direction], [Run Time] and [Reminder Volume] only for [Normal] or [Cycled].

3.10. Voice Assistant

The voice assistant can be used as an auxiliary input interface. You can issue specific control commands to the monitor via voice. After the monitor recognizes the command, it performs the corresponding operation function.

The method of operating the voice assistant as following:

- 1. Select Voice Assistant quick key, open the voice assistant function, the icon
 - " will display on the top of status bar.

- Say the wake-up words. At this time, the top status bar will display the " icon, indicating that the monitor is in the state of voice recognition control command.
- 3. When the control command is spoken, the monitor will recognize the command and perform the corresponding operation, at this time, " icon disappears and the " ucon is displayed.



WARNING:

■ The voice input function is only used as the auxiliary input function of monitor.

The specific operation shall be subject to the touch screen of mouse operation.



CAUTION:

- Every time before you say a control command, you must say the wake-up word to wake up the voice assistant at first. After waking up (the top status bar displays the " icon), if the control command is not spoken within 7 seconds, the voice assistant will enter the sleep state again (top The status bar displays the " icon), you need to re-execute the "Wake-up" → "Control Command" voice operation.
- User can configure the voice commands that need to be supported according to actual application or for risk considerations. (Click the "☑" icon to pop up a list of voice commands, select the commands that need to be supported.)
- When using the voice assistant, try to be as close as possible to the monitor.
- Please avoid using the voice assistant in noisy environments.



NOTE:

■ The specific wake-up words and supported voice commands will show in the voice assistant screen.

3.11. Shutting off the Monitor

Please follow the below steps to shut off the monitor:

- 1. Confirm that the patient monitoring is finished.
- 2. Disconnect the cables and sensors form the monitor.
- 3. Confirm that the monitoring data is stored or cleared.
- 4. Press the power switch for several second, the monitor interface will pop up the shutdown dialog box, click OK to shutting off the monitor.



Although not recommended, you can press and hold the power on/off switch for 5 seconds to forcibly shut down the monitor when it could not be shut down normally or under some special situations. This may cause loss of data of the patient monitor.

ॐNOTE:

- The AC power supply of the monitor is not cuff off through the power switch.

 To completely disconnect the power supply, unplug the power cord.
- In case of a temporary power failure, the monitor retains patient data before shutdown, including patient monitoring data and configuration data.

Chapter 4 Patient Management

4.1. Discharging a Patient

Before monitoring a new patient, discharge the previous patient. After the patient is discharged, all patient data, including patient information, trend data, and physiological alarm information is be deleted from the monitor, the technical alarms are reset, and monitor settings returns to their defaults (current configuration or user-specified configuration). For more information, see *5.2 Setting Default Configuration*.

After discharging a patient, the monitor automatically admit a new patient.



Always discharge the previous patient before starting monitoring a new patient.
Failure to do so can lead to data being attributed to the wrong patient.



■ Discharging a patient deletes all history data of current patient in the monitor.

Discharge a patient manually using either of the following methods:

- ♦ Select **[Discharge Patient]** quick key.
- ◆ Select the patient information area at the top left of the screen→Select

 【Discharge Patient】
- ◆ Select 【Patient Management】 quick key→Select 【Discharge Patient】
- ◆ Select Main Menu from Patient column to select Discharge Patient
 In the pop-up dialog box to select:
- > 【OK】: All patient data, including patient information, trend data, and physiological alarm information, are cleared. The technical alarm status is reset and the system reverts to its default configuration and enters into the standby screen.
- > 【Cancel】: Exit the operation of discharging patient data and return to the main interface.

4.2. Admitting a Patient

The monitor admits a new patient in the following situations:

- After discharging a patient, the monitor automatically admit a new patient.
- From the standby screen, select [Discharge Patient] to admit a new patient.

Always inputs patient information as soon as the patient is admitted. For more information please refer to *4.3.2 Editing Patient Information* for details.



- 【Patient Type】 and 【Paced】 will always contain a value, regardless of whether the patient is fully admitted or not. If you do not specify settings for these fields, the patient monitor uses the default settings from the current configuration, which might not be correct for your patient.
- For paced patients, you must set 【Paced】 to 【Yes】. If it is incorrectly set to 【No】, the patient monitor could mistake a pace pulse for a QRS and fail to alarm when the ECG signal is too weak.
- For non-paced patients, you must set 【Paced】 to 【No】.

4.3. Managing Patient Information

4.3.1. Patient Management menu

Use any of the following methods to enter the **[Patient Management]** menu:

- > Select patient information area at the top left corner of the screen.
- > Select [Patient Management] quick key.
- Select [Main Menu] → from [Patient] column to select [Patient Management].

4.3.2. Editing Patient Information

Edit patient information after a patient has been admitted, or when patient information is incomplete, or when you want to change patient information.

To edit patient information, follow this procedure:

- Enter the 【Patient Management】 menu. For more information, please refer to 4.3.1 Patient Management menu.
- Select patient type according to the actual situation: [Adult], [Pediatric],
 [Neonate].
- 3. Edit patient information as required.

If your monitor is connected with the barcode scanner, you can enter the medical record number by scanning the barcode.



- The setting of patient type determines the algorithm used by the monitor to process and calculate certain measurements, as well as the safety limit and alarm limit range applicable to certain measurements.
- The monitor reloads the configuration when the patient type is changed.

4.3.3. Setting the Display Item

You can set whether to display and edit patient room number, middle name, race, age, and so on in the **[Patient Management]** menu by following these steps:

- Select 【Main Menu 】 quick key→from 【System 】 column to select 【Maintenance】→input password→Enter.
- 2. Select 【Patient Management】 submenu→ 【Field】 submenu.
- 4. Select customizes the patient information section, if necessary, and enters the name of the section.

4.4. Transferring Patient Data

The P1 monitor has the function of uninterrupted monitoring. That is, P1 can realize patient transferring between different monitors. Patient transferring contains: patient information, patient measurement setup (such as: alarm limit), parameter measurement setup and patient trend data.



- Do not discharge a patient before the patient is successfully transferred.
- After a patient is successfully transferred, check if the settings of monitor (especially patient category, paced status, alarm limits settings, and etc) are appropriate for this patient.

■ NOTE:

■ The system automatically switches on the HR alarm and lethal arrhythmia alarm after transferring the patient data.

4.4.1. Setting the Data Transfer Strategy

When P1 is connected to the monitor, P1 automatically uploads the data to the monitor if the patient information in the monitor is consistent with those of in P1. However, the monitor needs a data transfer strategy if the patient information in the monitor is not consistent with those of in the P1. To set the data transfer strategy, follow these procedures:

- Select 【 Main Menu 】 quick key→from 【 System 】 column to select 【 Maintenance】 →input password→Enter.
- 2. Select 【Patient Management】 submenu→ 【Transfer】 submenu.
- 3. Set 【Data Transfer Strategy】.
 - ◆ 【Always Ask】: Monitor will ask for strategy each time it detects inconsistent information.
 - ◆ 【Continue Patient in Module】: The monitor continues to use the patient data of P1 when it detects inconsistent information. At the same time, discharge the patient, and automatically admits a new patient and copies the data of P1.
 - ♦ 【Continue Patient in Monitor】: The monitor continues to use the

patient data in the monitor when it detects inconsistent information. At the same time, delete all patient data of P1 and copies the data in the monitor to the P1.

4.4.2. Transferring Patient Data via P1



■ While transferring a patient with EMS, it must be sure that the EMS is battery-powered and with enough capacity.

To transfer the patient data via P1, insert the P1 into the module racks of the monitor.

- P1 automatically uploads the data to the monitor if the patient information and ID number in the monitor is consistent with those of in P1.
- If the patient information in the monitor is not consistent with those of in the P1 and 【Data Transfer Strategy】 is set to 【Always Ask】 (see 4.4.1 Setting the Data Transfer Strategy for details), the monitor prompts the 【Select Patient】 menu automatically. In this case, you need to select an operation (see the following table) according to the actual situation.

Operations	Description	Examples of applications
【 Continue Patient	Continue to use the patient data in	1. Replace P1 during
in Monitor]	the monitor. At the same time, patient monitoring	
	delete all patient data of P1 and	2. After the patient is
	copies the data in the monitor to	admitted, connect the
	the P1.	P1.
Continue Patient	Continue to use the patient data	You are monitoring a
in Module	of P1. At the same time,	patient using P1, but the
	discharge the patient, and	patient needs to be
	automatically admits a new	transferred, e.g. from a
	patient and copies the data of P1.	ward (original monitor) to
		the operating room

		(destination monitor).
[New Patient]	If it is confirmed that the patient	Inserting P1 into the
	information in the monitor and P1	monitor before admitting a
	are not correct, this option will	new patient. However, the
	clear all patient data on the	monitor and/or P1 have
	monitor and P1 at the same time,	stored the previous
	and the monitor and P1 will admit	patient's data and settings.
	a patient again. In this case, you	
	need to re-enter the patient	
	information. And the monitor will	
	restore the settings according to	
	the patient category.	
【Same Patient】	Select this option if the patient	A patient monitored with
	information in the monitor and P1	the P1 is moved to another
	are different but you are sure that	department and again
	it is the same patient. This merges	moved back.
	the patient's trend data in the	However, the patient
	monitor and P1 and copies the	information stored in the P1
	settings in P1 to the monitor as	was altered before
	well.	connected to original
		monitor.



If you select **[**Apply Module Settings **]**, the P1 settings can be transferred to the monitor along with the patient data. See 4.4.4 Transferring the P1 settings.

4.4.3. Setting the duration of data transferred by P1

When using P1 to transfer the patient data, you can set the data transfer duration through the following steps:

1. Select 【 Main Menu 】 quick key→from 【 System 】 column to select

【Maintenance】 →input password→Enter.

- 2. Select 【Patient Management】 submenu→ 【Transfer】 submenu.
- 3. Set [Data Transfer Duration].

4.4.4. Transferring the P1 settings

When transferring the P1 data, the P1 settings can also be transferred. You can set the transferring of P1 settings through following steps:

- Select 【 Main Menu 】 quick key→from 【 System 】 column to select 【 Maintenance 】 →input password→Enter.
- 2. Select 【Patient Management】 submenu→ 【Transfer】 submenu.
- 3. Set **[Upload Module Settings]**.

4.5. Connecting to a Central Monitoring System

You can connect the monitor to the central monitoring system (CMS) manufactured by Biolight through wired LAN or wireless LAN. When connected to the CMS, the system provides the following functions:

- > The monitor can transmit patient information, real-time monitoring or measurement data, alarm limits, alarm levels, alarm messages, prompts, and various settings to the central monitoring system.
- The central monitoring system and the monitor display synchronously, and some functions can be controlled bidirectionally. For example, change patient information, receive patient data, cancel patient data, start or stop NIBP measurement, etc.
- Alarm delay to the central monitoring system: the alarm delay time from this equipment to the central monitoring system is $\leq 2s$.

For more information on the CMS, refer to the Central Monitoring System's instructions for use. For more information about connection of monitor and CMS, refer to the *27.2.5 Connecting the Central Monitoring System (CMS)* in this manual.

Chapter 5 Managing Configurations

5.1. Introduction

When performing continuous monitoring on a patient, the clinical professional often needs to adjust the monitor's settings according to the patient's condition. The collection of all these settings is called a configuration. The configuration of the monitor includes: parameter configuration, alarm configuration and monitor configuration. You can change some settings from a certain set of configuration and then save the changed configuration as a user configuration.



■ The configuration management function is password protected. The configuration management tasks must be performed by clinical professionals.

5.2. Selecting default configuration

The default configuration can be the factory default configuration or a user configuration that has been stored.

You can use the following steps to select the default configuration:

- Select 【Main Menu】 quick key→from 【Configuration】 column to select
 【Set Default Config】 →input maintenance password→Enter.
- 2. Select **[Factory Default]** or **User-defined configuration.**

When selecting user-defined configuration, only one user configuration that has been stored under the current patient type can be selected.

5.3. Saving current settings

Current settings can be saved as user configuration. Up to 10 user configurations can be saved. The steps to protect the current settings are as follows:

- Select 【Main Menu】 quick key→from 【Configuration】 column to select
 【Save User Config】 →enter maintenance password→Enter.
- 2. In the popup dialog box of **[Save User Config]**, input the configuration name.

3. Select **[OK]**

5.4. Deleting a configuration

You can delete the saved user configurations by following these steps:

- Select 【Main Menu】 quick key→from 【Configuration】 column to select
 【Delete User Config】 →input maintenance password→Enter.
- Select the configuration you want to remove:
 In the Delete User Config Imenu, the currently existing user configuration on the monitor is displayed.
- 3. Select **[Delete]** button.
- 4. In the popup dialog box to select **[OK]**.



■ The current configuration in use cannot be deleted.

5.5. Transferring a configuration

The monitor provides configuration transfer function. You can use a USB drive to transfer the configuration from one monitor to another monitor that needs the same settings without having to reset them item by item. The monitor supports transferring the monitor configuration with USB disk.

5.5.1. Exporting a configuration

You can export the monitor's current user configuration to a USB drive by following these steps:

- 1. Connect the USB drive to the monitor's USB port.
- Select 【Main Menu】 quick key→from 【Configuration】 column to select
 【Export User Config】 →input maintenance password→Enter.
- 3. Select configuration that needs to be exported.
- 4. Select [OK].

5.5.2. Importing a configuration

You can import the user configuration to the monitor through a USB drive by following these steps

- 1. Connect the USB drive to the monitor's USB port.
- Select 【Main Menu】 quick key→from 【Configuration】 column to select
 【Import User Config】 →input maintenance password→Enter.
- 3. Select configuration that needs to be imported.
- 4. Select [OK].

5.6. Loading current configuration

You may make changes to some settings during operation. However, these changes or the pre-selected configuration may not be appropriate for the newly admitted patient. Therefore, the monitor allows you to load a desired configuration so as to ensure that all the settings are appropriate for your patient.

To load a configuration by following these steps:

- Select 【Main Menu】 quick key→from 【Configuration】 column to select 【Load Current Config】.
- 2. Select configuration that needs to be loaded.
- 3. Select [OK].

5.7. New Patient Usage Configuration

When receiving patients, you can select loading the nearest configuration or the specified configuration. You can set the default configuration by following these steps:

- Select 【Main Menu】 quick key→from 【Configuration】 column to select
 【New Patient Config】 →input maintenance password→Enter.
- 2. Select [Default] or [Current].
 - ➤ 【Default】: The monitor loads the default configuration specified by the user when it receives the patient, please refer to 5.2 Selecting Default Configuration.
 - > **[Current]**: The monitor loads the nearest configuration when it receives the patient.

5.8. Monitor Boot Usage Configuration

When the monitor starts, you can select whether the monitor loads the nearest configuration or the specified configuration. You can set the default configuration by following these steps:

- Select 【Main Menu】 quick key→from 【Configuration】 column to select
 【Boot Config】 →input maintenance password→Enter.
- 2. Select [Default] or [Current].
 - Configuration.
 The monitor loads the default configuration specified by the user the user when it starts. Please refer to 5.2 Selecting Default Configuration.
 - **Current**: The monitor loads the nearest configuration when it starts.

5.9. Setting password valid time

If you access the configuration management menu and use the password to access alarm-related settings, you can set a valid time for the password, beyond which you will need to re-enter the password.

Please follow the steps below:

- Select 【 Main Menu 】 quick key→from 【 System 】 column to select
 【 Maintenance 】 →input maintenance password→Enter.
- 2. Select [Authorization] submenu.
- 3. Set [Retention Time].

Chapter 6 User Interface

6.1. Interface Style

You can set the style of the interface as needed.

6.1.1. Changing the Screen Layout

You can select the parameters and waveforms you want to view in the **Layout** window. For details, please refer to 3.7.2 Setting Display Screen.

6.1.2. Selecting Screen

The conventional screen is the most commonly used clinical monitoring screen for the monitor, and the monitor enters the normal screen after being turned on. You can also select the screen type as needed, the steps are as follows:

- 1. Enter **[Screen Select]** interface in one of the following ways:
 - > Select 【Screen Setup】 quick key→Select 【Screen Select】 submenu.
 - Select 【Main Menu】 quick key→from 【Display】 column to select 【Screen Select】.
- 2. Select screen types as needed.

6.1.3. Setting Big Font Screen

- 1. Enter **Screen Layout** interface in one of the following ways:
 - > Select **[Screen Setup]** quick key—Select **[Screen Layout]** submenu.
 - ➤ Select 【Main Menu】 quick key→from 【Display】 column to select 【Screen Layout】.
- 2. Select **[Big Font]** submenu.
- 3. Click on each locale to display the parameters you want to display.

6.1.4. Changing Parameter Color

The steps for setting colors of parameter values and waveforms are as follows:

- 1. Enter [Param Color] interface in one of the following ways:
 - > Select 【Screen Setup】 quick key→Select 【Param Color】 submenu.
 - ➤ Select [Main Menu] quick key→from [Parameter] column to select [Param Color].
- Select [Current Select] submenu to set the colors of parameter values and waveforms.
- 3. Select [All] submenu to set the colors of all parameter values and waveforms.

6.2. Dynamic Trend Screen

6.2.1. Entering Dynamic Trend Screen

The Dynamic Trend window is located to the left of the waveform area, showing the trend of a series of parameters in a recent period of time. You can enter the Dynamic Trend screen in any of the following ways:

- ➤ Select [Screen Setup] quick key→Select [Screen Select] submenu→Select [Dynamic Trend].
- Select [Main Menu] quick key→from [Display] column to select [Screen Select] →Select [Dynamic Trend].

In the Dynamic Trend window, the parameter label name of the trend is displayed above each trend curve, and the trend scale is displayed on the left. Trend times are displayed at the bottom of the window.

6.2.2. Setting the Trend Time

Follow these steps to set the trend time:

- 1. Enter the Dynamic Trend window.
- 2. Select the Dynamic Trend area, open **[Dynamic Trend]** menu.
- 3. Select Trend Length.

6.2.3. Exiting Dynamic Trend Screen

You can exit the Dynamic Trend screen by any of the following methods:

➤ Select 【Screen Setup 】 quick key→Select 【Screen Select 】 submenu→Select the screen you need to enter.

➤ Select [Main Menu] quick key—from [Display] column to select [Screen Select] →Select the screen you need to enter.

6.3. OxyCRG screen

The OxyCRG screen graphically displays high-resolution trend curves and compressed waveforms of HR, SpO_2 , and RR.

6.3.1. Entering OxyCRG screen

You can enter the OxyCRG screen by any of the following methods:

- Select [Screen Setup] quick key→Select [Screen Select] submenu→Select
 [OxyCRG View].
- Select [Main Menu] quick key→from [Display] column to select [Screen Select] →Select [OxyCRG View].

The OxyCRG screen shows two trend curves and a compression waveform.

6.3.2. Select Display Parameters and Scales

Follow the steps below to set the parameters of the OxyCRG screen:

- 1. Enter OxyCRG screen.
- 2. Select [Setup].
- 3. Set [Trend 1], [Trend 2], [Compressed Wave] separately.
- 4. Select **[Scale]** submenu, set the scales of each parameter. If you want to use the default scaleplate of the system, select the **[Default Scale]** on the OxyCRG screen.

6.3.3. Setting the Trend Time

Follow the steps below to set the Trend Time:

- Enter OxyCRG screen.
- 2. Select **Zoom**.

6.3.4. Entering OxyCRG Review Screen

You can review the 48-hour trend curve and compression waveform on the OxyCRG

Review Screen. Follow these steps to go to the OxyCRG Review Screen:

- 1. Enter OxyCRG screen.
- 2 Select [Review]

6.4. Other Bed Observation

You can check the alarm status and real-time physiological data of patients on other remote monitors in the LAN on the monitor. A remote monitor (such as a bedside monitor) is also called other bed monitor. You can monitor the alarms of up to 16 other beds at the same time, and you can also view the waveform of 1 other bed from the current monitor.

You can monitor the alarm of other bed through the alarm monitoring area of [Remote View] interface.



■ Can view the monitor through the remote monitor. You can check the alarm and waveform of this monitor from 5 remote monitors at the same time.

6.4.1. Other Bed Screen

Through the **【Other Bed View】** screen, you can check the real-time parameters and waveforms of a remote monitor and monitor the alarm of other monitors.

6.4.1.1. Entering Other Bed Screen

Entering other bed screen by any of the following methods:

- ♦ Select 【Remote View 】 quick key.
- ◆ Select [Screen Setup] quick key→Select [Screen Select] submenu→Select [Other Bed View].
- ◆ Select [Main Menu] quick key→from [Display] column to select [Screen Select] →Select [Other Bed View].

6.4.1.2. Other Bed Interface

Below is other bed observation interface:



- Other bed observation area: Display the patient information, alarm status, information, waveform and parameters of the selected bed. You can move the interface down to browse the contents of the interface.
- 2. Other bed monitoring area
 - > Displays all remote monitors being monitored.
 - Display the equipment number of other bed monitor, and indicate other bed monitor's alarm status with different background colors:
 - Red: Indicates that other bed monitor is giving high priority physiological or technical alarm, and the high priority alarm is the highest level alarm in the current alarm of the bed.
 - Yellow: Indicates that other bed monitor is giving medium priority physiological or technical alarm. The medium priority alarm is the highest level alarm in the current alarm of the bed.
 - Cyan: Indicates that the monitor is giving low priority physiological or technical alarm, and the low priority alarm is the highest level alarm in the current alarm of the bed.
 - · Green: Indicates that the monitor is connected successfully and no

alarms have occurred.

- Gray: Indicates that the monitor is not connected successfully.
- Gray with : ": Indicates that the monitor is disconnected during the connection process.

6.4.1.3. Adding Other Bed

Only add other bed monitor, the device can monitor alarm of other bed. If you have added other bed monitor, you can up to a choice of 16 beds. Add other bed as follows:

- Select other bed observation interface area, in the pop-up window 【Bed View Settings (Bed number)】 to select 【Bed】 submenu.
- 2. Select the device number of the monitor to be observed in the list.
 - The setup interface mainly displays the device number, IP, and patient information of the networked monitor.
 - Select **[Show Offline Bed]** to display the device numbers of all monitors.

6.4.1.4. Delete Bed

If you no longer need to monitor the remote monitor, you can remove it as follows:

- Select other bed observation interface area, in the pop-up window Bed View
 Settings (Bed number) to select Bed submenu.
- 2. Cancel the device number of the monitor in the list. If you want to delete all beds, you can select **[Delete All]**.

6.4.1.5. Display Main Bed

In the other bed monitoring area of its bed observation window, select a bed and then other bed observation window will display the real-time monitoring interface of the monitor. This selected bed is called the main bed.

6.4.1.6. Alarm information Display

You can follow these steps to view the current real-time alarm information of the main bed:

1. Enter [Alarm] interface by one of the following methods:

- Click on the alarm information display area of the bed observation area, and the alarm interface will pop up.
- Select other bed observation interface area, in the pop-up window
 [Bed View Settings (Bed number)] to select [Alarm] submenu.
- 2. In **[Alarm]** submenu to check the current physiological and technical alarm information of the main bed.

6.4.1.7. Reset Other Bed Alarm

In the **【Bed View Settings (Bed number)** window—select **【Reset Remote Alarm** which in the **【Alarm** submenu, the alarm of the corresponding remote monitor (main bed) is reset. Only when this function is turned on can you reset other bed alarm. For detailed, please refer to 7.13.1 Other Bed Alarm Reset.

6.4.1.8. Selecting Waveform

Other bed observation area can display up to 4 waveforms. Following these steps to select the label name of the waveform you want to observe:

- Select other bed observation interface area, in the pop-up window Bed View
 Settings (Bed number) to select Wave submenu.
- 2. In proper order to select [First Wave], [Second Wave], [Third Wave] and [Fourth Wave], then select the label name of the waveform in the pop-up list. If you select is [Close], then the display of one waveform will be turned off.

6.4.1.9. Selecting Parameters

Other bed observation can display all the online parameters. Select the parameter label names you want to observe as follows:

- Select other bed observation interface area, in the pop-up window [Bed View Settings (Bed number)] to select [Param] submenu.
- 2. Open the parameter labels you want to observe in the displayed list of online parameters.



■ The data displayed will delay in other bed observation window. Don't rely on other bed observation window for real-time data.

6.5. Big Font Screen

You can enter big numerics screen in either of the following ways:

- ➢ Select [Screen Setup] quick key→Select [Screen Select] submenu→Select
 [Big Font Screen].
- Select [Main Menu] quick key→from [Display] column to select [Screen
 Select] →Select [Big Font Screen].

In the setting window of **[Big Font Screen]**, you can select 6 parameters to observe according to your needs. For parameters with waveform, one waveform is displayed at the same time.

6.6. Freezing Waveforms

During monitoring the patient, you can freeze the waveform on the screen and then review it to carefully observe the patient during this time. You can also export the frozen waveform through the recorder.

6.6.1. Entering Freezing Status

Under the non-freezing condition, select [Freeze] quick key, and then pop-up [Freeze] menu.



2. All waveforms are frozen, that is, the waveforms are not refreshed. The data in the parameter area is refreshed normally.

6.6.2. Waveform Review

On the freezing waveforms screen you can operate the following:

In the frozen status, you can select the control icon to browse the frozen waveforms: the frozen waveform will move to the left or right correspondingly. At the same time, each waveform is marked with a time scale, and the freezing time is recorded as [0s]. As the waveform moves to the right, the time scale will be gradually changed to

[-1s], [-2s], [-3s]......

Icon	Function
<<	Up to the fist page
<<	Up to the previous page
<	Up to the previous second
>	Down to the next second
>>	Down to the next page
>>	Down to the last page

You can set the speed of the frozen waveform as needed.

6.6.3. Releasing Freezing

Under freezing condition, you can select button X in the upper right corner of the freezing menu to release the freezing condition.

6.6.4. Recording Freezing Waveforms

Select the **【Record】** button in **【Freeze】**, the recorder will output the waveform selected and the parameter value at the Freezing time. The recorder can output up to 3 waveforms at one time. For the setting of frozen waveforms, please refer t to **26.6.1 Selecting** the recorded waveforms.

6.7. SpO₂ Screen

For neonatal patients, if you only care about the patient's SpO₂ and PR, you can use the SpO₂ Screen. The SpO₂ Screen uses large fonts to display the data of SpO₂ related parameters, and real-time body temperature and NIBP values.



■ The SpO₂ Screen is only applicable to neonates.

6.7.1. Entering the SpO₂ Screen

Choose any of the following methods to enter the SpO₂ Screen:

- Use two fingers to slide the touch screen left and right to switch to the SpO₂ Screen.
- Select [Screen Setup] quick key → select [Screen Select] submenu → select [SpO₂ Screen].
- Select 【Main Menu】 quick key→from 【Display】 column to select 【Screen Select】 → select 【SpO₂ Screen】.

6.7.2. SpO₂ Screen Display

The figure below is a schematic diagram of the SpO_2 Screen. The graphics displayed on your monitor may be slightly different.



- 1. Trend Table: Display the trend of SpO₂, PR, PI.
- 2. SpO₂ statistics area: Display SpO₂ segment statistics data.
- 3. NIBP parameter area: Display the measurement value and alarm limit of NIBP.
- 4. Temp parameter area: Display the measurement value and alarm limit of temp.
- 5. SpO₂ parameter area: Display the measurement value and alarm limit of SpO₂, PR

and PI. The dashboard displays information about the alarm limit. \triangle pointer indicates the current measurement value.

6. Pleth waveform

6.7.3. SpO₂ Screen Operation

You can enter the parameter setting menu through the SpO₂ Screen. The operation method is as follows:

- Click SpO₂ statistical histogram to enter 【SpO₂ Statistics】 setup menu, set SpO₂ segment result values and target segment.
- Click the measurement values (SpO₂ and PI), the dashboard or Pleth waveform to enter 【SpO₂】 setup menu.
- Click the measurement or the dashboard of PR to enter 【PR】 setup menu.
- Click the measurement of Temp to enter the 【TEMP】 setup menu.
- Click the measurement of NIBP to enter the 【NIBP】 setup menu.

6.7.4. Setting

You can select different SpO_2 module and Temp module, set the interval of trend data and SpO_2 statistics time.

6.7.4.1. Selecting SpO₂ parameter

When monitoring the patient using two SpO₂, you can set the SpO₂ parameters you want to display in the following ways:

- 1. Select o to enter the SpO₂ Screen Setting menu.
- 2. Select **[Setup]** menu.
- 3. Set the ${ { { [\![SpO_2]\!] }}}$ to ${ { { { [\![SpO_2]\!] }}}}$ or ${ { { { { [\![SpO_2]\!] }}}}}$.

6.7.4.2. Selecting Temp parameter

When monitoring the patient using multiple temp modules, you can set the Temp parameters you want to display in the following ways:

1. Select **o** to enter the **SpO₂ Screen Setting** menu.

- 2. Select [Setup] menu.
- 3. Set the **[Temp]**.

6.7.4.3. Setting the interval of trend data

You can set the interval of trend data by following ways:

- 1. Select **o** to enter the **SpO₂ Screen Setting** menu.
- 2. Select [Setup] menu.
- 3. Set the **[Trend Interval]**.

6.7.4.4. Setting the SpO₂ statistics duration

You can set the SpO₂ statistics time by following ways:

- 1. Select o to enter the SpO₂ Screen Setting menu.
- 2. Select [Setup] menu.
- 3. Set the **[Statistics Duration]**.

Chapter 7 Alarm

7.1. Introduction

This chapter introduces the alarm function and the settings of the monitor.

7.2. Safety Information



- A potential hazard can exist if different alarm presets are used for the same or similar equipment in any single area, e.g. an intensive care unit or cardiac operating room.
- Alarm Settings for different monitors in the same area may vary to suit the condition of the patient being monitored. Before starting the patient monitoring, check whether the alarm setting is suitable for the patient, and always open certain necessary alarm limits, and ensure that the alarm limit setting is suitable for the patient.
- Setting the alarm limit beyond the measurement range or to the limit value may invalidate the alarm system. For example, high oxygen level can make premature infants infect crystalline post fibroplasias. If the SpO₂ alarm high limit is set at 100%, it is equivalent to disconnecting the upper limit alarm.
- When the alarm sound is turned off, even if a new alarm is triggered, the monitor will not emit an alarm sound. Therefore, the user must carefully select whether to turn off the alarm sound. Check patient status frequently after turning off alarm or alarm sound.
- For patients who cannot be continuously treated by medical staff, the alarm settings must be made according to the patient's condition.
- Do not rely solely on an audible alarm system to monitor a patient. There may be risks in adjusting the alarm sound to a lower volume, which may impede operator recognition of alarm. The alarm volume should be large enough in the current monitoring environment and the actual clinical condition of the patient should be paid close attention.



NOTE:

■ When the alarm system is powered off, the monitor will save the alarm information before power interruption, and the stored alarm information will not change.

7.3. About the alarm

7.3.1. Alarm Categories

The monitor has two different types of alarms: physiological alarms and technical alarm.

- Physiological alarms: Physiological alarms are triggered by a monitored parameter value that violates set alarm limits or an abnormal patient condition.
- > Technical alarms: Technical alarms are triggered by a device malfunction due to improper operation or system problems. The problems may result in system abnormal operation or irresponsible monitoring parameters.

Apart from the physiological and technical alarm messages, the monitor will also display some information related to system status or patient status.

7.3.2. Alarm Priority

By severity, the patient monitor's alarms can be classified into three categories:

- ➤ High priority: Indicate that the patient is in a life threatening situation or a severe device malfunction, and an emergency treatment is necessary.
- Medium priority: Indicate that your patient's vital signs appear abnormal, a severe device malfunction or an improper operation, and an immediate treatment is required.
- Low priority: Indicate that the patient's vital signs appear abnormal, a severe device malfunction or an improper operation, the user needs to know the current situation.
- Prompt: Prompt patient and system status information.

7.3.3. Alarm Indicators

When an alarm occurs, the patient monitor will indicate it to the user through visual or audible alarm indications:

Alarm signa	l	High priority alarm	Medium priority alarm	Low priority alarm	Prompt	Note
Alarm Lamp		The lamp quickly flashes red with 1.4Hz ~2.8Hz, Duty cycle 20%-60%.	The lamp slowly flashes yellow with 0.4Hz~ 0.8Hz, Duty cycle 20-60%.	The lamp turns cyan without flashing, Duty cycle 100%.	/	/
Alarm Tone Mode	ISO	DO-DO-DO-D O-DO DO-DO-DO DO-DO	DO-DO-DO -	DO-	/	/
Alarm Inform	nation	White words, Red background	Black words, Yellow background	Black words, Cyan background	White words	Displayed in the top information area, click on the alarm information to view the alarm information list.
Alarm level s	symbol	***	**	*	/	Displayed in front of alarm information.
Parameter ala	nrm	Red background, flashes	Yellow background, flashes	Cyan background, flashes	/	/



■ When multiple alarms of different priorities occur simultaneously, the alarm lamp and alarm tone are prompted according to the highest level of all current alarms.

■ When there are multiple alarms in the same area at the same time, the alarm messages are displayed circularly on the screen.

7.3.4. Alarm Status Symbols

In addition to the alarm methods described in section *Alarm Indicators*, the following alarm icons will appear on the screen to indicate different alarm states:



Indicates an alarm for a parameter is off or the alarm system is off.



Indicates all the alarms are paused.



Indicates the alarm sound is off.



Indicates alarms are reset.

7.4. Viewing physiological alarms list

The steps of viewing physiological alarms are as follows:

- 1. Select physiological alarms area to enter **[Alarm Informations]** window.
- 2. Select [Phy. Alarm] submenu.

7.5. Viewing technical alarms list

The steps of viewing technical alarms list are as follows:

- 1. Select technical alarms area to enter [Alarm Informations] window.
- 2. Select [Tec. Alarm] submenu.

7.6. Viewing the graphical display of technical alarms (Alarm View)

In the technical alarms list, if "---" is prompted behind the alarm information, the alarm contains help information of pictures to help you identify the cause of the alarm. The steps of viewing the help information of alarms are as follows:

- 1. Select technical alarms area to enter [Alarm Informations] window.
- 2. Select **Tec. Alarm** submenu.
- 3. Select the alarm you need to view in the alarms list.

7.7. Setting Alarm

You can set the alarm properties centrally. Select [Alarm Setup] quick key or select the corresponding button from the [Alarm] column, in the main menu to set alarm.

7.7.1. Setting Parameter Alarm

The steps to centrally set the properties of the parameter alarm are as follows:

- 1. Enter **[Limit]** interface in any of the following ways:
 - > Select [Alarm Setup] quick key.
 - > Select [Main Menu] quick key→from[Alarm]column to select[Limit].
- 2. Select parameter submenu to set the alarm according to the required. You can also set the alarm for individual parameters from the parameters menu.

7.7.2. Changing Alarm Setup Protection Mode

You can change the password protection mode of the alarm settings and arrhythmia settings as follows:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】→input maintenance password→Enter.
- Select [Authorization] submenu.
- 3. Change the password protection mode of the alarm settings.
 - No Password : Change alarm setups to not be password protected.
 - Password : Change alarm switch, alarm limit and alarm level to be protected by password.

If you use password to access alarm and arrhythmia alarm related settings, you can set the valid time of the password, beyond which you need to re-enter the password. For details, please refer to 5.9 Setting Password Valid Time.

7.7.3. Setting Alarm Sound Properties

7.7.3.1. Setting Alarm Volume

1. Enter **[Setup]** interface in either of the following ways:

- > Select 【Alarm Setup】 quick key→Select 【Setup】 submenu.
- > Select [Main Menu] quick key→from [Alarm] column to select [Setup].
- 2. Set 【Alarm Volume】. The alarm volume range is X-10, in which X is the minimum volume, depending on the set minimum alarm volume, and 10 is the maximum volume.
- 3. Set [High Alarm Volume].
- 4. Set [Reminder Volume].



NOTE:

- When the alarm volume is set to 0, the alarm tone will be turned off, and an alarm audio off icon will appear on the screen.
- When the alarm volume is set to 0, the setting of high level alarm volume is invalid.

7.7.3.2. Setting the minimum alarm volume

The minimum alarm volume determines the minimum alarm volume setting. The steps are as follows:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】→input maintenance password→Enter.
- 2. Select 【Alarm】 submenu→ 【Sound】 submenu.
- 3. Select [Minimum Alarm Volume].



NOTE:

- You can set the minimum alarm volume to 0 only when you are connected to the CMS. If the monitor is not connected to the CMS, the minimum alarm volume can only be set to 1.
- When the CMS is connected, if the minimum alarm volume is set to 0, the minimum alarm volume will be automatically changed to 2 when the CMS is disconnected.

7.7.3.3. Setting Alarm Tone Mode

The setting steps are as follows:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】→input maintenance password→Enter.
- Select 【Alarm】 submenu→ 【Sound】 submenu→ 【Alarm Sound】, you can select 【ISO】.

7.7.3.4. Setting Alarm Tone Interval

You can set the alarm tone interval. The steps are as follows:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】 →input maintenance password→Enter.
- 2. Select 【Alarm】 submenu→ 【Sound】 submenu.
- 3. Set [High Alarm Interval], [Med Alarm Interval] and [Low Alarm Interval]
 - \blacktriangleright **[High Alarm Interval]**: 3~15s, and the default value is 10s.
 - \triangleright [Medium Alarm Interval]: 3~30s, and the default value is 20s.
 - Low Alarm Interval $16 \sim 30$ s, and the default value is 20s.

7.7.3.5. Setting Reminder Volume

When the alarm volume is 0, alarm reset or the alarm is off, the monitor can provide periodic alarm prompt tone to remind you that there is still an activated alarm in the current system. This function is turned on by default.

You can set the alarm tone as follows:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】 →input maintenance password→Enter.
- 2. Select 【Alarm】 button→ 【Pause /Reset】 submenu.
- 3. Set [Alarm Pause duration]. You can set [Alarm Pause duration] to [1min], [2min], [3min] or [Permanent], the default is [2min].
- 4. Set [Alarm Off Reminder] switch.
 - > **(On)**: The monitor provides an alarm tone according to the set interval.

- > **(Off)**: The monitor does not provide an alarm tone.
- Set [Reminder Interval]. You can set [Reminder Interval] to [1min],
 [2min], [3min], [5min] or [10min], the default is [5min].

7.7.3.6. Setting Alarm Tone Enhancement

The monitor provides an alarm tone enhancement function. If the alarm exceeds the set time and is not confirmed, the alarm volume can be automatically enhanced.

The steps to set the alarm tone enhancement are as follows

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】 →input maintenance password→Enter.
- 2. Select 【Alarm】 → 【Sound 】.
- 3. Set [Auto Increase Volume] to [3 Steps], [2 Steps], [1 Steps] or [Off].
 - ➤ 【3 Steps】: After the alarm occurs, the alarm volume will be automatically increased to level 3, if the set time is not confirmed.
 - > **[2 Steps]**: After the alarm occurs, the alarm volume will be automatically increased to level 2, if the set time is not confirmed.
 - > **[** 1 Steps]: After the alarm occurs, the alarm volume will be automatically increased to level 1, if the set time is not confirmed.
 - > **(Off)** After the alarm occurs, the set time is not confirmed, and the alarm volume remains unchanged.
- 4. Set [Increase Volume Delay], select sound enhanced delay time.

7.7.4. Setting Alarm Delay Time

For the over-limit alarm of continuous measurement parameters, the alarm delay time can be set. If the condition of triggering alarm disappears within the delay time, the monitor will not alarm.

Set the delay time for the alarm by following these steps:

Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】 →input maintenance password→Enter.

- 2. Select 【Alarm】 submenu→ 【Other】 submenu.
- 3. Set [Alarm Delay].

The delay time of the apnea alarm is not affected by the alarm delay time setting. You can set the delay time of the apnea alarm separately.

7.7.4.1. Setting Apnea Alarm Delay Time

Steps to set apnea alarm delay time are as follows:

- 1. Enter **[Setup]** interface in either of the following ways:
 - > Select 【Alarm Setup】 quick key→Select 【Setup】 submenu.
 - ➤ Select [Main Menu] quick key→from [Alarm] column to select [Setup].
- 2. Select [Apnea Delay] to set apnea alarm delay time.

7.7.5. Setting Alarm Waveform Length

You can set the length of the waveform needs to be output when an alarm occurs, the setting steps are as follows:

- 1. Enter **[Setup]** interface in either of the following ways:
 - > Select 【Alarm Setup】 quick key→Select 【Setup】 submenu.
 - ➤ Select [Main Menu] quick key→from [Alarm] column to select [Setup].
 - ➤ Select [Main Menu] quick key→from [Report] column to select [Record Setup].
- 2. Set [Alarm Record Duration].

7.7.6. Setting CMS and eGateway Disconnect Alarm Switch

You can set whether to alarm when the monitor and the CMS or eGateway are disconnected. This function is enabled by default. The setting method is as follows:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】→input maintenance password→Enter.
- 2. Select 【Alarm】 submenu→【Other】 submenu.

3. Open or Close 【CMS/eGW Disconnected】.

When the 【CMS/eGW Disconnected】 switch is turned on, the technical alarm will be generated when the monitor and the CMS/eGateway are disconnected after successful connection.

7.8. Alarm Pause

When the alarm is paused, it has the following characteristics:

- > Shield all physiological alarms within the set time.
- > The technical alarm sound is paused, but the alarm light and alarm information are still displayed.
- Display the remaining time of alarm paused in the physiological alarm information area.
- Display the alarm paused icon in the information area.

After reaching the alarm pause time, the monitor will automatically exit the alarm pause state. You can also click **[Alarm Pause]** quick key to manually cancel the alarm pause.

7.8.1. Setting Alarm Pause Time

Alarm pause time can set to: [1min], [2min], [3min] and [Permanent], the default is 2 minutes. The steps to set the alarm pause time are as follows:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】→input maintenance password→Enter.
- 2. Select [Alarm] submenu→ [Pause /Reset] submenu.
- 3. Set [Alarm Pause Duration] .

7.8.2. Turn off all the Alarm

If **【Alarm Pause Duration 】** is set to **【Permanent】** (Refer to section 7.8.1 Setting Alarm Pause Time), you can press **【Alarm Pause 】** quick key to turn off all alarms. When the alarm is turned off, it has the following characteristics:

- No physiological alarm lamps flash and no physiological alarms are sounded.
- > The technical alarm sound is turned off, but the alarm light and alarm information are still displayed

- Display "Alarm Off" in the physiological alarm information area and the background color is red.
- Display alarm off icon in status area.

To exit the alarm off state, click [Alarm Pause] quick key again.



WARNING:

Pausing or turning off the alarm may cause the patient to be in danger, please handle it carefully.

7.9. Alarm Reset

Click on [Alarm Reset] quick key to reset the alarm system, and the alarm reset icon will appear in the system status information area.



NOTE:

■ In the alarm reset state, if a new alarm is generated, the alarm reset icon disappears and the alarm system is reactivated.

7.9.1. Physiological Alarm Reset

After the physiological alarm is reset, the sound of the currently existing physiological alarm is shielded, and the other alarm states remain unchanged.

7.9.2. Technical Alarm Reset

When the technical alarm is reset, it has the following characteristics:

- > The technical alarm that can be completely cleared is cleared. The monitor will not have any alarm indication for the cleared technical alarm.
- > Technical alarm that can clear sound and light is displayed as prompt message.
- The sound of the technical alarm that cannot be cleared is shielded. For the indication of the technical alarm after the alarm is reset, please refer to **D.2**Technical Alarm.

7.10. Latching Alarms

The physiological alarms are classified into "Latching" and "Non-latching".

- Non-latching alarms: After the condition that triggered the alarm of a parameter disappears, the system will not make any prompt for this alarm of this parameter.
- Latching alarms: Even if the condition that caused the physiological alarm disappears, the alarm signal will still be "Latched", and the time of the last triggering of the alarm will be displayed behind the alarm information in the information area.
- You can choose to individually lock the visual signal or simultaneously lock the visual and audible signals.
- For visual latching, after the alarm condition disappears, the visual signal of the alarm, including the alarm light, the alarm information and the background color remain unchanged, and the alarm information text is followed by the time of last triggering the alarm.
- For audible latching, the system still emits an alarm tone after the alarm condition disappears.

The steps to latch the physiological alarm are as follows:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】 →input maintenance password→Enter.
- 2. Select 【Alarm】 submenu→ 【Latching】 submenu.
- 3. Select how you want to latch the alarms. Alarm latching rules are as follows:
 - You can separately select visual latching.
 - Latching audible alarm signal simultaneously latches visual signal corresponding to the alarm level.
 - When a low priority alarm is latched, the high priority alarm is also automatically locked. For example, if you select the low priority alarm, the medium priority alarm and the high priority alarm will also be latched simultaneously.



CAUTION:

- Changing of alarm priority may affect the latching status of corresponding alarm. Please determine if you need to reset the latching status for the specific alarm when you have changed its alarm priority.
- When the alarm system is reset, the latched physiological alarms are cleared.
- Do not set all alarm status to latching alarm signals when used in the intensive care unit.

7.11. Nurse Call

The nurse call function means that when the alarm set by the user occurs, the monitor can output a signal to the nurse call system, call the nurse. The monitor provides a nurse call connector, and the monitor is connected to the nurse call system of the hospital through the randomly provided nurse call cable. After the system is connected, the connector can implement the nurse call function.

The nurse call function must be valid only if the following conditions are met:

- ◆ The nurse call function is turned on.
- ♦ A user-defined alarm occurs.
- ◆ The monitor is not alarm paused or off.

7.11.1. Changing Nurse Call Settings

To set the type and priority of alarms that are sent to the nurse call system, follow this procedure:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】→input maintenance password→Enter.
- 2. Select 【Alarm】 submenu→ 【Nurse Call】 submenu.
- 3. Select **[Signal Type]** to set the type of nurse call signal.
 - ◆ 【Pulse】: The nurse call signal is a pulse signal and each pulse lasts one second. When multiple alarms simultaneously occur, only one pulse signal is outputted. If an alarm occurs but the previous one is not cleared, a new pulse signal will also be outputted.

- ◆ 【Continuous】: The nurse call signal lasts until the alarm ends. That is to say the duration of a nurse call signal is equal to that of the alarm condition.
- 4. Select **[Trigger Type]** to set the work mode of the nurse call relay.
- Select 【Alarm Priority】 to set the priority of alarms sent to the nurse call system.
- 6. Select **【Alarm Type 】** to set the type of alarms sent to the nurse call system.



WARNING:

■ Do not rely exclusively on the nurse call system for alarm notification.

Remember that the most reliable alarm notification combines audible and visual alarm indications with the patient's clinical condition.

7.12. Intubation Status

The monitor provides the intubation status function during RESP, CO₂, AG and RM monitoring. In this state, the physiological alarms related to RESP, CO₂, AG and RM are shielded, and the alarm off icon is displayed in the parameter area. During the intubation process of general anesthesia surgery, the intubation status can be selected to shield unnecessary alarms.

7.12.1. Entering Intubation Status

To enter the intubation status, choose either of the following ways:

- ◆ Select 【Intubation Status】 quick key.
- ◆ From the bottom of the 【RESP】, 【CO₂】, 【AG】 or 【RM】 menu to select 【Intubation Status】 button.
- ◆ Select [Main Menu] quick key→from [Alarm] column to select [Intubation Status].

7.12.2. Setting Intubation Status Time

The default intubation time is 2 minutes. To change the time, following this procedure:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】→input maintenance password→Enter.
- 2. Select 【Alarm】 submenu→【Other】 submenu.
- 3. Set [Intubation Duration].

7.12.3. Exiting the Intubation Status

To exit the intubation status, choose either of the following ways:

- > Select [Intubation Status] quick key.
- From the bottom of the [RESP], [CO₂], [AG] or [RM] menu to select [Exit Intubation Status] button.
- ➤ Select [Main Menu] quick key→from [Alarm] column to select [Exit Intubation Status].

7.13. Other Bed Alarm

Enter other bed observation interface, and when the monitored bed monitor has an alarm triggered, the alarm light and alarm sound are prompted according to the highest level of all alarms of the current monitor and other bed monitor. You can view and manage other bed alarm. The alarm delay time from the device to other bed is ≤2s.

7.13.1. Other Bed Alarm Reset

You can reset other bed alarm on the monitor. The steps to enable it to reset the bed alarm are as follows:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】→input maintenance password→Enter.
- 2. Select 【Alarm】 submenu→ 【Remote View】 submenu.
- 3. Open [Reset Remote Bed's Alarms].

And then 【Bed View Settings (bed number)】 window→ 【Reset Remote Alarm】

button in the 【Alarm】 submenu will be activated. Click on 【Reset Remote Alarm】 button, other bed alarm will be reset.



 Only when the "Alarm Reset By Other Bed" function of the remote monitor is enabled, can you reset other bed alarm on this monitor

7.13.2. Authorizing the Alarm Reset to Other Devices

Alarms on your monitor can be reset by remote devices if you enable this function. To do so, follow this procedure:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】→input maintenance password→Enter.
- 2. Select 【Alarm】 submenu→ 【Remote View】 submenu.
- 3. Open [Alarm Reset By Other Bed] switch.

7.13.3. Switching Off the Remote Device Disconnection Alarm

The monitor can provide an alarm if remote devices are disconnected. By default, the function is enabled. To disable the alarm, follow this procedure:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】→input maintenance password→Enter.
- 2. Select 【Alarm】 submenu→ 【Remote View】 submenu.
- 3. Switch off [Remote Disconnected Alarm].

7.14. Detecting Alarm

The monitor automatically performs a self-test at startup. Check that the alarm lamp illuminates, one after the other, in red, yellow, and cyan, and that an alarm tone is heard. This indicates that the visible and audible alarm indicators functions correctly.

To further test individual measurement alarms, perform measurements on yourself or using a simulator. Adjust alarm limits and check that appropriate alarm behavior is observed

7.15. Actions When an Alarm Occurs

When an alarm occurs, please refer to the following steps to take proper actions:

- 1. Check the patient's condition.
- 2. Confirm the alarming parameter or alarm category.
- 3. Identify the source of the alarm.
- 4. Take proper action to eliminate the alarm condition.
- 5. Check if the alarm is eliminated.

For more information, please refer to *D Alarm Message*.

Chapter 8 ECG

8.1. Introduction

The electrocardiogram (ECG) measures the electrical activity of the heart and displays it on the monitor as waveforms and parameters. The monitor provides 3-lead, 5-lead, 6-lead, and 12-lead ECG monitoring, arrhythmia analysis, ST segment analysis and OT/OTc measurements.

8.2. Safety Information



WARNING:

- This equipment is not intended for direct cardiac application.
- Make sure the conductive parts of electrodes and associated connectors for applied parts, including the neutral electrode, should not contact any other conductive parts including earth.
- Use defibrillation-proof ECG cables during defibrillation.
- Do not touch the patient or metal devices connected to the patient during defibrillation.
- To reduce the hazard of burns during high-frequency surgical procedure, ensure that the monitor's cables and transducers never come into contact with the electrosurgery unit (ESU).
- To reduce the hazard of burns during use of high-frequency surgical unit (ESU), the ECG electrodes should not be located between the surgical site and the ESU return electrode.



CAUTION:

- Only use parts and accessories specified in this manual. Follow the instructions for use and adhere to all warnings and cautions.
- Regularly inspect the electrode application site to ensure skin quality. If there are signals of allergies, replace the electrodes or change the application site.

■ Interference from a non-grounded instrument near the patient and electrosurgery interference can cause problems with the waveform.



NOTE:

■ Due to the asynchrony of ECG signal sampling characteristics and sampling rate, the digital system will produce a perceptible modulation effect from one cycle to the next, especially when the electrocardiogram is measured by children.

8.3. ECG Display

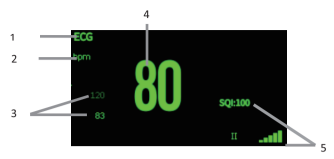
The following figures show the ECG waveform and parameter areas. Your display may be configured to look slightly different.

♦ Waveform Display



- (1). ECG lead label
- (2). ECG waveform gain
- (3). ECG filter mode
- (4). ECG waveform speed
- (5). Paced status: If **[Paced]** is set to **[Yes]**, **[Second Paced]** is set to **[No]**, **[Second Paced]** is displayed.
- (6). Notch frequency
- (7). Alarm message: Display only the highest level of alarm information.
- (8). Pace pulse mark: If **[Paced]** is set to **[Yes]**, the pace pulse markers are shown on each ECG waveform when the patient has a paced signal.

♦ Parameter Display



- (1) Parameter label
- (2) HR unit
- (3) HR alarm limit: If the HR alarm is turned off, the alarm close icon is displayed here.
- (4) HR value
- (5) ECG signal quality index: Indicates the signal quality of the primary calculation lead.



NOTE:

The ECG parameter area and waveform area are configured to be different for different lead type and ECG settings.

8.4. Preparing for ECG Monitoring

8.4.1. Preparing the Patient Skin

Proper skin preparation is necessary for good signal quality at the electrode sites, as the skin is a poor conductor of electricity. To properly prepare the skin, choose flat areas and then follow this procedure:

- 1. Shave hair from skin at chosen electrode sites.
- 2. Gently rub skin surface at sites to remove dead skin cells.
- 3. Thoroughly cleanse the site with a mild soap and water solution.
- 4. Dry the skin completely before applying electrodes.

8.4.2. Applying Electrodes

To connect ECG cables, follow this procedure:

- Check that electrode packages are intact and not expired. Make sure the electrode gel is moist. If you are using snap electrodes, attach the snaps to the electrodes before placing electrodes.
- 2. Place the electrodes on the prepared sites. Make sure that all electrodes have good skin contact.
- 3. Connect the leadwires to the patient cable.
- 4. Plug the patient cable into the ECG connector.



NOTE:

- Store the electrodes in room temperature.
- Open the electrode package immediately prior to use.
- Never mix patient electrode types or brands. This may lead to problem due to impedance difference.
- When applying the electrodes, avoid bones close to skin, obvious layers of fat, and major muscles. Muscle movement can result in electrical interference.

8.4.3. Lead Wire Color Code

The following table lists the 5-lead labels and colors for AHA and IEC standards:

Lead	IEC		AHA	
Lead	Label	Color	Label	Color
Right arm	R	Red	RA	White
Left arm	L	Yellow	LA	Black
Right leg (neutral)	N/RF	Black	RL	Green
Left leg	F	Green	LL	Red
Chest	С	White	V	Brown

The following table lists the 6-lead labels and colors for AHA and IEC standards:

Lead	IEC		AHA	
Lead	Label	Color	Label	Color
Right arm	R	Red	RA	White

Lead	IEC			AHA	
Leau	Label	Color	Label	Color	
Left arm	L	Yellow	LA	Black	
Right leg (neutral)	N/RF	Black	RL	Green	
Left leg	F	Green	LL	Red	
Chest 1	Ca	White	Va	Brown	
Chest 2	Cb	White	Vb	Brown	

The following table lists the 12-lead labels and colors for AHA and IEC standards:

Lead	IEC		AHA	
Leau	Label	Color	Label	Color
Right arm	R	Red	RA	White
Left arm	L	Yellow	LA	Black
Right leg (neutral)	N/RF	Black	RL	Green
Left leg	F	Green	LL	Red
Chest1	C1	White/Red	V1	Brown/ Red
Chest 2	C2	White/Yellow	V2	Brown/Yellow
Chest 3	C3	White/Green	V3	Brown/Green
Chest 4	C4	White/Brown	V4	Brown/Blue
Chest 5	C5	White/Black	V5	Brown/ Orange
Chest 6	C6	White/ Purple	V6	Brown/ Purple

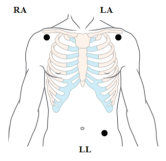
8.4.4. ECG Electrode Placements

In this section, we adopt the AHA standard to illustrate electrode placement.

8.4.4.1. 3-lead Electrode Placement

Taking the AHA standard as an example, the 3-lead electrode placement position is as shown:

- RA placement: directly below the clavicle and near the right shoulder.
- ◆ LA placement: directly below the clavicle and near the left shoulder.
- ◆ LL placement: on the left lower abdomen.

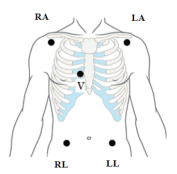


8.4.4.2. 5-lead and 6-lead Electrode Placement

Taking the AHA standard as an example, the 5-lead electrode placement position is as shown:

- RA placement: directly below the clavicle and near the right shoulder.
- ◆ LA placement: directly below the clavicle and near the left shoulder.
- RL placement: on the right lower abdomen.
- ◆ LL placement: on the left lower abdomen.
- ◆ V placement: on the chest.

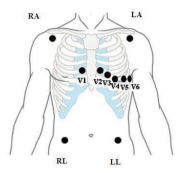
For 6-lead placement, you can use the position for the 5-lead placement, but with two chest leads. The two chest leads (Va and Vb) can be positioned at any two of the V1 to V6 positions. For more information, please refer to *8.4.4.3 12-lead Electrode Placement*.



8.4.4.3. 12-lead Electrode Placement

The electrode placement position of 12-lead includes limbs and chest. The limb

electrodes should be placed on the soft skin. The standard electrode placement position is shown below:



8.4.4.4. Electrode Placement for Surgical Patients

While placing electrodes for a surgical patient, the type of surgery should be considered, for instance, as to a chest surgery, the chest lead electrodes can be placed at sides or backside of chest. Moreover, while using a surgical electrotome, in order to reduce the influence of artifacts to ECG waveform, the electrodes can be placed at left and right shoulders, close to left and right sides of abdomen; the chest lead electrodes can be placed at left side of chest midst. Do not place the electrodes on the upper arm. Otherwise, the ECG waveform will be very small.



WARNING:

- To reduce the hazard of burns during use of electrosurgical units (ESU), the ECG electrodes should not be located between the surgical site and the ESU negative electrode plate.
- Never entangle the ESU cable and the ECG cable together.
- When using ESU, never place ECG electrodes near to the negative electrode plate of the ESU, as this can cause a lot of interference on the ECG signal.

8.4.5. Selecting ECG Lead Type

To select ECG lead type, follow this procedure:

- 1. Select ECG parameter area or waveform area to enter the **【ECG】** menu.
- 2. Select **[Setup]** submenu.
- 3. Set **[Lead Type]** according to the lead type you are going to use.
 - Lead Type is set as [Auto], the monitor automatically detects the lead type.

8.4.6. Checking Paced Status

It is important to correctly set the paced status before you start monitoring ECG. The paced symbol is displayed when **[Paced]** is set to **[Yes]**. The pace pulse markers " | " are shown on each ECG waveform when the patient has a paced signal. If **[Paced]** is set to **[No]** or the patient's paced status is not selected, the symbol will be shown in the ECG waveform area.

To change the paced status, follow this procedure:

- 1. Select ECG parameter area or waveform area to enter the **[ECG]** menu.
- Select [Paced] submenu.
- 3. Set **[Paced]** to be **[Yes]** or **[No]**. If you did not set the paced status, the monitor issues a prompt tone when pace pulse is detected. At the same time, the paced symbol **[Masses]** flashes and the message prompt "Suspected Pacing Signal". Check and set the patient's paced status.



WARNING:

- For paced patients, you must set [Paced] to [Yes]. If it is incorrectly set to [No], the monitor could mistake a pace pulse for a QRS complex and fail to alarm when the ECG signal is too weak. On ventricular paced patients, episodes of ventricular tachycardia may not always be detected. Do not rely entirely upon the system's automated arrhythmia detection algorithm.
- False low heart rate or false asystole alarms may result with certain pacemakers because of pacemaker artifacts, such as electrical overshoot of the pacemaker overlapping the true QRS complexes.
- Do not rely entirely on rate meter alarms when monitoring patients with

pacemakers. Always keep these patients under close surveillance.

- The auto pacer recognition function is not applicable to pediatric and neonatal patients.
- For non-paced patients, you must set 【Paced】 to 【No】.

8.4.7. Enabling Pacer Rejection

The pace pulse rejection function is disabled by default. To enable this function, follow this procedure:

- 1. Select ECG parameter area or waveform area to enter the **【ECG】** menu.
- 2. Select **[Paced]** submenu.
- 3. Switch on [Pacer Reject].



NOTE:

- When pace pulses are detected, the pace pulse marks "|" are shown on the ECG waveforms. [Pacer Reject] setting has no impact on the display of pace pulse marks "|".
- You can switch on 【Pacer Reject】 only when 【Paced】 is set to 【Yes】.

8.5. ECG Settings

8.5.1. Selecting ECG Screen

When monitoring ECG, you can choose the screen as desired.

- For 3-lead ECG monitoring, only normal screen is available.
- ◆ For 5-lead ECG monitoring, besides the normal screen, it can be selected to display 7 waveforms.
- ◆ For 6-lead ECG monitoring, besides the normal screen, it can be selected to display 8 waveforms.
- ◆ For 12-lead ECG monitoring, besides the normal screen, it can be selected to display 12 waveforms.

To choose the screen type, follow this procedure:

- 1. **[Screen Select]** interface in one of the following ways:
 - > Select 【Screen Setup】 quick key→Select 【Screen Select】 submenu.
 - Select [Main Menu] quick key→from [Display] column to select [Screen Select].
- 2. Select [ECG Screen].

8.5.2. Setting ECG Alarm

To set ECG alarm properties, follow this procedure:

- 1. Select the ECG parameter area or waveform area to enter the **【ECG】** menu.
- 2. Select [Alarm] submenu.
- 3. If the alarm settings are password protected, enter the password. For details, please refer to 7.7.2 Changing Alarm Setup Protection Mode.
- Set alarms as needed.

8.5.3. Setting ECG calculating Lead

You can set the label name of the ECG calculation lead as follows:

- 1. Select ECG parameter area or waveform area to enter the **【ECG】** menu.
- 2. Select **[Setup]** submenu.
- 3. Select **[ECG 1]** or **[ECG 2]** to set label name of ECG calculating Lead.



WARNING:

■ Only when you switch on 【Multi-lead Analysis】 can you set 【ECG 2】.

8.5.4. Setting Multi-lead Analysis

When multi-lead analysis function is switched on, the **[ECG 2]** participate in the calculation of HR, the steps to set up the multi-lead analysis switch are as follows:

- 1. Select ECG parameter area or waveform area to enter the **【ECG】** menu.
- 2. Select [Setup] submenu.
- 3. Switch on or off [Multi-lead Analysis].



■ 【ECG 1 list he key calculation lead; 【ECG 2 list he auxiliary calculation lead. Only when the ECG 【Lead Type lis 5/6/12 lead can you set 【Multi-lead Analysis 】.

8.5.5. Setting ECG Waveform

8.5.5.1. Setting ECG Waveform Gain

If the ECG waveform is too small or clipped, you can change its amplitude by selecting an appropriate gain setting. To do so, follow this procedure:

- 1. Select ECG parameter area or waveform area to enter the **[ECG]** menu.
- 2. Select [Gain] submenu.
- 3. Set the size of each ECG waveform. If you select [Auto], the monitor automatically adjusts the gain of the ECG waveforms.

8.5.5.2. Setting ECG Waveform Speed

To change ECG waveform speed, follow this procedure:

- 1. Select ECG parameter area or waveform area to enter the **[ECG]** menu.
- 2. Select **Setup** submenu.
- 3. Set [Wave Speed].

8.5.5.3. Setting ECG Filter Mode

To set the ECG waveform filtering mode, follow this procedure:

- 1. Select ECG parameter area or waveform area to enter the **[ECG]** menu.
- 2. Select **Setup** submenu.
- 3. Set [Filter Mode].
 - ◆ 【Diagnose】: Use when diagnostic quality is required. The unfiltered ECG waveform is displayed so that changes such as notch on R-wave,

ST elevation or depression, etc.

- ♦ **Monitor** : Use under normal measurement conditions.
- ♦ 【Operation】: Use when the signal is distorted by high frequency or low frequency interference. High frequency interference usually results in large amplitude spikes making the ECG signal look irregular. Low frequency interference usually leads to wandering or rough baseline. The surgery filter reduces artifacts and interference from electrosurgical units. Under normal measurement conditions, selecting 【Operation】 may suppress the ORS complexes.
- ◆ 【ST】: It is recommended to use in ST segment analysis.

Filter status in various ECG modes:

Filter ECG mode	Drift filter	EMG filter	Notch Filter
Diagnose	Weak	Weak	Optional
Monitor	Moderate	Moderate	On
Operation Intense		Intense	On
ST	Weak	Moderate	Optional



- Under the mode of 【Operation 】 and 【Monitor 】, the state of the filter cannot be regulated. Only under the state 【Diagnose 】 and 【ST】 can adjust the notch filter status. Please select 【Monitor 】 during monitoring a patient, select 【Operation 】 under the state of great interference.
- The diagnose mode has passed the distortion test.

8.5.5.4. Setting Notch Filter

The notch filter can eliminate power frequency interference. Follow the steps below to turn the notch switch on or off:

- 1. Select ECG parameter area or waveform area to enter the **[ECG]** menu.
- 2. Select **[Setup]** submenu.
- 3. Switch on or off [Notch Filter].



NOTE:

 Only the [Filter Mode] is set to [Diagnose Mode] or [ST] can you switch on or off [Notch Filter], other mode is enabled by default.

8.5.5.5. Setting Notch Filter Frequency

According to the mains frequency of your country, you can set the frequency of the notch to **[50Hz]** or **[60Hz]**. If you need to change the **[Notch Frequency]**, please contact the manufacturer maintenance personnel.

8.5.6. Setting Smart Lead Switch

This monitor provides the function of switching main lead automatically. When switch on **[Smart Lead]** (Smart lead auto switchover), the current smart leads are automatically identified by the algorithm, and the host automatically switches the smart leads according to the identification of the algorithm.

Steps of switching off smart lead function are as follows:

- 1. Select ECG parameter area or waveform area to enter the **[ECG]** menu.
- 2. Select **[Setup]** submenu.
- 3. Switch off [Smart Lead].

8.5.7. Setting the Priority of the ECG Lead Off Alarm

The steps to set the alarm level for ECG lead off alarms are as follows:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】→input maintenance password→Enter.
- 2. Select 【Alarm】 submenu→【Other】 submenu.
- 3. Set [ECG Lead Off Alarm Level].

8.5.8. Adjusting the QRS Volume

The QRS volume is determined by **[Alarm Source]** in the ECG or PR alarm setting menu. Which parameter (HR or PR) is set to **[Alarm Source]** and the QRS volume is sounded according to which parameter's rhythm.

The volume of QRS sound can be set, the steps are as follows:

- 1. Select ECG parameter area or waveform area to enter the **[ECG]** menu.
- 2. Select **[Setup]** submenu.
- 3. Set [ORS Volume]

When valid SpO₂ measurements are available, the monitor adjusts the pitch tone of QRS volume based on the SpO₂ value. For detail, please refer to 10.7.8 Setting Pitch Tone.

8.5.9. Setting Multi-lead Signal Quality

The signal quality of the ECG waveform provides two display modes. The monitor displays the signal quality of the main calculated lead waveform by default. You can set the signal quality of the multi-lead waveform as required. The setting steps are as follows:

- 1. Select ECG parameter area or waveform area to enter the **【ECG】** menu.
- 2. Select [Setup] submenu.
- 3. Switch on [Multi-lead Signal].

Multi-lead Signal Quality: The color of the ECG signal of all leads is indicated by the waveform color respectively. The five colors of white, red, orange, yellow and green respectively correspond to the five signal quality levels of extreme bad, bad, general, good and excellent.

When switch off [Multi-lead Signal],

Main-lead Signal Quality: The signal quality of the main calculation lead is indicated by a triangular diagram of 5 grids, and 1 to 5 grids respectively correspond to five signal quality levels of extreme bad, bad, general, good and excellent. The signal quality is displayed above the icon (SQI) value, which unit is "%".

8.5.10. Setting ECG Standard

Select the ECG standard according to the leads you are using. To select the ECG

standard, follow this procedure:

- Select 【 Main Menu 】 quick key→from 【 System 】 column to select
 【 Maintenance 】 →input maintenance password→Enter.
- 2. Select 【Module】 submenu→ 【ECG】 submenu.
- 3. Set [ECG Standard] to [AHA] or [IEC].

8.5.11. Multi-parameter joint analysis function

The Multi-parameter joint analysis function analyzes the ECG waveform and a Pleth wave signal together to achieve more accurate measurement results through the mutual correction of HR and PR. The source of Pleth wave preferentially uses the Pleth wave, and it also can be derived from the arterial IBP wave. See 10.8.3 Setting PR Source for the setting details. To set the Multi-parameter joint analysis function, you can follow below steps:

- 1. Select ECG parameter area or waveform area to enter the **【ECG】** menu.
- 2. Select **[Setup]** submenu.
- 3. Set [MultiParam Ana] to [ON] or [OFF].

When **[MultiParam Ana]** is **[ON]**, ECG parameter area provides ECG wave and Pleth wave signal quality and joint status indicator:

- > The quality of ECG wave and Pleth wave signals are excellent, and ECG and Pleth wave signals are analyzed independently.
- The quality of Pleth wave signal is poor, and PR parameter calculation may be inaccurate. The ECG waveform signal will be used to correct the PR parameter, and the quality of the ECG signal will be framed.
- The quality of ECG wave signal is poor, and HR parameter calculation may be inaccurate. The Pleth wave signal will be used to correct the HR parameter, and the quality of the Pleth wave signal will be framed.

8.6. Arrhythmia Monitoring

Arrhythmia monitoring is applicable for adult, pediatric and neonatal patients.

8.6.1. Safety Information



WARNING:

- Arrhythmia may affect heart rate. When monitoring arrhythmia patients, do not rely entirely on the alarm information calculated by heart rate, but always place the patients under close surveillance.
- Arrhythmia function is applicable for detecting certain ventricular and atrial arrhythmias, not all atrial or supraventricular arrhythmias. Sometimes, it may detect wrong arrhythmia. Therefore, doctors must combine more clinical manifestations to analyze arrhythmia information.



\ CAUTION:

- Since the arrhythmia detection algorithm sensitivity and specificity is less than 100%, sometimes there may be some false arrhythmias detected and also some true arrhythmia events may not be detected. This is especially true when the signal is noisy.
- The amplitude of ECG waveform will affect the arrhythmia detection and heart rate calculation sensitivity.
- If the QRS amplitude is too low, the monitor may not be able to calculate the heart rate and false asystole may occur.
- Arrhythmia detection may not be available during ECG relearning. Therefore, the patient's state should be closely observed during ECG relearning and within a few minutes after completion.

8.6.2. Arrhythmia Events

This section lists all arrhythmia events and their criteria.

Arrhythmia Events	Description
Asystole	There is no fluctuation or very small and slow waveform for 6
	seconds.
Vent Fib/Tach	Ventricular fibrillation waveform for 4 seconds.

V-Tach	More than 5 (including 5) ventricular waveforms were detected		
	continuously, and the heart rate was greater than the ventricular		
	tachycardia heart rate limit.		
Vent Brady	More than 3 (including 3) ventricular waveforms were detected		
	continuously, and the heart rate was less than the ventricular		
	bradycardia limit.		
Extreme Tachy	Non-ventricular rhythm and the heart rate are greater than the extreme		
	tachycardia limit.		
Extreme Brady	Non-ventricular rhythm and the heart rate are less than the extreme		
	bradycardia limit.		
R on T	Ventricular premature beats appear on the T wave of the previous		
	cardiac cycle.		
Tachy	Non-ventricular rhythm and the heart rate are greater than the		
	tachycardia limit.		
Brady	Non-ventricular rhythm and heart rate less than bradycardia limit.		
Nonsustained V-Tach	Three or four consecutive ventricular waveforms and the heart rate are		
	greater than the ventricular tachycardia heart rate limit.		
Vent Rhythm	More than 5 (including 5) ventricular waveforms were detected		
	continuously, and the heart rate was less than the ventricular		
	tachycardia heart rate limit and greater than the ventricular bradycardia		
	heart rate limit.		
PNC	One cardiac leak and one pacing pulse were detected.		
PNP	One cardiac leak was detected, but no pacing pulse was detected.		
Pause	No heartbeat is detected within 1.75× of the average R-R interval		
	(when the heart rate is less than 100), or no heartbeat is detected within		
	1 second (when the heart rate is more than 100) and the current RR		
	interval is greater than 4 seconds and less than 6 seconds.		
Pauses/min High	The number of Pause per minute is greater than the decision limit.		
Run PVCs	For 3 or 4 consecutive ventricular waveforms, the heart rate is less		
	than the ventricular tachycardia heart rate limit and greater than the		
	ventricular bradycardia heart rate limit.		
Couplet	Two consecutive ventricular waveforms.		
Bigeminy	Dominant rhythm of N, V, N, V.		
Trigeminy	Dominant rhythm of N, N, V, N, N, V.		
Frequent PVCs	The number of PVC per minute is greater than the decision limit.		
PVC	Occasional ventricular premature beat.		

Missed Beat	No heartbeat is detected within 1.75× of the average R-R interval (when the heart rate is less than 100bpm), or no heartbeat is detected within 1 second (when the heart rate is more than 100bpm) and the current RR interval is less than 4 seconds.	
A-Fib	RR interval of normal cardiac beats is irregular and there is no P wave.	
A-Fib End	No atrial fibrillation was detected within the delay time after the end of atrial fibrillation.	
ECG Noise	There is too much noise to analyze the waveform.	
Irregular Rhythm	Always an irregular rhythm.	
Irregular Rhythm End	No irregular rhythm was detected within the delay time after the end of the irregular rhythm.	

8.6.3. Arrhythmia alarm settings

Use the following steps to set arrhythmia related alarms:

- 1. Select ECG parameter area or waveform area to enter **[ECG]** menu.
- 2. Select 【ARR】 submenu→ 【Alarm】 submenu.
- 3. If the arrhythmia setting is protected by a password, enter the password. For details, please refer to 7.7.2 *Changing Alarm Setup Protection Mode*.
- 4. Set each arrhythmia alarm as required.



NOTE: The alarm level for lethal arrhythmia is always high and cannot be changed by the user.

8.7. ST Monitoring

ST segment of ECG waveform refers to the phase from the end of ventricular depolarization to the beginning of ventricular repolarization, or from the end of QRS complex (point J) to the beginning of T wave. ST segment analysis is mostly used to monitor the oxygen supply and myocardial viability of patients. ST segment analysis function is applicable to adults, pediatric and neonatal patients.

8.7.1. Safety Information



WARNING:

- Factors such drugs, metabolism or conduction disorders may affect ST values.
- Since ST is calculated by a fixed delay after point J, it may be affected by changes in heart rate.
- The data accuracy of ST algorithm has been tested, and its clinical significance should be decided by doctors.
- The monitor provides ST segment change information, and the clinical opinion of this information should be decided by the doctor.

8.7.2. Enabling ST Monitoring

The ST segment analysis function is disabled by default. Please enable ST segment analysis according to the following steps:

- 1. Select ECG parameter area or waveform area to enter **[ECG]** menu.
- 2. Select **ST** submenu→ **Setting** submenu.
- 3. Switch on the **【ST Analysis】**. The following clinical situations may make it difficult to obtain reliable ST monitoring:
 - ◆ Lead with low noise cannot be obtained.
 - ◆ Arrhythmia leading to irregular baseline exists, such as atrial fibrillation/atrial flutter.
 - ◆ The patient is continuously performing ventricular pacing.
 - ◆ The patient has left bundle branch block.

When these situations exist, you should consider turning off the ST segment analysis function.

8.7.3. Displaying ST parameter

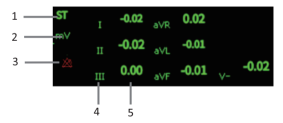
The method of displaying ST parameters and waveforms is as follows:

- 1. Enter the **Screen Layout** page in one of the following ways:
 - ◆ Select [Screen Setup] quick key—select [Screen Layout]

submenu.

- ◆ Select 【Main Menu】 quick key→from 【Display】 column to select 【Screen Layout】.
- Click on the location in the parameter area where ST parameters need to be displayed, and select 【ECG】→【ST】. Depending on the type of lead you are using, the ECG parameter area displays different ST parameters:
 - ♦ When using 3-lead monitoring, an ST parameter value is displayed in the ECG parameter area but not in the ST parameter area.
 - When using 5-lead monitoring, the ST parameter area displays 7 ST parameter values, namely ST-I, ST-II, ST-III, ST-aVR, ST-aVL, ST-aVF and ST-V respectively.
 - ◆ When 6-lead monitoring is used, the ST parameter area shows the same values of 8 ST parameters, namely ST-I, ST-II, ST-III, ST-aVR, ST-aVL, ST-aVF, ST-Va and ST-Vb.
 - ◆ When 12-lead monitoring is used, the ST parameter area displays 12 ST parameter values, namely ST-I, ST-II, ST-III, ST-aVR, ST-aVL, ST-aVF, ST-V1, ST-V2, ST-V3, ST-V4, ST-V5, and ST-V6.

Take 5-lead as an example, the ST parameter area is shown as follows:



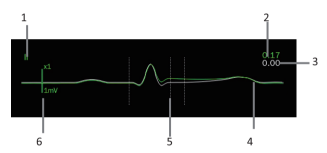
- (1) Parameter label
- (3) ST alarm off symbol
- (2) ST unit
- (4) Lead label
- (5) ST numerics: a positive value indicates ST segment elevation, and a negative value indicates ST segment depression.

8.7.4. Displaying ST Segment in Waveform Area

The steps for displaying ST segment in waveform area are as follows:

- 1. Enter the **[Screen Layout]** page in one of the following ways:
 - ◆ Select 【Screen Setup 】 quick key→select 【Screen Layout 】
 - ◆ Select 【Main Menu】 quick key→from 【Display】 column to select 【Screen Layout】.
- Click on the waveform area where you need to display ST segment, and select
 【ECG】→【ST Segment】 from the list.

The ST waveform area displays the current ST segment waveform and baseline waveform, the current ST value and baseline value. Generally, the current ST segment and parameter values are displayed in green, while the baseline segment and parameter values are displayed in white.



- (1) ST lead
- (2) The current ST value
- (3) ST baseline value
- (4) The current ST segment (green) and baseline ST segment (white)
- (5) ST segment measurement position line
- (6) Scale

8.7.5. Entering ST View

ST View displays a complete QRS segment of each ST lead. You can enter 【ST View】 to view these ST segments. The color of the current ST segment and ST value is the same as that of ECG waveform, usually green. ST baseline segment and baseline

value are white.

You can select the ST waveform area to enter the **[ST View]** page or enter the **[ST View]** page through the following steps:

- Select ECG parameter area, waveform area or ST parameter area to enter [ECG] menu.
- 2. Select **[ST]** submenu.
- 3. Select **[ST View]** from the bottom of the menu.

8.7.6. Saving the ST Baseline

ST analysis requires valid samples. Set an ST baseline when ST values become stable. If you do not set a baseline, the monitor will automatically save a set of baselines about 5 minutes after a valid ST measurement appears. You can also manually update the baseline by selecting **[Set Baseline]** in the lower left corner of the **[ST View]** interface.

You can also make the following settings under the ST interface:

- Select [Show Baseline] or [Hide Baseline] to show or hide ST baseline segments and parameter values.
- Select [Show Mark] or [Hide Mark] to show or hide ST reference point, J point and ST point positions.



■ Changing the ST baseline will affect the ST alarm.

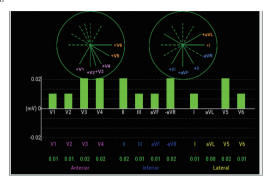
8.7.7. Entering ST Graphic Window

The steps to enter the ST Graphic window are as follows:

- Select ECG parameter area, waveform area, ST parameter area or ST waveform area to enter 【ECG】 menu.
- Select [ST View] submenu.
- 3. Select **[ST Graphic]** from the bottom of the **[ST View]** menu.

The following figure shows ST Graphic. The height of the bar represents the ST

value of the corresponding ST lead. The color of the bar indicates the ST alarm status: green indicates that the ST value is within the normal range; Cyan, yellow and red indicate that the ST value exceeds the alarm limit. The alarm color corresponds to the level of ST alarm.



8.7.8. ST Setup

8.7.8.1. Setting ST Alarm

ST alarm is set as follows:

- Select ECG parameter area, waveform area or ST parameter area to enter [ECG] menu.
- 2. Select **ST** submenu→ **[Alarm]** submenu.
- 3. Set the properties of ST alarm as required.

8.7.8.2. Showing ISO Point, J Point and ST Point Marks

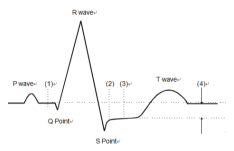
The ISO point, J point and ST point position marks are not displayed by default on the ST segment in the waveform area. To display these marks, the steps are as follows:

- Select ECG parameter area, waveform area, ST parameter area or ST waveform area to enter 【ECG】 menu.
- 2. Select 【ST】 submenu→ 【Setting】 submenu.
- 3. Switch on **[ST Mark]**.

8.7.9. Adjusting ST Measurement Point

8.7.9.1. ST Point, ISO Point and J Point

The ST value for each beat complex is the vertical difference between the isoelectric (ISO) point and the ST point. The 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 the J point is a fixed distance away from the ST point, it can be useful to help you correctly position the ST point.



(1) ISO Baseline Point (2) J Point (3)ST Measurement Point (4) ST Value

8.7.9.2. Setting ST Point, J Point and ISO Point



CAUTION:

When you start monitoring or the patient's heart rate or ECG waveform has obvious changes, it may affect the length of QT interval, thus affecting the position of ST points, so the positions of ISO and ST points need to be adjusted. Incorrect setting of ISO point or ST point may lead to false ST segment depression or elevation. Please always ensure that the location of ST measurement point is suitable for the patients under monitoring.

The steps for setting ST, J and ISO points are as follows:

- Select ECG parameter area, waveform area, ST parameter area or ST waveform area to enter 【ECG】 menu.
- 2. Select 【ST】 submenu→【Adjust】 submenu.

3. Select **ST Point** to set the position of ST Point.

The setting of 【Auto Adjust】 defines the method of adjusting the ISO point and J point. When the 【Auto Adjust】 switch is turned on, the module automatically adjusts the positions of ISO and J points according to the current waveform. When the 【Auto Adjust】 switch is off, you can manually adjust the positions of 【ISO】 and 【J】 through "+" and "-".

- ◆ The ISO point (isoelectric) position is given relative to the R-wave peak. Position the ISO point in the middle of the flattest part of the baseline (between the P and Q waves).
- ◆ The J point position is given relative to the R-wave peak and helps locating the ST point. Position the J point at the end of the QRS complex and the beginning of the ST segment.
- ◆ The ST point is located at a fixed distance relative to the J point, and the J point is moved so that the ST point is located in the middle of the ST segment. The ST point can be located at the positions of J+0, J+20, J+40, J+60, and J+80

8.8. QT/QTc Monitoring

QT interval is the time from the beginning of QRS complex to the end of T wave, that is, the whole period of ventricular action potential depolarization (QRS interval) and repolarization phase (ST-T). QT test can help you to judge long QT interval syndrome.

QT interval is negatively correlated with heart rate. As heart rate increases, the QT interval shortens, while at lower heart rates QT interval gets longer. We can use several formulas to correct QT interval according to heart rate. The QT interval corrected by heart rate is called QTc.

QT/QTc monitoring is applicable for adults, pediatric and neonatal patients.

8.8.1. QT/QTc measurement limitation

The following conditions may affect the accuracy of OT measurement:

- The amplitude of R wave is too low.
- Excessive ventricular heartbeat.
- RR interval is unstable.

- High heart rate causes P wave to invade the end of the previous T wave.
- T wave is too flat or t wave boundary is unclear.
- The existence of U wave makes the end of T wave difficult to define.
- > QTc measurement is unstable.
- In the presence of noise, asystole, ventricular fibrillation, and ECG lead off.

In the above situation, you need to select leads with good T wave amplitude, no visible flutter, and no dominant U wave or P wave. In some cases, such as left and right bundle branch block or cardiac hypertrophy, QRS complex may widen. If a long QTc is observed, this should be confirmed to ensure that it is not caused by QRS broadening.

QT measurement cannot be performed in the presence of bigeminy rhythm because normal cardiac beats are not included in the analysis when they are followed by ventricular beats.

QT measurement cannot be performed when the heart rate is extremely high (adults over 150bpm, pediatric and neonatal over 180bpm). When the heart rate changes, it can take several minutes for the QT interval to stabilize. In order to obtain reliable QTc calculation results, it is important to avoid areas where the heart rate changes.

8.8.2. Enabling the QT/QTc Monitoring

QT/QTc monitoring function is off by default, and you need to turn it on before performing QT/QTc monitoring. Enable QT/QTc monitoring as follows:

- 1. Select ECG parameter area or waveform area to enter **[ECG]** menu.
- 2. Select 【QT】 submenu→【Setting】 submenu.
- 3. Switch on **[QT Analysis]**.

8.8.3. Displaying QT/QTc Parameter

The method for displaying QT parameters and waveforms is as follows:

- 1. Enter the 【Screen Layout】 page in one of the following ways:
 - ◆ Select 【Screen Setup】 quick key→select 【Screen Layout】submenu.
 - ◆ Select 【Main Menu】 quick key→from 【Display】 column to select 【Screen Layout】.
- Click on the location in the parameter area where QT parameters need to be displayed and select 【ECG】→【QT】.



NOTE: QTc value is calculated based on QT-HR, not ECG-HR calculation

leads. You can enter QT View to view QT-HR. For details, please refer to 8.8.4 Entering QT View.

The QT parameter area is displayed as follows. Depending on the settings, the display of your monitor may be different.



- (1) QTc alarm limit (if QTc alarm is off, the alarm off icon is displayed here)
- (2) Parameter Unit
- (3) Parameter Label
- (4) QTc value
- (5) ΔQTc value (the difference between the current value of QT_C and the baseline value; if ΔQTc alarm is off, the alarm off icon is displayed on the right side of the value)
- (6) QT value

8.8.4. Entering QT View

QT View displays the current QT parameter values and waveforms, as well as baseline/reference QT parameter values and waveforms. The steps to enter 【QT View】 are as follows:

- 1. Select QT parameter area to enter 【QT】 menu.
- 2. Select the 【QT View】 button at the bottom of the menu.



The following figure shows an example of QT View:

- ◆ The current waveform is displayed at the top of the view, and the color is the same as the ECG waveform, usually green.
- ◆ The baseline segment is displayed below in white.
- ◆ The starting point of QRS complex and the ending point of T wave are marked with vertical lines.
- In some cases, the algorithm may not be able to give QT measurement results because the waveform does not meet the requirements. At this time, the reason that cannot be analyzed will be displayed below the QT parameter area in QT View. In addition, a prompt message "QT cannot be analyzed" will be displayed in the technical alarm information area of the main interface.
- Select the lead label at the lower left of QT View, switch leads, and highlight the waveforms of the corresponding leads.

8.8.5. Setting the QT Baseline

Setting QT baseline is helpful to quantify QTc changes. After QT valid values appear, if you do not set QT baseline within 5 minutes, the monitor will automatically set QT baseline.

The steps for manually setting QT baseline are as follows:

- 1. Select the **[Set Baseline]** button below OT View.
- 2. Set the current QT parameter value as the baseline. The baseline value will be used to calculate the Δ QTc value. After the new QT baseline is set, the

original baseline will be discarded. The baseline will be cleared when the patient is released.

Select **[Show Baseline]** or **[Hide Baseline]** to show or hide QT baseline waveform.



■ Changing QT baseline will affect \triangle QTc value and \triangle QT alarm.

8.8.6. QT Setting

8.8.6.1. Setting the QT Alarm

QT alarm is set as follows:

- 1. Select QT parameter area to enter **[QT]** menu.
- 2. Select [Alarm] submenu.
- 3. Set properties of QTc and \triangle QTc alarm.

8.8.6.2. Selecting QTc Formula

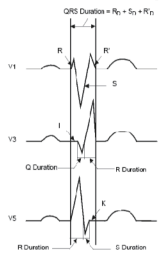
The monitor uses Hodges formula by default to correct QT interval according to heart rate. If you need to select other QTc formulas, the steps are as follows:

- Select 【 Main Menu 】 quick key→from 【 System 】 column to select
 【 Maintenance 】 →input maintenance password→Enter.
- 2. Select 【Module】 submenu→ 【ECG】 submenu.
- 3. Select [QTc Formula].
 - ♦ Hodges: $QTc = QTc + 1.75 \times (HeartRate 60)$
 - Bazett: $QTc = QT \times \left(\frac{\text{Heat Rate}}{60}\right)^{\frac{1}{2}}$
 - Fridericia: $QTc = QT \times \left(\frac{\text{Heart Rate}}{60}\right)^{\frac{1}{3}}$
 - Framingham: $QTc = QT + 154 \times \left(1 \frac{60}{Heart Rate}\right)$

8.9. Isoelectric Segments

Between the global onset and offset of the QRS-complex, signal parts with a duration of more than 6 ms and amplitudes not exceeding $20\mu V$ for at least three samples should be defined as isoelectric segments – I-wave before the global QRS-ONSET and K-wave after the global QRS-OFFSET.

Isoelectric parts (I-wave) after global QRS-ONSET or before global QRS-OFFSET (K-wave) are excluded in the duration measurement of the respective adjacent waveform.



8.10. ECG Relearning

Changes in ECG templates may result in erroneous arrhythmia alarms or/and inaccurate heart rates.

The monitor provides ECG relearning function. ECG relearning enables the monitor to learn new ECG templates to correct arrhythmia alarms and heart rate values. After ECG relearning is completed, the monitor stores the QRS wave form obtained by learning as a template as the normal ECG wave form of the patient. During ECG monitoring, when you suspect abnormal arrhythmia alarm, you may need to start an ECG relearning.

8.11. Calibrating ECG

The ECG signal may be inaccurate due to hardware or software problems. As a result, the ECG waveform amplitude becomes greater or smaller. In that case, you need to

calibrate the ECG module. To do so, follow this procedure:

- Select ECG parameter area or waveform area, set [Filter Mode] to [Diagnose].
- Select 【 Main Menu 】 quick key→from 【 System 】 column to select
 【 Maintenance 】 →input maintenance password→Enter.
- 3. Select 【Module】 submenu→ 【ECG】 submenu.
- 4. Select [Calibrate], the square wave signal will appear on the screen to compare the amplitude of the square wave with the scale. The error range should be within 5%. The ECG calibration must be completed by the maintenance personnel.

8.12. Defibrillation Synchronization

The module provides an analog out connector to output defibrillation synchronization signal. If a defibrillator is connected, it receives a synchronization pulse (100 ms, +5 V) through the analog out connector each time an R-wave is detected.

The steps to set defibrillation synchronization are as follows:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】→input maintenance password→Enter.
- 2. Select [Module] submenu→ [Auxiliary Output] submenu.
- 3. Set the defibrillation synchronization signal as needed.



CAUTION:

- Improper use of a defibrillator may cause injury to the patient. The operator should determine whether to perform defibrillation or not according to the patient's condition.
- According to AAMI specifications the peak of the synchronized defibrillator discharge should be delivered within 60 ms of the peak of the R-wave. The signal at the ECG output on the monitors is delayed by maximum of 25ms.

8.13. ECG Troubleshooting

This section lists the problems that might occur. If you encounter problems when using the monitor or accessories, check the table below before requesting for services. If the problem persists after you have taken corrective actions, contact your service personnel.

Corrective Actions		
1. Check that electrodes are not detached or dry. Replace with fresh and		
moist electrodes if necessary.		
2. Check that leadwires are not defective. Replace leadwires if necessary.		
3. Check that patient cable or leadwires are routed too close to other		
electrical devices. Move the patient cable or leadwires away from		
electrical devices.		
Use ESU-proof ECG cables.		
Inadequate skin preparation, tremors, tense subject, and/or poor electrode		
placement.		
1. Perform skin preparation again and re-place the electrodes. For more		
information, see 8.4.1 Preparing the Patient Skin		
2. Apply fresh, moist electrodes. Avoid muscular areas.		
1. Check that cables are properly connected.		
2. Check that electrodes are not detached or dry. Perform skin preparation		
again as described in 8.4.1 Preparing the Patient Skin and apply fresh		
and moist electrodes		
3. Check that the patient cable or leadwires are not damaged. Change		
them if necessary.		
1. Check that electrodes are not dry. Perform skin preparation again and		
replace the electrodes. For more information, see 8.4.1 Preparing the		
Patient Skin.		
2. Check for excessive patient movement or muscle tremor. Reposition the		
electrodes. Replace with fresh and moist electrodes if necessary.		
1. Check that the ECG gain is not set too low. Adjust the gain as required.		
For more information, see 8.5.5 Setting ECG Waveforms.		
2. Perform skin preparation again and re-place the electrodes. For more		

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Problem	Corrective Actions
	information, see 8.4.1 Preparing the Patient Skin.
	3. Check electrode application sites. Avoid bone or muscular area.
	4. Check that electrodes are not dry or used for a prolonged time. Replace
	with fresh and moist electrodes if necessary.
No ECG Waveform	1. Check that the ECG gain is not set too low. Adjust the gain as required.
	For more information, see 8.5.5 Setting ECG Waveforms.
	2. Check that the leadwires and patient cables are properly connected.
	Change cable and leadwires.
	3. Check that the patient cable or leadwires are not damaged. Change
	them if necessary.
Base Line Wander	1. Check for excessive patient movement or muscle tremor. Secure
	leadwires and cable.
	2. Check that electrodes are not detached or dry and replace with fresh and
	moist electrodes if necessary. For more information, see 8.4.1 Preparing
	the Patient Skin.
	3. Check for ECG filter setting. Set ECG Filter mode to 【Monitor】.



NOTE: Physiological alarm and technical alarm information refer to D Alarm

Message.

Chapter 9 Respiration Rate (RESP)

9.1. Introduction

Impedance respiration is measured across the thorax. When the patient is breathing, the volume of air changes in the lungs, resulting in impedance changes between the electrodes. Respiration rate (RR) is calculated from these impedance changes, and a respiration waveform appears on the patient monitor screen.

Resp monitoring is applicable for adult, pediatric and neonatal patients

9.2. Safety Information



WARNING:

- If you do not set the detection level for the respiration correctly in manual detection mode, it may not be possible for the monitor to detect apnea. If you set the detection level too low, the monitor is more likely to detect cardiac activity, and to falsely interpret cardiac activity as respiratory activity in the case of apnea.
- The respiration measurement does not recognize the cause of apneas. It only indicates an alarm if no breath is detected when a pre-adjusted time has elapsed since the last detected breath. Therefore, it cannot be used for diagnostic purpose.
- If operating under conditions according to the EMC Standard IEC 60601-1-2 (Radiated Immunity 3V/m), field strengths above 3V/m may cause erroneous measurements at various frequencies. Therefore, it is recommended to avoid the use of electrically radiating equipment in close proximity to the respiration measurement unit.
- The impedance respiration measurement may cause rate changes in Minute Ventilation Rate Responsive Pacemakers. Set the pacemaker rate responsive mode off or disable the impedance respiration measurement on the monitor.
- When using the electrosurgery unit, ensure proper contact of the ESU return electrode to the patient to avoid burns at monitor measurement sites. Also ensure that the ESU return electrode is near the operating area.



CAUTION:

- Only use parts and accessories specified in this manual, and obey all warnings and cautions.
- Respiration monitoring is not for use on the patients who are very active, as this will cause false alarms.

9.3. RESP Display



(1) Parameter label

(2) RESP waveform gain

(3) RESP lead

(4) Resp waveform

- (5) RR unit
- (6) RR Alarm limits: If the respiration rate is turned off, the alarm off icon will be displayed here.
- (7) RR value

(8) RR source

9.4. Preparing for RESP Monitoring

9.4.1. Preparing the Patient skin

Follow this procedure to prepare the patient:

- 1. Shave hair from skin at chosen sites.
- 2. Gently rub skin surface at sites to remove dead skin cells.
- 3. Thoroughly cleanse the site with a mild soap and water solution.
- 4. Dry the skin completely before applying the electrodes.



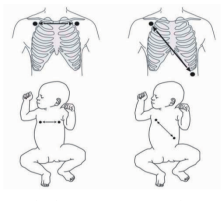
CAUTION:

Proper skin preparation is necessary for good signal quality at the electrode site,
 as the skin is a poor conductor of electricity

9.4.2. Placing the Electrodes

As the Respiration measurement adopts the standard ECG electrode placement, you can use different ECG cables. Since the respiration signal is measured between two ECG electrodes, if a standard ECG electrode placement is applied, the two electrodes should be RA and LA, or RA and LL.

For more information, see 8.4.2 Applying Electrodes.



RA-LA RA-LL



CAUTION:

- Correct electrodes placement can help to reduce cardiac overlay: avoid the liver area and the ventricles of the heart in the line between the respiratory electrodes. This is particularly important for neonatals.
- Some patients with restricted movements breathe mainly abdominally. In these cases, you may need to place the left leg electrode on the left abdomen at the point of maximum abdominal expansion to optimize the respiratory wave.
- In clinical applications, some patients (especially neonatals) expand their chests laterally, causing a negative intrathoracic pressure. In these cases, it is better to

place the two respiration electrodes in the right midaxillary and the left lateral chest areas at the patient's maximum point of the breathing movement to optimize the respiratory waveform.

■ Regularly inspect the electrode application site to ensure skin quality. If there are signals of allergies, replace the electrodes or change the application site.



NOTE:

- Store the electrodes at room temperature. Open the electrode package immediately prior to use.
- Check that the electrode packages are intact and not expired. Make sure the electrode gel is moist.

9.5. RESP Settings

9.5.1. Setting the RESP Alarm

To set the RESP alarm properties, follow this procedure:

- 1. Select the Resp parameter area or waveform area to enter the **【RESP】** menu.
- 2. Select [Alarm] submenu.
- 3. If the alarm setting is protected by password, enter the password. For detail, please refer to 7.7.2 Changing Alarm Setup Protection Mode.
- 4. Set alarms as needed.

9.5.2. Selecting RR Source

You can select RR source, follow this procedure:

- 1. Select the RESP parameter area or waveform area to enter the **[RESP]** menu.
- 2. Select [Setup] submenu.
- 3. Set [RR Source]. When you select [Auto], the system automatically selects the RR source according to the priority. The priority of the RR source is CO₂, RM, ECG and SpO₂. When the current RR source does not have valid measurement, the system automatically switches the [RR Source] to [Auto].

9.5.3. Selecting Respiration Lead

You can set up respiration lead to get the best respiratory waveform. The steps to set up breathing leads are as follows:

- 1. Select the RESP parameter area or waveform area to enter the **[RESP]** menu.
- 2. Select **[Setup]** submenu.
- Set 【RESP Lead 】. If the respiratory waveform is still poor after adjusting the
 respiration lead or the respiration rate measurement is suspected to be
 inaccurate, you can adjust the electrode position.

9.5.4. Setting RESP Waveform Gain

You can adjust the RESP waveform gain to better view the waveform amplitude. The steps to set the RESP waveform gain are as follows:

- 1. Select the RESP parameter area or waveform area to enter the **[RESP]** menu.
- 2. Select **[Setup]** submenu.
- 3. Set [Gain].

9.5.5. Setting the RESP Waveform Speed

To set the RESP waveform speed, follow this procedure:

- 1. Select the RESP parameter area or waveform area to enter the **[RESP]** menu.
- 2. Select **[Setup]** submenu.
- 3. Set [Wave Speed].

9.5.6. Setting the Auto Detection Switch

To set the auto detection switch, follow this procedure:

- 1. Select the RESP parameter area or waveform area to enter the **【RESP】**menu.
- 2. Select **[Setup]** submenu.
- 3. Switch on or off [Auto Threshold Detection].
 - > If **[Auto Threshold Detection]** is switched on, the monitor automatically adjusts the RESP waveform detection level, or threshold.
 - **\rightarrow (Auto Threshold Detection)** is switched off, you have to manually

adjusts the RESP waveform threshold. For more information, see 9.5.7. Manually Adjust the RESP Waveform Detection Threshold.

9.5.7. Manually Adjust the RESP Waveform Detection Threshold

Use the manual detection mode in the following situations:

- The respiration rate and the heart rate are close.
- Patients have intermittent mandatory ventilation.
- Respiration is weak. Try repositioning the electrodes to improve the signal.

To set the Resp waveform threshold to the desired level, follow this procedure:

- 1. Select the RESP parameter area or waveform area to enter the **[RESP]** menu.
- 2. Select [Threshold] submenu.
- 3. Select the up and down arrows below **[Threshold]** to define the Resp waveform threshold. Once set, the detection level will not adapt automatically to different respiration depths. It is important to remember that if the depth of breathing changes, you may need to change the detection level.

9.5.8. Setting the Respiration Filter

Turn on the respiration filter function can filter out the interference in the respiration waveform. The steps to set the respiration filter switch are as follows:

- 1. Select the RESP parameter area or waveform area to enter the **【RESP】** menu.
- 2. Select **[Setup]** submenu.
- 3. Turn on or turn off 【RESP Filter】.

9.6. RESP Troubleshooting

For more information, see D Alarm Message

Chapter 10 SpO₂

10.1. Introduction

Pulse Oxygen Saturation (SpO₂) monitoring is a non-invasive technique used to measure the amount of oxygenated haemoglobin and pulse rate by measuring the absorption of selected wavelengths of light. The light generated in the emitter side of the probe is partly absorbed when it passes through the monitored tissue. The amount of transmitted light is detected in the detector side of the probe. When the pulsative part of the light signal is examined, the amount of light absorbed by the haemoglobin is measured and the pulse oxygen saturation can be calculated. This device is calibrated to display functional oxygen saturation.

You can simultaneously measure SpO_2 using the P1 or MPS-P module and the SpO_2 plug-in module. The measurement value from the the P1 or MPS-P module is labeled SpO_2 and the measurement value from the SpO_2 plug-in module is labeled SpO_2L .

The monitor can support Masimo SpO_2 plug-in module or Nellcor SpO_2 plug-in module.

10.2. Safety Information



WARNING:

- Use only SpO₂ sensors specified in this manual. Follow the SpO₂ sensor's instructions for use and adhere to all warnings and cautions. The SpO₂ probe specified in the manual has been tested with the monitor and meets the requirements of ISO80601-2-61.
- Before use, the operator needs to verify the compatibility between the monitor, probe and cable. Otherwise, it may cause injury to the patient.
- When a trend toward patient deoxygenation is indicated, analyze the blood samples with a laboratory co-oximeter to completely understand the patient's condition.
- Do not use SpO₂ sensors during magnetic resonance imaging (MRI). Induced current could potentially causes burns. The sensor may affect the MRI image,

and the MRI unit may affect the accuracy of the oximetry measurements.

- Prolonged continuous monitoring may increase the risk of undesirable changes in skin characteristics, such as irritation, reddening, blistering or burns. Inspect the sensor site every 2 hours and move the sensor if the skin quality changes. Change the application site every 4 hours. For neonates, or patients with poor peripheral blood circulation or sensitive skin, inspect the sensor site more frequently.
- Functional testers cannot be used to evaluate the accuracy of pulse oximetry probes and pulse oximetry monitors.



CAUTION:

Use only specified accessories in this manual. Follow the instructions for use and adhere to all warnings and cautions.



NOTE:

- Functional test equipment or SpO₂ simulators can be used to evaluate pulse rate accuracy.
- Functional testing equipment or oximetry simulators should not be used to verify the accuracy of oximetry monitors and pulse oximetry probes.
- The accuracy of oximetry monitor and pulse oximetry probe needs to be verified by clinical data.
- The SpO₂ probe and extension cord match with this monitor were confirmed and tested with the conformity of ISO 80601-2-61.
- The monitor does not provide automatic generation of SpO_2 self-detection alarm signals. Operators need to use SpO_2 simulator for detection.

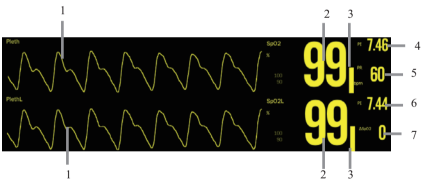
10.3. SpO₂ Measurement Limitations

If you doubt the SpO₂ measurements, check the patient's vital signs first, then check the monitor and SpO₂ sensor. The following factors may influence the accuracy of

measurements:

- There is excessive illumination from light sources such as a surgical lamp, a brilirubin lamp, or sunlight;
- Excessive patient movement;
- Diagnostic test;
- Low perfusion;
- Electromagnetic interference, such as MRI device;
- Electrosurgical equipment;
- > Concentration of nonfunctional hemoglobin, such as carbonyl hemoglobin (COHb) and methemoglobin(MetHb);
- The presence of certain dyes, such as methylene blue or indigo carmine;
- Improper placement or incorrect use of pulse oximeter probe;
- > Shock, anemia, hypothermia or use of vasoconstrictor drugs, which can cause blood flow in the arteries to drop to unmeasurable levels.

10.4. SpO₂ Display



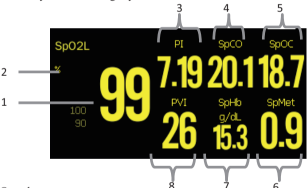
- Pleth waveform (Pleth/PlethL): The amplitude of the Pleth/PlethL waveform can directly reflect the strength of the patient's pulse signal. The Pleth waveform is not normalized.
- (2) SpO₂ value (SpO₂/ SpO₂L): Percentage of oxygenated hemoglobin in relation to the sum of oxyhemoglobin and deoxyhemoglobin.
- (3) Pleth bar: Proportional to the intensity of the pulse.
- (4) Perfusion index (PI): Gives the numerical value for the pulsatile portion of the measured signal caused by arterial pulsation. PI indicates the signal strength of

SpO₂ and also partially indicates the signal quality.

- ◆ Above 1 is optimal;
- Between 0.3 and 1 is acceptable;
- ◆ Below 0.3 indicates low perfusion. When 0.3≤PI<1, the PI values will be displayed in yellow background; PI<0.3, PI values will be displayed in red background, and SpO₂ parameter values are displayed as hollow words. Reposition the SpO₂ sensor or find a better site. If low perfusion persists, choose another method to measure oxygen saturation if possible.
- (5) Pulse rate: the number of pulses detected per minute (from the pleth waveform)
- (6) $\Delta \operatorname{Sp}O_2$: $(\Delta \operatorname{Sp}O_2 = |\operatorname{Sp}O_2 \operatorname{Sp}O_2L|)$

> For Masimo module

The Masimo module is intended to monitor the SpO₂, SpMet, PVI, SpHb, SpOC, PI, SpCO of patients. The following figure shows the SpO₂ display screen, the display on your monitor may be looked slightly different.



- 1. SpO₂ value;
- 2. SpO₂ unit;
- 3. PI (Perfusion Index) value and unit;
- 4. SpCO (Carboxyhemoglobin Saturation) value and unit.
- 5. SpOC (Oxygen Content) value and unit;
- 6. SpMet (Methemoglobin Saturation) value and unit;
- 7. SpHb (Total Hemoglobin) value and unit;
- 8. PVI (Pleth Variability Index) value and unit;

■ Parameters Description

- a) Carboxyhemoglobin Saturation (SpCO): SpCO is a value that represents the percentage of carboxyhemoglobin saturation within the blood.
- b) Methemoglobin Saturation (SpMet): SpMet is a value that represents the percentage of methemoglobin saturation within the blood.
- c) 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.
- d) Total Hemoglobin (SpHb): SpHb is a measure of the total hemoglobin concentration in arterial blood.
- e) Oxygen Content (SpOC): SpOC is a measure of the total oxygen content present in the blood.

10.5. SpO₂ Module

Accuracy confirmation of SpO_2 module measurement: SpO_2 accuracy has been confirmed in human experiments by comparing with the reference values of arterial blood samples measured by the blood gas analyzer. Compared with the blood gas analyzer measurement, the SpO_2 measurement is in accordance with the statistics distributed.

10.6. Monitoring Procedure

- Select an appropriate sensor according to the module type, patient category and weight.
- 2. Clean the contact surface of the reusable sensor.
- 3. Remove colored nail polish from the application site.
- 4. Apply the SpO₂ sensor to the patient according to the instruction for use of the sensor.
- 5. Select an appropriate extension cable according to the connector type and plug the cable into the SpO₂ connector.
- 6. Connect the SpO_2 sensor to the extension cable.



■ Do not apply sensor too tightly as this results in venous pulsation which may

severely obstruct circulation and lead to inaccurate measurements.

- At elevated ambient temperatures be careful with measurement sites that are not well perfused, because this can cause burns after prolonged application.
- Avoid placing the sensor on extremities with an arterial catheter, an NIBP cuff or an intravascular venous infusion line.
- For neonatal patients, make sure that all sensor connectors and adapter cable connectors are outside the incubator. The humid atmosphere inside can cause inaccurate measurements.
- \blacksquare SpO₂ can be measured at up to 2 locations at the same time.

10.7. Setting SpO₂

10.7.1. Setting SpO₂ Alarm

To change the SpO₂ alarm settings, follow this procedure:

- 1. Select the SpO_2 parameter area or waveform area to enter the $[SpO_2]$ menu.
- 2. Select [Alarm] submenu.
- 3. If the alarm setting is protected by password, enter the password. For detail, please refer to 7.7.2 Changing Alarm Setup Protection Mode.
- 4. Set alarms as needed.

If use SpO_2 plug-in module to measurement SpO_2L , you can set alarm properties for ΔSpO_2 .

10.7.2. Nellcor SpO₂ Alarm Delay (Sat-Seconds)

With traditional alarm management, high and low alarm limits are set for monitoring SpO_2 . During monitoring, once SpO_2 exceeds alarm limit, an audible alarm immediately sounds. When the patient SpO_2 fluctuates near an alarm limit, the alarm sounds each time the limit is violated. Such frequent alarms can be distracting.

If your monitor is configured with a Nellcor SpO₂ module, then you can set Sat-Seconds (the delay time of SpO₂ alarm) to reduce such alarms. The method of calculation is as follows: the percentage points of the SpO₂ saturation falling outside the alarm limit is multiplied by the number of seconds remaining outside the limit. This can

be stated as the equation:

$$Sat$$
-Seconds = $Points \times Seconds$

Only when the Sat-Seconds limit is reached, the monitor gives a Sat-Seconds alarm. For example, the figure below demonstrates the alarm response time with a Sat-Seconds limit set at 50 and a low SpO₂ limit set at 90%. In this example, the patient SpO₂ drops to 88% (2 points) and remains there for 2 seconds. Then it drops to 86% (4 points) for 3 seconds, and then to 84% (6 points) for 6 seconds. The resulting Sat-Seconds are:

% SpO ₂	Seconds	Sat-Seconds
2×	2=	4
4×	3=	12
6×	6=	36
Total Sat-Seconds=		52

After approximately 10.9 seconds, a Sat-Second alarm would sound, because the limit of 50 Sat-Seconds would have been exceeded.

Saturation levels may fluctuate rather than remaining steady for a period of several seconds. Often, the SpO_2 of patient may fluctuate above and below the alarm limit, re-entering the non-alarm range several times. During such fluctuation, the monitor integrates the number of SpO_2 points, both positive and negative, until either the Sat-Seconds limit is reached, or the patient SpO_2 re-enters the non-alarm range and remains there.

10.7.3. Setting the Sat-Seconds (Only for Nellcor SpO₂)

You can set the Sat-Seconds through follow this procedure:

- 1. Select the SpO_2 parameter area or waveform area to enter the $[SpO_2]$ menu.
- 2. Select [Alarm] submenu.
- 3. Set [Sat-Seconds].

10.7.4. Setting Sensitivity (Only for BLT SpO₂)

The SpO_2 value displayed on the monitor screen is the average of data collected within a specific time. The higher the sensitivity, the quicker the monitor responds to changes in the patient's oxygen saturation level. Contrarily, the lower the sensitivity, the

slower the monitor responds to changes in the patient's oxygen saturation level, but the SpO_2 measurement is more stable. For critically ill patients, selecting shorter averaging time will help understanding the patient's state.

To set the sensitivity, follow this procedure:

- 1. Select the SpO_2 parameter area or waveform area to enter the $[SpO_2]$ menu.
- 2. Select [SpO₂ Setup] submenu.
- 3. Select [Sensitivity] and then toggle between [High], [Med] or [Low].

10.7.5. Setting Averaging Time (Only for Masimo SpO₂ and Nellcor SpO₂)

The SpO_2 value displayed on the monitor screen is the average of data collected within a specific time. The shorter the averaging time, the quicker the monitor responds to changes in the patient's oxygen saturation level, but the lower the measurement accuracy. Contrarily, the longer the averaging time, the slower the monitor responds to changes in the patient's oxygen saturation level, but the higher the measurement accuracy. For critically ill patients, selecting shorter averaging time will help understanding the patient's state.

To set the averaging time, follow this procedure:

- 1. Select the SpO₂L parameter area or waveform area to enter the **[SpO₂L]** menu.
- 2. Select [SpO₂L Setup] submenu.
- 3. Set the [Averaging Time].

10.7.6. Setting NIBP measurement on the same limb

When monitoring SpO_2 and NIBP on the same limb simultaneously, you can switch on **[NIBP Simul]** to lock the SpO_2 alarm status until the NIBP measurement ends. If you switch off **[NIBP Simul]**, low perfusion caused by NIBP measurement may lead to inaccurate SpO_2 readings and therefore cause false physiological alarms.

To set the **[NIBP Simul]**, follow this procedure:

- 1. Select the SpO₂ parameter area or waveform area to enter the **[SpO₂]** menu.
- 2. Select [Alarm] submenu.
- 3. Set [NIBP Simul] as to [On] or [Off].

10.7.7. Changing the Speed of Pleth Waveform

To set the sweep speed of Pleth waveforms, follow this procedure:

- 1. Select the SpO_2 parameter area or waveform area to enter the $\llbracket SpO_2 \rrbracket$ menu.
- 2. Select [SpO₂ Setup] submenu.
- 3. Set **[Wave Speed]** to the appropriate value. The larger the value, the faster the scanning speed and the wider the waveform.

10.7.8. Setting Pitch Tone

The monitor can adjust the QRS tone according to the SpO₂ value. The pitch tone function is on by default. The steps to turn off the pitch tone function are as follows:

- 1. Select the SpO_2 parameter area or waveform area to enter the $[SpO_2]$ menu.
- 2. Select [SpO₂ Setup] submenu.
- 3. Switch off [Pitch Tone].

10.7.9. Setting PI Display

You can switch on or off PI display by following these steps:

- 1. Select the SpO_2 parameter area or waveform area to enter the $[SpO_2]$ menu.
- 2. Select **[SpO₂ Setup]** submenu.
- 3. Set **(Display PI)** as to **(On)** or **(Off)**.

10.7.10. Setting the Masimo SpO₂

When your monitor is equipped with Masimo SpO_2 module, you can set the following contents:

10.7.10.1. Setting Sensitivity mode

To set the sensitivity mode, follow these procedures:

- 1. Select the SpO_2 parameter area or waveform area to enter the $[SpO_2]$ menu.
- 2. Select [SpO₂ Setup] submenu.
- Select [Sensitivity] in the SpO₂ setting menu with the options of [Max],
 [Normal] or [APOD].

- > [Max]: This mode should be used for the sickest patients, where obtaining a reading is most difficult. The mode is recommended during procedures and when clinician and patient contact is continuous.
- [Normal]: This mode provides the best combination of sensitivity and probe-off detection performance. The mode is recommended for the majority of patients.
- > 【APOD】: This 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 mode is useful for patients that are at particular risk of the sensor becoming detached (pediatric, combative, etc.)

10.7.10.2. Setting Alarm Delay Time

To set the alarm time, follow these procedures:

- 1. Select the SpO₂ parameter area or waveform area to enter the **[SpO₂]** menu.
- 2. Select [Alarm] submenu.
- 3. Select [Alarm Delay], and you can select the alarm delay time as required or you can also set to [Off].

10.7.10.3. Setting FastSat mode

The FastSat mode is clinically applicable during procedures when detecting rapid changes in oxygen saturation is paramount such as induction, intubation and sleep studies. To set the FastSat mode, follow these procedures:

- 1. Select the SpO_2 parameter area or waveform area to enter the $[SpO_2]$ menu.
- 2. Select [SpO₂ Setup] submenu.
- 3. Select **[On]** or **[Off]** to enable or disable the **[FastSat]** mode.

10.7.10.4. Setting SmartTone

To set the SmartTone, follow these procedures:

- 1. Select the SpO_2 parameter area or waveform area to enter the $[SpO_2]$ menu.
- 2. Select [SpO₂ Setup] submenu.

- 3. Select **(On)** or **(Off)** to enable or disable the **(SmartTone)**.
 - When you set it to **[On]**, it will allow the audible pulse beep to beep when the pleth shows signs of motion.
 - The pulse beep is suppressed during signs of motion when SmartTone is set to [Off].

10.7.10.5. Setting SpHb mode

While monitoring Hemoglobin levels, there are two blood sample sources from which Hemoglobin readings can be obtained: arterial and venous. To set the SpHb mode, follow these procedures:

- 1. Select the SpO_2 parameter area or waveform area to enter the $[SpO_2]$ menu.
- 2. Select [SpO₂ Setup] submenu.
- 3. Select [Arterial] or [Venous].

10.7.10.6. Setting SpHb Average

To set the SpHb average time, follow these procedures:

- 1. Select the SpO_2 parameter area or waveform area to enter the $\llbracket SpO_2 \rrbracket$ menu.
- 2. Select [SpO₂ Setup] submenu.
- 3. Set **SpHb Average Time**.

10.7.10.7. Setting SpHb Precision

To set the SpHb precision, follow these procedures:

- 1. Select the SpO_2 parameter area or waveform area to enter the $\P SpO_2$ menu.
- 2. Select 【SpO₂ Setup 】 submenu.
- 3. Set **SpHb Precision** as required.

10.7.10.8. Setting SpHb Unit

To set the SpHb unit, follow these procedures:

- 1. Select the SpO_2 parameter area or waveform area to enter the $\llbracket SpO_2 \rrbracket$ menu.
- 2. Select [SpO₂ Setup] submenu.

3. Set **SpHb Unit**.

10.7.10.9. Setting Waveform Mode

To set the SpHb unit, follow these procedures:

- 1. Select the SpO_2 parameter area or waveform area to enter the $[SpO_2]$ menu.
- 2. Select [SpO₂ Setup] submenu.
- Select [Wave Mode] as required, and select whether the SpO₂ waveform contains respiratory or not.

10.8. Setting PR

10.8.1. Setting PR Alarm

You can set PR alarm by following these steps:

- 1. Select the SpO_2 parameter area or waveform area to enter the $[SpO_2]$ menu.
- 2. Select [PR Alarm] submenu.
- 3. Set alarms as needed.

10.8.2. Setting QRS Volume

If the alarm source is set to PR, the QRS tone is derived from PR measurements. To set the QRS volume, follow this procedure:

- 1. Select the SpO_2 parameter area or waveform area to enter the $[SpO_2]$ menu.
- 2. Select [PR Setup] submenu.
- 3. Set **[QRS Volume]** to the appropriate value.

If the SpO_2 value is effective, the monitor also adjusts the QRS tone (Pitch tone) according to the SpO_2 value. For information, see 10.7.8 Setting Pitch Tone.

10.8.3. Setting PR Source

Current pulse source is displayed in the PR parameter area. The PR from current pulse source has the following characteristics:

PR is monitored as system pulse and generates alarms when you select PR as the active alarm source; PR is stored in the monitor's database and reviewed in the graphic/tabular trends; in trend graphs, as the PR curve is in the same color with that of the PR source, it is unlikely to distinguish the PR source.

To set which pulse rate as PR source, follow this procedure:

- 1. Select the SpO_2 parameter area or waveform area to enter the $[SpO_2]$ menu.
- 2. Select [PR Setup] submenu.
- 3. Select **[PR Source]**, and select a suitable PR source in the drop-down list.

The drop-down list of **【PR Source】** displays the currently valid PR source from top to bottom according to the priority level. When you select **【Auto】**, the system will automatically select the first option in the list as the PR source. If the PR source you set does not exist, the system will automatically switch **【PR Source】** to **【Auto】**. When you select **【IBP】**, the system will automatically use the first pressure label in the list as the PR source.

10.9. SpO₂ Troubleshooting

This section lists the problems that might occur. If you encounter the problems when using the equipment or accessories, check the table below before requesting for services. If the problem persists, contact your service personnel.



NOTE: For the physiological and technical alarm messages, see D Alarm

Message.

Problem	Corrective Actions	
Do not display SpO ₂ numeric area	1. Check that the SpO ₂ is set to display in the 【Screen	
or waveform area on the main	Layout menu. For more information, see 3.7.2 Setting	
screen.	Display Screen.	
	2. Check that if the SpO ₂ parameter switch is enabled. For	
	more information, see 3.7.1 Setting Parameters.	
	3. Check that the cable connections of SpO_2 sensor and the	
	extension cable are tight. Replace the SpO2 sensor or the	
	extension cable if needed.	

Dashes "" display in place of	1. Check that the cable connections of SpO ₂ sensor and the
numerics	extension cable are tight. Replace the SpO ₂ sensor or the
	extension cable if needed.
	2. Reconnect the SpO ₂ sensor if the alarm SpO₂ sensor Off
	appears.
	3. Check the PI value. If the PI value is too low, adjust the
	SpO ₂ sensor, or apply the sensor to the site with better
	perfusion.
	4. Move the sensor to the place with weaker light, or cover
	the sensor with shade.
Low amplitude SpO ₂ signal	1. The SpO ₂ sensor and NIBP cuff are placed on the same
	limb. Change a monitoring site if necessary.
	2. Check the PI value. If the PI value is too low, adjust the
	SpO ₂ sensor, or apply the sensor to the site with better
	perfusion.
	3. Check the sensor and its application site.
SpO ₂ value is inaccurate	1. Check the patient's vital signs.
	2. Check for conditions that may cause inaccurate SpO ₂
	readings. For more information, see 10.3 SpO ₂
	Measurement Limitations.
	3. Check the monitor, the MPS-P module, SpO ₂ module or
	if the function of sensor is normal.

Chapter 11 Temperature (TEMP)

11.1. Introduction

The thermistor is applied on continuous temperature measurement, which is based on the principle that electrical resistance of the thermistor changes as temperature changes. Thermistors measure the resistance change and use it to calculate the temperature.

The monitor supports at least 8 channels of temperature measurement. When measuring temperature on two different sites, can calculate the difference between two measured sites ($\triangle T$).

Temperature monitoring is intended for adult, pediatric and neonatal patients.

Measuring mode is direct mode.

11.2. Displaying the TEMP Parameter Area

To display the Temp parameters area, follow this procedure:

- 1. Enter **[Screen Layout]** interface in either of the following ways:
 - ◆ Select [Screen Setup] quick key→Select [Screen Layout] submenu.
 - ◆ Select【Main Menu]quick key→from【Display]column to select【Screen Layout].
- 2. Select you want to display the parameter area of the temperature parameters, and then from the popup list select 【TEMP】.

11.3. TEMP Display

The following figure shows the TEMP parameter area for temperature monitoring. Your display may be configured to look different.



- (1) Parameter label
- (2) TEMP unit

- (3) TEMP alarm limits: If TEMP alarm is turned off, the alarm closing icon is displayed here.
- (4) TEMP value
- (5) TEMP Difference (ΔT): TEMP Difference between two temperature sites. It displays only when ΔT is switched on.

11.4. Preparing for TEMP Monitoring

Please follow these steps to prepare TEMP measurement:

- 1. According to the type of patient and the measurement site, select the appropriate temperature probe.
- 2. Insert the probe or extension cable into the temperature probe connector. If a disposable probe is used, connect the probe and extension cable.
- 3. Refer to the instructions for using the probe and connect the probe to the patient.

11.5. TEMP Settings

11.5.1. Setting TEMP Alarm

To set the temperature alarm, follow this procedure:

- 1. Select the TEMP parameter area to enter the **【TEMP】** menu.
- 2. Select [Alarm] submenu.
- 3. If the alarm setting is protected by password, enter the password. For detail, please refer to 7.7.2 *Changing Alarm Setup Protection Mode*.
- 4. Set alarms as needed.

11.5.2. Setting TEMP Label

Select the temperature label according to the measurement site. To do so, follow this procedure:

- 1. Select the TEMP parameter area to enter the **【TEMP】** menu.
- 2. Select **Setup** submenu.
- 3. Set the TEMP label name according to the measurement site.

11.5.3. Displaying the Temperature Difference

To display the temperature difference between two measurement sites monitored by the same temperature module, switch on corresponding ΔT . To do so, follow this procedure:

- 1. Select the TEMP parameter area to enter the **【TEMP】** menu.
- 2. Select **Setup** submenu.
- 3. Switch on ΔT .

11.5.4. Setting TEMP Unit

You can change the unit of TEMP by following the steps below:

- 1. Select the TEMP parameter area to enter the **【TEMP】** menu.
- 2. Select [Setup] submenu.
- 3. Set TEMP [Unit].

11.6. TEMP Troubleshooting

This section lists the problems that might occur. If you encounter the problems when using the equipment or accessories, check the table below before requesting for services. If the problem persists, contact your service personnel.



NOTE: For the physiological and technical alarm messages, see D Alarm

Message.

Problem	Corrective Actions	
Do not display TEMP parameter area on the main screen.	1.	Check if the display of the TEMP parameter is set in 【Screen Setup】 menu.
	2.	Check that if the TEMP parameter switch is enabled. For more information, see <i>3.7.1 Setting Parameters</i> .
	3.	Check that if the connections of the temperature probe and the extension cable are tight.
Measurement fails / "" is	1.	If you are using a disposable probe, check whether

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displayed in the Temp parameter		the probe is tightly connected to the extension cable.
area.	2.	Try using a known good probe in case the sensor is
		damaged

Chapter 12 NIBP

12.1. Introduction

The monitor uses the oscillometric method for measuring the non-invasive blood pressure (NIBP). NIBP measurement is based on the principle that pulsatile blood flow through an artery creates oscillations of the arterial wall. The oscillometric device uses a blood pressure cuff to sense these oscillations that appear as tiny pulsations in cuff pressure. The oscillometric devices measure the amplitude of pressure changes in the occluding cuff as the cuff deflates from above systolic pressure. The amplitude suddenly increases as the pulse breaks through the occlusion in the artery. As the cuff pressure decreases further, the pulsations increase in amplitude, reach a maximum (which approximates to the mean pressure), and then diminish. The oscillometric method measures the mean pressure and determines the systolic and diastolic pressures.

NIBP monitoring is applicable for adult, pediatric, and neonatal patients.

The monitor supports BLT NIBP module and SunTech NIBP module.



NOTE:

- Blood pressure measurements determined with this device are equivalent to those obtained by a trained observer using the cuff/stethoscope auscultatory method or an intra-arterial blood pressure measurement device, within the limits prescribed by IEC 80601-2-30.
- NIBP measurement can be performed during electro-surgery and discharge of defibrillator.

12.2. Safety Information



WARNING:

Be sure to select the correct patient category setting for your patient before NIBP measurement. Do not apply the higher adult settings for pediatric or neonatal patients. Otherwise, it may present a safety hazard.

- Do not measure NIBP on patients with sickle-cell disease or on the limb where skin damage has occurred or is expected.
- Use clinical judgment to determine whether to perform frequent automatic blood pressure measurements on patients with severe blood clotting disorders because of the risk of hematoma in the limb fitted with the cuff.
- Do not use the NIBP cuff on a limb with an intravenous infusion or arterial catheter in place. This could cause tissue damage around the catheter when the infusion is slowed or blocked during cuff inflation.
- Do not apply cuff on the arm on the side of a mastectomy.
- Continuous cuff pressure due to connection tubing kinking may cause blood flow interference, and resulting in harmful injury to the patient.
- NIBP reading can be affected by the measurement site, the position of the patient, exercise, or the patient's physiologic condition. If you doubt the NIBP measurements, determine the patient's vital signs by alternative means, and then verify that the monitor is working correctly.
- In automatic or continuous measurement mode, if prolonged, cuff friction with limb may lead to purpura, ischemia, and neuropathy. In patient care, color, temperature, and sensitivity of distal extremities should be frequently examined. Check more frequently when making automatic or STAT measurements. If any abnormalities are observed, the site of the cuff should be changed or the NIBP measurement should be stopped. Auto NIBP measurements with one and two minute intervals are not recommended for extended periods of time.
- NIBP diagnostic significance must be decided by the physician.
- As the monitor uses an alternative small-bore connector design different from those specified in the ISO 80369 series, there is a possibility that a misconnection can occur between the monitor and a medical device using a different alternative small-bore connector, which can result in a hazardous situation causing harm to the patient. Special measures need be taken by the user to mitigate these reasonable foreseeable risks.



CAUTION:

- Only use parts and accessories specified in this manual. Follow the instructions for use and adhere to all warnings and cautions.
- Accuracy of NIBP measurement depends on using a cuff of proper size. It is essential to measure limb circumference and choose a cuff with proper size.
- NIBP automatically calibrated every time when the monitor is turned on. If it is not turned off for a long time or if the pressure is not accurate during use, you can use the "Reset" function in the NIBP menu to calibrate. You need to remove the cuff and windpipe before calibration, which in order to connect the NIBP pressure sensor to the atmosphere.

12.3. NIBP Measurement Limitations

Measurements are impossible with heart rate extremes of less than 40bpm or greater than 240bpm, or if the patient is on a heart-lung machine. The measurement may be inaccurate or impossible in the following situations:

- Regular arterial pressure pulses are hard to detect;
- With excessive and continuous patient movement such as shivering or convulsions:
- With cardiac arrhythmias;
- With rapid blood pressure changes;
- With severe shock or hypothermia that reduces blood flow to the peripheries;
- > On an edematous extremity;



NOTE:

The effectiveness of this sphygmomanometer has not been established in pregnant, including pre- eclamptic patients.

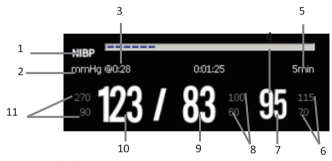
12.4. Measurement Modes

The monitor has the following NIBP measurement modes:

- Manual: Manually start a NIBP measurement.
- Auto: The monitor automatically and repeatedly performs NIBP measurements at set intervals.
- > STAT: With 5 minutes, the measurement is continuously performed, and then the monitor returns to the original mode.
- Sequence: The monitor measures automatically according to the set cycle length and interval.

12.5. NIBP Display

The NIBP display shows only numerics.



- (1) Parameter Label
- (2) NIBP Unit: mmHg or kPa
- (3) The last NIBP measurement time
- (4) Time to the next measurement (for Auto mode and Sequence only).
- (5) Measurement mode: The measurement interval time is displayed during Auto NIBP measurement, and the current measurement period and measurement interval time are displayed during Sequence measurement.
- (6) Mean pressure alarm limit
- (7) Mean pressure (displayed after measurement completed) or cuff pressure (displayed during the measurement)
- (8) Diastolic pressure alarm limit
- (9) Diastolic pressure
- (10) Systolic pressure
- (11) Systolic pressure alarm limit



- If NIBP measurement fails, "XX" is displayed; if NIBP measurement is not taken, "--" is displayed
- Outlined NIBP numerics indicate that the measurement exceeds the set time.
 So these NIBP values are not recommended for reference.

12.6. Preparing for NIBP Measurements

12.6.1. Preparing the Patient for NIBP Measurements

In normal use, perform NIBP measurement on a patient who is in the following position:

- Comfortably seated
- Legs uncrossed
- Feet flat on the floor
- Back and arm supported



NOTE:

- It is recommended that the patient calms down and relaxes as much as possible before performing the measurement and that the patient do not talk during the measurement.
- It is recommended to have the patient sit quietly for five minutes before taking the measurement.
- Other factors that have been shown to result in an overestimation of blood pressure are labored breathing, full bladder, pain etc.

12.6.2. Placing the NIBP Cuff

To place the NIBP cuff, follow this procedure:

1. Verify that the patient category setting is correct.

- 2. Connect the airpipe to the NIBP cuff connector of the device.
- 3. Select an appropriately sized cuff for the patient, and then wrap it around the limb directly over the patient's skin as follows:
 - a) Determine the patient's limb circumference.
 - b) Select an appropriate cuff by referring to the limb circumference marked on the cuff. The width of the cuff should be 40% (50% for neonates) of the limb circumference, or 2/3 of the length of the upper arm or the thigh. The inflatable part of the cuff should be long enough to encircle at least 50% to 80% of the limb.
 - c) Apply the cuff to the patient's upper arm or leg and make sure the Φ marking on the cuff matches the artery location. The cuff should fit snugly, but with enough room for two fingers to be placed between the cuff and the patient's arm (on adults), and loosely on neonates with little or no air present within the cuff. Otherwise it may cause discoloration and ischemia of the extremities. Make sure that the cuff index line falls within the range markings on the cuff.
 - d) Middle of the cuff should be at the level of the right atrium of the heart.
- 4. Connect the cuff to the air tubing. Avoid compression or restriction of pressure tubes. Air must pass unrestricted through the tubing.



CAUTION:

- A wrong cuff size and a folded or twisted bladder can cause inaccurate measurements.
- Do not touch or apply external pressure against the cuff and air tubing during NIBP measurement. This may cause inaccurate blood pressure values.
- Use care when placing the cuff on an extremity used for monitoring other patient parameters. Pressurization of the cuff may temporarily cause loss of function of simultaneously used monitoring equipment on the same limb.

12.7. Starting and Stopping NIBP Measurements

Start and stop NIBP measurement by selecting the NIBP quick key keys or from the NIBP menu:

Task	Via quick key	Via NIBP menu
Start a manual measurement	Select [NIBP Start/Stop] quick key	Select [Start]
NIBP Auto measurement	Select [NIBP Measure] quick key →Select interval time	Select 【Setup】 submenu→set 【Interval Time】
NIBP Sequence measurement	Select [NIBP Measure] quick key →Select [Sequence] →Select [NIBP Start/Stop] quick key	Select [Sequence] submenu—set NIBP Sequence measurement—Select [Start]
Start STAT measurement	Select [NIBP STAT] quick key	Select [STAT]
Stop the current NIBP measurements	Select [NIBP Start/Stop] quick key	Select [Stop]
End Auto NIBP or Sequence measurement	Select [NIBP Stop All] quick key	Select [Stop All]
Stop STAT measurement	Select [NIBP Start/Stop] quick key	Select [Stop] or [Stop All]

12.8. Viewing the Dynamic blood pressure analysis

Dynamic blood pressure analysis can intuitively understand the patient's blood pressure changes and distribution over a period of time. The steps of viewing the dynamic blood pressure analysis are as below:

- 1. Select the NIBP parameter area to enter the 【NIBP】 menu.
- 2. Select [Analysis] submenu.

12.9. NIBP Settings

12.9.1. Setting the NIBP Alarm

To set the NIBP alarm properties, follow this procedure:

- 1. Select the NIBP parameter area to enter the [NIBP] menu.
- 2. Select [Alarm] submenu.
- 3. If the alarm setting is protected by password, enter the password. For detail, please refer to 7.7.2 *Changing Alarm Setup Protection Mode*.
- 4. Set alarms as needed.

12.9.2. Setting the Initial Cuff Inflation Pressure

You can manually set the initial inflation pressure of the cuff, follow this procedure:

- 1. Select the NIBP parameter area to enter the **[NIBP]** menu.
- 2. Select **[Setup]** submenu.
- 3. Set 【Initial Pressure】: Select the appropriate cuff pressure value as needed.

12.9.3. Setting the NIBP Interval

For Auto NIBP measurement, you need to set the interval between two NIBP measurements. To set the NIBP interval, follow this procedure:

- 1. Select the NIBP parameter area or waveform area to enter the **[NIBP]** menu.
- 2. Select **Setup** submenu.
- 3. Set [Interval].

12.9.4. Selecting NIBP Start Mode

Start mode defines how NIBP auto mode works. To set the 【Start Mode】, follow this procedure:

- 1. Select the NIBP parameter area a to enter the [NIBP] menu
- 2. Select **Setup** submenu.
- 3. Set [Start Mode].
 - ♦ 【Clock】: After the first measurement, the monitor automatically

synchronizes NIBP automatic measurements with the real time clock. For example, if **【Interval】** is set to **【30min】**, and you start NIBP automeasurement at 10:03, the next measurement will be taken at 10:30, and then at 11:00, 11:30, and so on.

◆ 【Interval】: After the first measurement, the monitor automatically repeats measurements at set interval. For example, if 【Interval】 is set to 【30min】, and you start NIBP auto measurement at 10:03, the next measurement will be taken at 10:33, and then at 11:03, 11:33, and so on.

12.9.5. Enabling the NIBP End Tone

The monitor can issue a reminder tone at the completion of NIBP measurement. The NIBP End Tone is off by default. To switch on the NIBP end tone, follow this procedure:

- 1. Select the NIBP parameter area a to enter the **[NIBP]** menu
- 2. Select [Setup] submenu.
- 3. Switch on [NIBP End Tone].

12.9.6. Setting NIBP Sequence Measurement

NIBP sequence measurements can consist of up to 5 measurement periods: A, B, C, D, and E. You can set the measurement duration for each period and the interval between NIBP measurements in each period separately. The steps to set up the NIBP measurement sequence are as follows:

- 1. Select the NIBP parameter area a to enter the **[NIBP]** menu.
- 2. Select [Sequence] submenu.
- 3. Set each sequence measurement to **[Duration]** or **[Interval Time]** separately.

12.9.7. Setting NIBP Unit

You can change the NIBP units by following these steps:

- 1. Select the NIBP parameter area a to enter the **[NIBP]** menu.
- 2. Select **[Setup]** submenu.
- 3. Set NIBP [Unit].

12.9.8. Setting NIBP Invalid Time

NIBP measurements become outline fonts after a preset time. This avoids older NIBP values being misinterpreted as current measurements. To set the timeout period, follow this procedure:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】 →input maintenance password→Enter.
- 2. Select **Module** submenu→ **Other** submenu.
- 3. Set [NIBP Invalid Time].

12.9.9. Displaying the NIBP List

To display multiple sets of the latest NIBP measurements, follow this procedure:

- 1. Enter **[Screen Layout]** submenu by either of the following ways:
 - ◆ Select 【Screen Setup】 quick key→Select 【Screen Layout】 submenu.
 - ◆ Select 【Main Menu】 quick key→from 【Display】 column to select
 【Screen Layout】.
- 2. In the desired parameter area, select $\{NIBP\} \rightarrow \{NIBP List\}$.

12.10. Assisting Venous Puncture

You can use the NIBP cuff to cause sub-diastolic pressure to block the venous blood vessel and therefore help venous puncture. To assist venous puncture, follow this procedure:

- 1. Select the NIBP parameter area.
- 2. Select **[Setup]** submenu;
- 3. Set **[Venipuncture Pressure]**.
- 4. Select **[Venipuncture]** at the bottom of the menu.
- 5. Puncture vein and draw blood sample.
- 6. Select [NIBP Start/Stop] quick key or [Venipuncture] button to manually deflate the cuff. When performing a venipuncture, observe the inflation pressure and the remaining time of the venipuncture in the NIBP parameter area.

12.11. NIBP Maintenance

12.11.1. NIBP Leakage Test

The NIBP leakage test checks the integrity of the system and of the valve. The NIBP leakage test should be performed once every two years or when you doubt the NIBP measurements. The NIBP leakage test should be performed by qualified service personnel only.

12.11.2. NIBP Calibration

The NIBP accuracy test should be performed once every two years or when you doubt the NIBP measurements. The NIBP accuracy test should be performed by qualified service personnel only.

12.12. NIBP Troubleshooting

For more information, see D Alarm Message.

Chapter 13 IBP

13.1. Introduction

You can use P1/MPS-P module and IBP plug-in module to measure IBP. The monitor can provide 8 channels of IBP measurement results (use one P1/MPS-P module and 3 IBP plug-in modules). IBP measurement is applicable for adult, pediatric and neonatal. Pulmonary wedge pressure (PAWP) measurement is applicable for adult and pediatric.

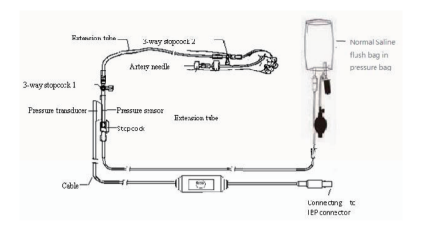
13.2. Safety information



WARNING:

- Use only IBP transducers specified in this manual. Never reuse disposable pressure transducers.
- The operator should avoid contact with conductive parts of the accessories when being connected or applied.
- 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.
- When using accessories, the operating temperature should be taken into consideration. For more information, see instructions for use of accessories.
- All invasive measurement involves risks to the patient. Use aseptic technique and perform according to manufacturer's instructions during measurement.
- Mechanical shock to the IBP sensor may cause severe shifts in zero balance and calibration, and cause erroneous readings.

13.3. IBP measurement



13.3.1. IBP Monitoring Procedure

Please make an IBP measurement following below steps:

- Plug one end of the IBP sensor cable into the monitor's IBP cable connector and the other end link to the IBP sensor.
- Refers to the IBP sensor manufacturer's instructions for exhausting air in the IBP sensor, it make ensure no air bubbles in the sensor's entire tube.
- Connecting IBP sensor to the patient, which makes sure the sensor and heart at the same level.
- 4. Selecting correct pressure label based on the measured pressure. Specifically, please refer to *13.5.2 Change the pressure label*.
- 5. Refers to *13.3.2 IBP Sensor Zero* for zeroing. During this process, the sensor keeps stationary and the valve is open to the atmosphere.



CAUTION:

- Before IBP measurements, it should make sure all IBP sensors are zeroed properly.
- Before IBP measurement, it makes sure no air bubbles in the IBP sensor which result in erroneous pressure readings.
- When intracranial pressure (ICP) measurements put on a sitting patient, the

sensor should be in line with the top of the patient's ear. Incorrect position can result in erroneous pressure readings.

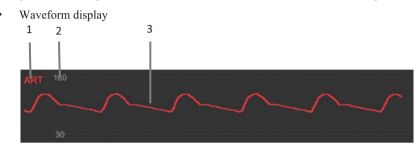
13.3.2. IBP sensor Zero

To obtain accurate pressure readings, the monitor requires a valid zero point. Zeroing the sensor at the hospital's specified frequency, and zeroing must be performed in the following cases:

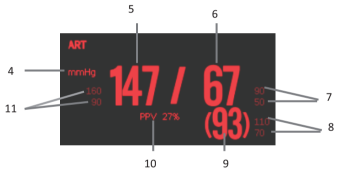
- Every time reconnecting IBP sensor and IBP sensor cable.
- The monitor needs restart.
- Suspecting the monitor's pressure reading is inaccurate.
- ◆ When the monitor displays the message 【Need to zeroing】, Please refer to the following steps to calibrating zero:
 - 1. Connecting IBP sensor, sensor cable and module.
 - Closing the 3-way stopcock (nearby the sensor end) to the patient's valve, and let the sensor pass through the 3-way stopcock to the atmosphere.
 - 3. Zeroing the sensor using one way from the following methods:
 - Select the parameter area of the pressure (e.g. ART) and select theZero button.
 - > Click **[Zero]** quick key-select **[IBP zero]** submenu-select the pressure to zeroing.
 - 4. After successful zero, close the valve to the atmosphere and open the valve to the patient. During zero process, when pressure fluctuation or the pressure exceeds the zero pressure range, it may fail. If fails, the processing method as follows:
 - > Check valve position of the 3-way stopcock near the sensor end for ensure access to the atmosphere.
 - Perform zero against. Do not shake the IBP sensor and tubing during zero calibration.

13.4. IBP Display

IBP measurements display the waveforms of pressure and pressure values on the screen. For arterial pressure, IBP parameter area displays systolic pressure, diastolic pressure and mean pressure. For venous pressure, the IBP numeric area displays only the mean pressure. The figure below shows the waveform and numerics for the Art pressure.



Parameter Display



- (1) IBP label
- (2) Waveform scale
- (3) Waveform
- (4) Pressure unit: mmHg, kPa or cmH₂O
- (5) Systolic pressure
- (6) Diastolic pressure
- (7) Diastolic pressure alarm limit
- (8) Mean pressure alarm limit
- (9) Mean pressure
- (10) PPV measurement value
- (11) Systolic pressure alarm limit

13.5. Setting IBP

13.5.1. Setting the IBP alarm

You can set the alarm by following the steps below:

- 1. Select the IBP parameter area or waveform area to enter the IBP menu.
- Select [Alarm] submenu.
 If the alarm setting is protected by password, enter the password. For detail, please refer to 7.7.2 Changing Alarm Setup Protection Mode.
- 3. Set alarms as needed.

13.5.2. Change the pressure label

The pressure label is identifier for each type only, so the pressure label must be set up when making pressure measurements. You can choose a pressure label following these steps:

- Selecting IBP parameter area or waveform area where you need to change labels for enter corresponding IBP menu.
- 2. Select **Setup** submenu.
- 3. Select **[Label]** where the appropriate tag name in the list.

Label name	Description	Label name	Description
PA	Pulmonary artery pressure	CVP	Central venous pressure
Ao	Aortic pressure	LAP	Left atrial pressure
UAP	Umbilical arterial pressure	RAP	Right atrial pressure
BAP	Brachial arterial pressure	ICP	Intracranial pressure
FAP	Femoral arterial pressure	UVP	Umbilical venous pressure
ART	Arterial blood pressure	LV	Left ventricular pressure
P1 to P2	Non-specific pressure label		



NOTE:

■ The same label name cannot be used for different channels about IBP

13.5.3. Setting up display types about Extended Pressure

If current pressure label is set to the extended pressure (P1 or P2), you could select the type which displays in the parameter area, following these steps:

- Select the parameter area or waveform area about the extended pressure for entering the corresponding pressure menu.
- 2. Select **[Setup]** submenu.

3. Set [Measurement]:

- All: Corresponding pressure in the parameter area shows all the pressure: systolic pressure, diastolic pressure and mean pressure.
- ➤ **Mean only**: Corresponding pressure in the parameter area only shows the average pressure.
- ➤ **Auto:** The system will automatically shows the pressure is displayed in the parameter area or only the average pressure according to the measured value of the extended pressure.

13.5.4. Setting the pressure sensitivity

The blood pressure value displayed on the monitor is average calculation about the collected data over a period time. The higher the sensitivity, the faster the monitor responds when the patient's blood pressure value changes, but the measurement accuracy is lower. Inversely, the lower the sensitivity, the slower the response of the monitor when the patient's blood pressure value changes, but the measurement accuracy is higher. When monitoring critically ill patients, setting up a higher sensitivity is useful for timely analysis.

You can set up the sensitivity of the current pressure, following these steps:

- Select the IBP parameter area or waveform area for enter the corresponding pressure menu.
- 2. Select [Setup] submenu.
- 3. Set [Sensitivity].

13.5.5. Setting the IBP Waveform

You could set up the IBP waveform, following these steps:

- Select the IBP parameter area or waveform area to enter the corresponding pressure menu.
- 2. Select **[Setup]** submenu.
- 3. Make the following settings for the IBP waveform:
 - > Set [Speed].
 - > Set **[Scale Type]**: If **[Auto]** is selected, the upper and lower scales of the IBP waveform will be automatically adjusted as the waveform amplitude changes.

13.5.6. Setting the PA-D instead of PAWP switch

You can select whether use PA-D value instead of PAWP value for hemodynamic calculation, and the methods are as below:

- 1. Select the PA parameter area or waveform area to enter the **PA** menu.
- 2. Select **[Setup]** submenu.
- 3. Set [Use PA-D as PAWP] to [ON] or [OFF].

See 25.4 Hemodynamic Calculations for details.

13.5.7. Turn on PPV measurement

PPV is the pulse pressure variation. When measuring arterial pressure (excluding PA), you can turn on PPV measurements, following these steps:

- Select the IBP parameter area or waveform area to enter the corresponding pressure menu.
- 2. Select [PPV] submenu.
- 3. Set [PPV Measurement] to [ON].

When **[PPV Measurement]** is setting to **[ON]**, the source of the PPV can be selected.



WARNING:

■ The monitor will calculate the PPV based on any arterial pressure value between heartbeats. The conditions of PPV measurement, and whether the

PPV numerical calculation has clinical significance or not, it is applicable or not. It must be judged by a doctor.

- Only a doctor can determine the clinical value of PPV information. According to recent scientific literature, the clinical relevance of PPV information is limited to controlled mechanical ventilation and to sedated patients without arrhythmias.
- The calculated PPV value may not be accurate under the following conditions:
 - a) Respiration rate is less than 8 rpm
 - b) During venting, the tidal volume is less than 8ml/kg
 - c) The patient has acute right ventricular dysfunction (i.e., pulmonary heart disease)
 - d) PPV measurements are only validated for adult patients

13.5.8. Changing the pressure unit:

You can change the unit of pressure by following these steps:

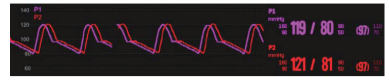
- Select the IBP parameter area or waveform area to enter the corresponding pressure menu.
- 2. Select [Setup] submenu.
- 3. Set IBP [Unit] if needed.

13.5.9. Overlapping IBP Waveforms

Please following steps below, set up the IBP waveform overlay display:

- 1. Enter the **[Screen Layout]** page, following method:
 - ◆ Select 【Screen Setup】 quick key → select 【Screen Layout】
 submenu
 - ◆ Select 【Main Menu】 quick key → select 【Screen Layout】 from the 【Display】 column.
- 2. Select 【IBP Overlap】 in the waveform parameter area and select the IBP waveform to be overlapped on the same line on the left side.
- Repeat the operation of step 2 at other locations with waveform parameter areas if needed.

Select
 ■ to exit Setup page. The overlapped IBP waveform can be displayed
 on the main interface.



You can open the 【IBP Overlap】 menu by selecting the IBP waveform area to be overlapped on the main screen. In the 【IBP Overlap】 menu, you can make the following settings:

- ◆ Scale
 - > Set the **[Left Scale]** for arterial pressure.
 - > Set the **[Right Scale]** for venous pressure.
- Set the **Grid** of the overlapped waveform area.
- ◆ Set the **【Wave Speed】** of the overlapped display waveform.

13.6. Calculating Cerebral Perfusion Pressure

The monitor can calculate the difference between mean arterial pressure (ART) and the intracranial pressure (ICP). The difference is cerebral perfusion pressure, which is labeled CPP. Therefore, the CPP value will be displayed on the screen only when the ART and ICP are displayed at the same time.

13.7. Measuring PAWP

Float a floating catheter with a ballon-tipped through the blood flow and wedge it into the small pulmonary artery to block the forward blood flow. At this time, the pressure measured at the tip of the cathether is the Pulmonary wedge pressure (PAWP). PAWP value is used to assess heart function, it mainly accepts the influence of fluid state, myocardial contractility and the integrity of valves and pulmonary circulation.

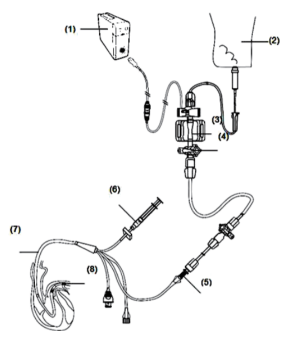
PAWP can reflect the changes in chest pressure during the entire respiration cycle. When the airway pressure and valve function are normal, PAWP is the end-diastolic pressure of the left ventricle. Therefore, the PAWP value measured at the end of the respiration cycle is the most accurate. At this time, the pressure in the chest is relatively constant, making the artifacts caused by respiration.



WARNING:

■ PAWP monitoring is not applicable for neonatal patients.

13.7.1. PAWP Device Connection



- (1) MPS-P/P1/IBP module
- (2) Flush bag
- (3) IBP sensor
- (4) 3-way valve
- (5) PA distal port
- (6) Balloon inflation valve
- (7) Floating catheter
- (8) Balloon

13.7.2. Preparing for monitoring PAWP

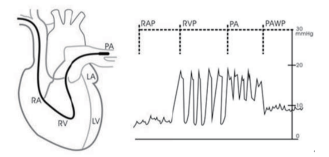
Please refer to the following steps to prepare PAWP measurement:

- Connect the IBP sensor, IBP cable and the module. For more information, see 13.3.1 IBP Monitoring Procedure.
- Connect the PA port of the floating catheter and the patient end of the IBP sensor following the manufacturer's instructions.
- 3. Zero the IBP sensor. For more information, see *13.3.2 IBP sensor Zero*.
- 4. Set the IBP label to **PA** since the PAWP is measured on PA. For more information, see *13.5.2 Change the pressure label*.

13.7.3. PAWP Monitoring Procedure

Please refer to the following steps to measure PAWP:

- Select the PA parameter area or waveform area on the main screen, and then select 【PAWP】 to enter the corresponding menu.
- 2. See the figure below, wedge the tip of floating catheter into the pulmonary artery by observing the PA waveform changes on the screen.



- 3. Select [Start].
- 4. Inflate the balloon and pay attention to PA waveform changes on the screen when the prompt message **【Ready for Balloon Inflation】** appears.
- 5. Deflate the balloon when the prompt message 【Ready for Balloon Deflation】 appears. If the PAWP waveform is stable yet the monitor still not show the prompt message 【Ready for Balloon Deflation】, select the 【Freeze】 to freeze the waveform, and deflate the balloon.

- 6. Select **(Accept)** to save the PAWP value.
- 7. If you need to start a new measurement, repeat the step 3 to step 6.

If the measurement fails or need to adjust the PAWP value, use the following buttons to adjust the PAWP waveform and values.

- Select the up or down arrow button to adjust the PAWP values.
- Select the left or right arrow button to view the frozen waveforms of 120 seconds.
- Select **(Accept)** to save the PAWP value.



WARNING:

- Prolonged inflation can cause pulmonary hemorrhage, infarction or both. Inflate the balloon for the minimum time necessary to get an accurate measurement.
- If the PAWP is greater than the PA (systolic), deflate the balloon and report the incident in accordance with hospital policy. Because the pulmonary artery could be accidentally ruptured, and the PAWP value obtained will not reflect the patient's hemodynamic state, but will merely reflect the pressure in the catheter or balloon.
- If the floating catheter inserts into the wedge position without inflation of the balloon, the PA waveform assumes a wedged appearance. Table appropriate action, in accord with standard procedures, to correct this situation.



NOTE:

The PA alarm is turned off automatically when entering the PAWP screen.

13.7.4. Setting the PAWP

Select the **【Setup】** on the PAWP screen to enter the **【PAWP setup】** menu. In the **【PAWP setup】** menu, you can make the following settings:

- Select 【Reference Waveform 1】 to set the ECG reference wave.
- Select 【Reference Waveform 2】 to set the RESP reference wave.
- Select **Speed** to set all waveform speed on PAWP screen.
- Select **Scale** to set the scale of PA waveform on PAWP screen.

13.7.5. Hemodynamic Calculation

Select **【Hemo Calcs 】** on the **【PAWP】** screen to enter **【Calculations】** menu. For more information, see 25.4 Hemodynamic Calculations.

13.8. IBP troubleshooting

This section describes problems you may encounter during using. You can refer to the following table for troubleshooting. If the problem persists, please contact maintenance staff.



NOTE: See D Alarm Message for physiological alarms and technical alarm

information.

Problem	Solution		
IBP parameter area and waveform	1. Check whether the display of IBP		
area cannot be found on the	parameters is set in the 【Screen Layout】		
interface	menu or not. For details, see 3.7.1 Setting		
	Parameters to be protected.		
	2. Check whether the IBP parameter switch is		
	turned on or not. For details, see 3.7.1 Setting		
	Parameters		
	3. Check the IBP cable, IBP sensor and		
	module are connected or not.		
	4. Check the valve position of the 3-way		
	stopcock is correct or not.		
	5. Confirm that the sensor has been zeroed.		

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	For details, see 13.3.2 IBP Sensor zero.
P1/P2 does not display systolic and	Set the displayed pressure to [All]. For details,
diastolic pressure measurements	see 13.5.3 Setting up display types about
	Extended Pressure.
IBP reading is unstable	1. Verify no air bubbles in the IBP sensor
	system.
	2. Check if the sensor is fixed.
	3. Perform zero against.
	4. Replace the sensor.
Zero failure	1. Check if the pipeline of the IBP sensor is
	open to the atmosphere.
	2. Perform zero against. Do not shake the IBP
	sensor and tubing during zeroing. For details,
	see 13.3.2 IBP Sensor zero.
	3. If zero still fails, replace the sensor.

Chapter 14 Impedance Cardiography (ICG)

14.1. Introduction

Impedance cardiography (ICG) is a safe, non-invasive method based on thoracic electrical bioimpedance (TEB) technology, which measures the level of change in impedance in the thoracic fluid, and generates ICG waveform, and calculates hemodynamic parameters.

Apply ICG monitoring only to patients in height of 120 to 230cm, weight of 30 to 155kg, and in age no less than 13.

14.2. Safety Information



WARNING:

- Do not allow patients to wear ICG sensors when undergoing Magnetic Resonance Imaging (MRI) due to risk of severe burns.
- The ICG sensor can only be applied to a patient at one time.
- The ICG module is not intended to be used while exposing the patient to high frequency current.
- Simultaneous use of high frequency electrosurgical equipment (ESU) during ICG monitoring may result in burns at the stimulation site and can also adversely affect measurement accuracy. Make sure the ESU return electrode is properly applied to the patient.



CAUTION:

Ensure that the sensor gel, sensors or lead wires do not contact any other conductive materials (including earth-grounded materials) during patient monitoring.

14.3. ICG Measurement Limitation

Factors which can affect the accuracy of ICG measurement, as following:

- Septic shock;
- Aortic valve regurgitation and defect of septum;
- Severe aortic sclerosis, aortic prosthesis;
- Severe hypertension (Mean>130 mmHg);
- Cardiac arrhythmia;
- Tachycardia with a heart rate higher than 200 bpm;
- The patient's weight and height are out of range: Patient heights below 120 cm (48") or above 230 cm (90"); Patient weights less than 30 kg (67 lbs.) or greater than 155 kg (341 lbs.);
- Aortic balloon or aortic balloon pump;
- 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.
- Simultaneous use of electrical cautery systems during surgical procedures;



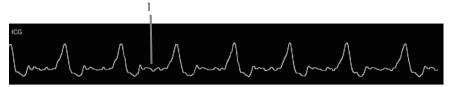
CAUTION:

■ The patient should maintain a static posture during ICG measurement. The measured ICG parameters can only be used when the quality of the ICG waveform signal is good enough and there are no artifacts.

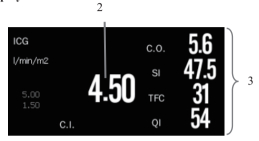
14.4. ICG Display

The ICG parameter area displays the value of a primary parameter and up to 4 secondary parameters. You can select the parameter for display in the **[Select Parameter]** page of the **[ICG]** menu. See *14.7.4 Selecting ICG Parameter* for details.

♦ Waveform Display



Parameter Display



- (1) ICG waveform
- (2) Primary parameter
- (3) Secondary parameter

14.5. Preparing for ICG Measurements

To prepare to monitor ICG, follow this procedure:

- 1. Prepare the patient's skin. For more information, see 14.5.1 Skin Preparation.
- 2. Place the ICG sensors on the patient. For more information, see *14.5.2 Placing the ICG Sensors*.
- 3. Connect one end of the patient cable to the ICG module.
- Connect the electrode wires of the patient cable to the sensors on the patient by matching the right and left electrode wire colors and numbers. For more information, see 14.5.3 Connecting ICG Cable.
- Input the patient information. For more information, see 14.7.2 Setting Patient Information.



WARNING:

Before monitoring patients with pacemakers, ensure that the function of the pacemaker cannot be influenced by the measuring current used for impedance cardiography. In the case of minute ventilation pacemakers the use of the ICG module is not allowed if the minute ventilation function of the pacemaker is activated.

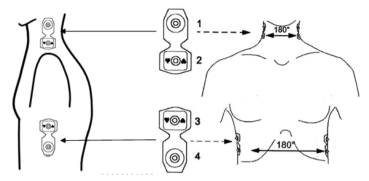
14.5.1. Skin Preparation

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 steps:

- 1. Select sites with intact skin, without impairment of any kind.
- 2. Clip or shave hair from sites as necessary.
- Gently abrade the skin and remove dead skin cells to improve the conductivity of the sensor site.
- 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.

14.5.2. Placing the ICG Sensors

In order to get good signals and accurate data, it is important to place ICG sensors on appropriate position, shown as follows:



To place the sensors:

- 1. A pair of sensors shall be placed below the earlobe on each side of the neck.
- 2. Another pair of sensors shall be placed on the midaxillary line at the xiphoid process level.

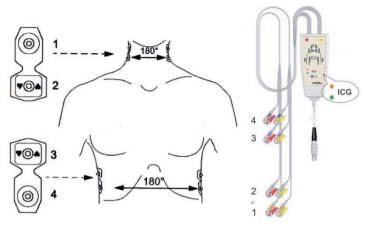


WARNING:

- The two sensors must be placed right on the opposite position (180°).
- The sensors must not have a direct contact to other electrically conductive materials.
- Do not reuse the disposable ICG sensor.

14.5.3. Connecting ICG Cable.

The ICG patient cable is used to connect the ICG module and the sensors on the patient. The left electrode wires (yellow) and right electrode wires (red) should be connected with the patient sensors by matching the number. See 14.5.2 Placing the ICG Sensors.



There is a signal indicating box on the patient cable. Two LED indicators (green and orange) on the signal indicating box are used to indicate the current status of the patient cable.

Green	Orange	Status description
	0	Measurement is running; sensor contact is good.
0	0	The patient cable is disconnected from the module or the monitor is turned off.
\Rightarrow	0	The patient cable is ready to use, but the measurement has not yet started.
0	Ď.	The patient cable is ready, but the module is in preparation.
•	•	Insufficient contact between sensors and patient: at least one lead wire is disconnected or not properly fixed; sensors are too dry (new sensors are necessary)

Note: ○ LED off

LED flashing

LED on

14.6. ICG Parameter

Select ICG parameter area, and select **[Hemodyna Param.]** in the ICG parameter setting menu, you can see the current hemodynamic parameters of patient, as following list:

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·m ² /cm ⁵	systemic vascular resistance index
TFI	Ω	thoracic fluid index
TFC	/kΩ	thoracic fluid content
HR	bpm	heart rate
DO ₂ I	mL/min/m ²	oxygen delivery index

Note:

- The HR reading is not obtained from ECG module, and it is obtained from ICG module.
- b) The output value is greater than the normal upper limit is indicated by an up arrow "↑"; the output value is lower than the normal lower limit is indicated by a down arrow "↓".
- c) Select [Record], the current page will be printed out.

14.7. ICG Setting

14.7.1. Setting the ICG Alarm

You can set the ICG alarm through following steps:

- 1. Select the ICG parameter area or waveform area to enter the ICG menu.
- 2. Select [Alarm] submenu.
 - If the alarm setting is protected by password, enter the password. For detail, please refer to 7.7.2 Changing Alarm Setup Protection Mode.
- 3. Set alarms as needed.

14.7.2. Setting Patient Information

You can set the patient information by following steps below:

- 1. Select the ICG parameter area or waveform area to enter the ICG menu.
- 2. Select **[Setup]** submenu.
- 3. Set [Height], [Weight], [Gender], [Age] and [Paced] of the patient. The patient information here is linked with the patient information in the system. When the above 5 information is modified, the patient information in the system will also be modified accordingly. Similarly, if the above 5 information in the system are changed, the corresponding information here will also be changed accordingly.
- Input the measurement values of 【SYS】, 【DIA】, 【MAP】, 【CVP】,
 【PAWP】, 【HB】 and 【SpO₂】. Each parameter has a default value, and the user can change it according to the actual situation.
- 5. Select 【Get Parameters】 below the menu, the system can obtain current monitoring parameter datas automatically. When required parameter value cannot obtain, you need to input it manually. For example, SYS, DIA and MAP can obtain through the measurement of IBP module (ART, P1 or P2). When SYS, DIA and MAP cannot obtain from IBP module, it can also be obtained from NIBP module. When these data cannot obtain from module, should input manually.

14.7.3. Setting the ICG waveform speed

You can set ICG waveform speed through following steps:

- 1. Select the ICG parameter area or waveform area to enter the ICG menu.
- 2. Select **[Setup]** submenu.
- 3. Set [Wave Speed].

14.7.4. Selecting ICG parameter

The ICG numeric area displays one primary parameter (C.I. by default) and four secondary parameters (SVRI, SVI, C.O. and TFC by default). You can also select your

desired primary and secondary parameters for display.

- 1. Select the ICG parameter area or waveform area to enter the **【ICG】** menu.
- 2. Select **[Setup Parameters]** submenu.
- 3. Select parameters to be displayed.

14.8. ICG Troubleshooting

This section lists the problems that might occur. If you encounter the problems when using the equipment or accessories, check the table below before requesting for services. If the problem persists, contact your service personnel.



NOTE: For the physiological and technical alarm messages, see D Alarm

Message.

Problem	Corrective Actions
Do not see ICG numeric area on the main screen.	 Check that the ICG is set to display in the 【Screen Layout】 menu. For details, see 3.7.2 Setting Display Screen. Check whether the ICG parameter switch is on or not. For details, see 3.7.1 Setting Parameters.
	3. Check if the patient type is correct.4. Check whether the ICG cable, ICG sensor and module are connected or not.

Chapter 15 Carbon Dioxide (CO₂)

15.1. Introduction

The monitor adopts infrared absorption technology to measure the carbon dioxide (CO₂) concentration in the breathing airway of patient. Because CO₂ molecule can absorb infrared light of special wavelength, and the amount of absorbed infrared light directly relates to the concentration of CO₂, therefore while the infrared light radiated from the infrared light source passing through the gas sample containing CO₂, part of energy will be absorbed by CO₂ in the gas. At another side of infrared light source, a photodetector is used to measure the remaining infrared energy and convert it to electric signal, which will be compared with the energy of infrared light source and adjusted so as to correctly reflect the CO₂ concentration in the gas sample.

There are two methods for measuring carbon dioxide in the patient's airway:

- 1. Mainstream: Uses a CO₂ sensor attached to an airway adapter directly inserted into the patient's breathing system.
- 2. Sidestream/Microflow: Takes a sample of the respiratory gas with a constant sample flow from the patient's airway and analyzes it with the CO₂ sensor.

If you use AG module to measure CO₂, see *Chapter 16 Anesthetic Gas (AG)*. CO₂ measurement is applicable for adult, pediatric and neonatal.

15.2. Safety information



WARNING:

■ When placing pipes such as sampling tubes, prevent the pipes from suffocating the patient's throat.



CAUTION:

- When the patient is being treated with aerosolized drugs, the measured EtCO₂ value may be inaccurate and is not recommended for use in this situation.
- The EtCO₂ value measured by the CO₂ module may differ from the CO₂ partial

pressure value measured by blood gas analysis.

■ The CO₂ module has an automatic alarm suppression function, and the CO₂ module performs a physiological alarm only after the respiratory wave is detected. When monitoring the patient with the CO₂ module, make sure the device is properly connected to the patient.

15.3. CO₂ Measurement Limitations

The following factors may affect the accuracy of the measurement:

- ◆ Leaks or internal venting of sampled gas;
- ◆ Mechanical shock;
- Cyclical pressure of up to 10kPa (100cmH₂O) and abnormal pressure change of the gas path;
- Other sources of interference (if any).

Measurement accuracy of the sidestream/microflow CO₂ module may be affected by the breath rate and inspiration/ expiration (I/E) ratio as follows:

- ◆ EtCO₂ value is within specification for breath rate≤60rpm and I/E ratio≤ 1:1.
- ◆ EtCO₂ value is within specification for breath rate≤30rpm and I/E ratio≤ 2:1.

15.4. Monitoring Procedure

15.4.1. Measure Using Mainstream CO₂ Module

- Attaching the CO₂ sensor cable
 Plug the cable of CO₂ sensor into CO₂ connector on the CO₂ plug-in module.
- 2. Selecting a proper 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
Pediatric/Adult (Disposable)	>4.0mm
Adult (Reusable)	>4.0mm
Neonatal/Pediatric (Disposable)	≤4.0mm

Neonatal (Reusable)	≤4.0mm

3. Attaching the airway adapter to the CO₂ sensor

Before attaching the airway adapter to the CO₂ 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.

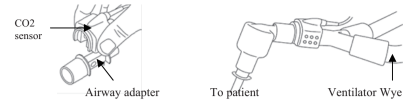
The monitor will display the "Sensor Warm Up..." message while the sensor and adapter warm to operating temperature. The message disappears when the sensor is ready for use.

Since the Masimo IRMA CO₂ mainstream module has a faster heating rate, there is no "Sensor Warm Up..." prompt.



CAUTION:

- The atmospheric pressure must be set to the correct value before using the mainstream CO₂ module. Incorrect atmospheric pressure settings can result in erroneous CO₂ readings (Masimo IRMA CO₂ mainstream modules do not function because they already have automatic atmospheric pressure compensation).
- Install the sensor above the adapter to avoid liquid build-up on the adapter window. Gas concentration at high concentrations at this location can hinder gas analysis.
- To avoid dead space, place the sensor and adapter as close as possible to the patient.
 - 4. Perform a zero, the details refer to 15.8 Zeroing.
 - After the zero is finished, take the Masimo IRMA CO₂ module as an example and connect the air circuit as shown below.



6. Check before use:

- ① Before connecting the airway adapter to the breathing circuit, check if the readings on the monitor are correct.
- ② Before connecting the airway adapter to the patient circuit, verify the gas reading and waveform through the display on the monitor.
- 3 After installing the mainstream CO₂ sensor on the airway adapter, please check the tightness of the patient circuit.
- ♦ Status indicated by LED on Masimo IRMA CO₂ sensor:

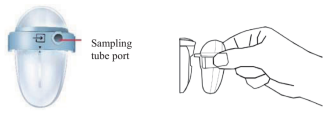


LED	The indicated status
Steady green light	System OK
Blinking green light	Zeroing in progress
Blinking red light	Check adapter

15.4.2. Measure Using Sidestream CO₂ module

Take Sidestream CO₂ plug-in module with water trap as an example. Please refer following steps to use sidestream CO₂ module to measure:

1. As shown below, install the water trap on the water trap holder of the sidestream CO₂ plug-in module, and connect one end of the sampling tube to the water trap.



2. Install another end of the sampling tube to patient.



CAUTION:

- Pay attention to the water level of water trap. If the highest water level reaches, please replace the water trap in time to prevent the module from entering by water.
- Please keep the sampling tube clean, and prevent the tube from clogging by dust.
- When using a sidestream CO₂ module, the exhaust holes on the module must be connected to the exhaust gas treatment system.

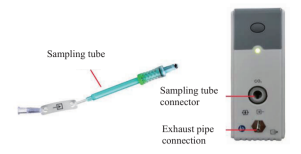


NOTE:

- Inserting the CO₂ module into slot, the monitor automatically starts the sampling pump. Removing the CO₂ module can turn the sample pump off.
- Water trap and sampling tubes are disposable, please use products provided or designated by manufacturer.

15.4.3. Measure Using Microflow CO₂ module

Connect one end of the sampling tube to the CO₂ plug-in module. If need to use
with CO₂ sensor, shall first connect CO₂ sensor cable, and then connect the catheter,
airway adapter or sampling tube to the sensor as required. Take Nomoline ISA CO₂
module as an example, as shown in the following figure:





NOTE:

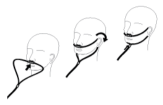
- Inserting the sampling tube into the sampling tube connector will automatically start the sampling pump. After the sampling tube is pulled out, the sampling pump stops.
- 2. If the sampling pump fails to turn on, or runs intermittently, perform a "Zero" procedure. (Refer to the *15.8 Zeroing*)
- Ensure that the CO₂ sensor exhaust tube vents gases away from the sensor environment.
- 4. Wait for the CO₂ sensor to warm up. The monitor will display the "Sensor Warm Up..." message for approximately 1minute while the sensor warms up to operating temperature. The message disappears when the sensor is ready for use.
- 5. Applying 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. Shown as follows:



2) 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. Shown as follows:



3) For non-intubated patients: Place the nasal cannula onto the patient. Shown as follows:



- 4) For patients prone to mouth breathing, use an oral-nasal cannula. Can be used as shown below:
 - > 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. Shown as follows:



When use the Nasal/Oral Cannula of Masimo ISA Capno, the cannula can be worn as shown below:



5) 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.



CAUTION:

- 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.
- Always disconnect the cannula, airway adapter or sampling tube from the CO₂ sensor when not in use.
- Pull the sampling tube out of the sampling tube connector when not in use.
- Do not insert the things other than sampling tube into receptacle of sampling tube.
- The sampling tubes are disposable. Please keep the sampling tube clean, and prevent the tube from clogging by dust. Under the conditions that the sampling gas temperature is 37°C, the indoor temperature is 23°C, and the sampling relative humidity is 100%, replace the sampling tube at least every 12 hours (if filter tip is used, it can be extended to 120 hours), or replace the sampling tube when the sampling tube is found to be leaking, damaged or contaminated.
- When using a microflow CO₂ module, the exhaust holes on the module must be connected to the exhaust gas treatment system.
- 6. Pre-use inspection (pre-use inspection must be performed before connecting the sampling tube to the patient's breathing circuit):
- ① Insert the sampling tube into the air inlet on the module.
- ② Check whether the LED aperture on the module is green and always bright. (a green light indicates that the system is normal)
- 3 Exhale toward the sampling tube and check whether CO₂ waveform and value are displayed on the monitor.
- ④ Block the sampling tube with your fingers and wait for 10s.
- 6 Check whether the blockage alarm is displayed, and whether the LED aperture on the module flashed red at the same time.
- ⑥ Under appropriate circumstances, check the tightness of the patient's breathing circuit connected with the sampling tube.

♦ Status indicated by LED aperture on Nomoline ISA CO₂ module:



LED aperture	The indicated status
Steady green light	System OK
Blinking green light	Zeroing in progress
Steady red light	Sensor error
Blinking red light	Check sampling tube

15.5. Display

♦ Waveform Display



♦ Parameter Display



5

- (1) CO₂ waveform
- (2) Parameter label
- (3) Unit of CO₂
- (4) Alarm limit of EtCO₂

- (5) EtCO₂ value
- (6) Inspired minimum CO₂ (FiCO₂)
- (7) Airway respiration rate (awRR)

15.6. Settings CO₂

15.6.1. Setting the CO₂ Alarm

- 1. Select the CO₂ parameter area or waveform area to enter the **CO₂** menu.
- 2. Select [Alarm] submenu.
 - ➤ If the alarm setting is protected by password, enter the password. For detail, please refer to 7.7.2 Changing Alarm Setup Protection Mode.
- Set alarms as needed.

15.6.2. Settings Apnea Alarm Time

You can set the apnea alarm time by following the steps below. The monitor will trigger the alarm when the patient's suffocation time exceeds the set time.

- 1. Select the CO₂ parameter area or waveform area to enter the **[CO₂]** menu.
- 2. Select [Alarm] submenu.
- 3. Set [Apnea Delay].



WARNING:

■ It is cannot judge the cause of respiratory apnea through CO₂ monitoring. If the monitor cannot detect the patient's breath again from the moment of the last breath to the pre-set time, the monitor only provides the alarm of respiratory apnea. Therefore, the alarm of respiratory apnea should not be used for the diagnosis of the patient.

15.6.3. Changing the CO₂ Unit

To change the CO₂ unit, follow this procedure:

- 1. Select the CO_2 parameter area or waveform area to enter the CO_2 menu.
- 2. Select [Setup] submenu.
- 3. Set CO₂ 【Unit】.

15.6.4. Setting the CO₂ Waveform

To set the CO₂ waveform, follow this procedure:

- 1. Select the CO₂ parameter area or waveform area to enter the 【CO₂】 menu.
- 2. Select **[Setup]** submenu.
- 3. Set CO₂ waveform [Wave Mode], [Wave Speed] or [Scale].

15.6.5. Setting CO₂ Corrections

Temperature, water vapor in the patient's breath, barometric pressure, and the proportions of O_2 , N_2O and Helium in the mixture all influence CO_2 absorption.

For mainstream and sidestream/microflow CO₂ module, you can set the CO₂ correction by following the steps below:

- 1. Select the CO₂ parameter area or waveform area to enter the **【CO₂】** menu.
- 2. Select **Setup** submenu.
- 3. Please set the following contents according to the actual situation before correction:
 - 【Gas Temperature】: Set the temperature of gas.
 - 【Barometric Pressure】: Set the atmospheric pressure.

The system default barometric pressure is 760mmHg, you can select [Main Menu] quick key—from [System] column to select [Maintenance] —in [Other] submenu to enter the barometric pressure value of the environment.

- 【Zero Gas Type 】: Select the gas type of zeroing, the options are 【Zero On Room Air 】 or 【Zero On N_2 】.
- $[O_2 \ Compensation]$: Select the concentration of oxygen. It can be set to a value between 0% and 100%. The default value is 16%.
- ——【Anesthetic Gas】: Select the concentration of anesthetic agent. It can be set to a value between 0.0% and 20.0%. The default value is 0.0%.
- 【Balance Gas】: Select the type of balance gas, the options are 【Room Air】, 【 N_2O 】 or 【Helium】.

When the most proportions of the mixture is air, select [Room Air] When the most proportions of the mixture is N_2O , select [N_2O]. When the most proportions of the

mixture is Helium, select [Helium].

The Masimo mainstream/sidestream CO₂ module supports automatic air pressure compensation and automatic temperature compensation. Manual setting is not required under normal conditions, but other gas interferences may exist in the gas circuit. In order to ensure accurate measurement of Masimo module, the following compensations can be manually set according to actual conditions:

— $[O_2 \ Concentration]$: Set the oxygen concentration. Can choose $0\% \sim 100\%$, the default value is 16%.

— $[N_2O \text{ Concentration }]$: Set the concentration of N_2O . Can choose $0\% \sim 100\%$, the default value is 0%.



WARNING:

■ Please set the CO₂ corrections according to actual situation, otherwise, the measured value may be inaccurate and away from actual value.

15.6.6. Operating mode

You can select the operating mode of the CO₂ module according to the actual situation of the module. The operation steps are as follows:

- 1. Select the CO₂ parameter area or waveform area to enter the **【CO₂】** menu.
- 2. Select [Setup] submenu.
- 3. Set [Operation Mode].
 - Measure: Select measurement mode is required when measuring with the CO₂ module.
 - ➤ Sleep: When not using CO₂ for monitoring, it is recommended to set the CO₂ module to sleep mode to increase the life of the CO₂ module.

15.7. Entering Intubation Status

When performing intubation during general anesthesia, you can enter the intubation mode in order to reduce unnecessary alarms. To enter the intubation mode, follow this procedure:

- 1. Select the CO₂ parameter area or waveform area to enter the 【CO₂】 menu.
- 2. Select [Intubation Status] button.

For the details of the intubation mode, see 7.12 Intubation Status.

15.8. Zeroing

■ Mainstream and Sidestream / Microflow CO₂ Module

While zeroing is recommended the first time a CO₂ sensor is connected to the monitor, it is only absolutely necessary when the message "Zero Required" is displayed.

Follow these steps:

- Ensure that the catheter 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).
- 2. Using any of the following methods to perform zeroing:
 - Select the CO₂ parameter area or waveform area, and then select 【Zero】 button.
 - Select 【Zero]quick key→Select【CO₂ Zero]submenu→select the CO₂ to zero.

The screen prompts [CO₂ Zero In Progress...], and the message disappears after the zeroing is completed.

Nomoline ISA CO₂ module will automatically zero when needed.



Δ CAUTION:

- Before zeroing, the side mainstream/ microflow CO₂ sensor must be connected with the sampling tube.
- Before zeroing, the mainstream CO₂ sensor must be connected with the airway adapter.
- Zeroing should not be performed for 20 seconds after the airway adapter or cannula is separated from the patient's airway. Wait a moment before zero correction to dissipate the remaining CO₂ in the adapter or cannula.
- Zeroing should not be performed when the airway adapter or cannula is connected to the patient's trachea.
- When the temperature is not stable, please do not adjust to zero.
- When CO₂ remains in the airway adapter or casing, zeroing will result in

inaccurate measurement or other error conditions. The time required for zeroing will also increase.

- When zeroing, do not rely on gas readings.
- Nomoline ISA CO₂ module do not require user to manually zeroing, and the user cannot successfully send the zeroing command to the module. Nomoline ISA CO₂ module will automatically zero when necessary and the sampling tube is not be inserted.

15.9. Calibration

The monitor has already been calibrated before leaving factory. User can directly apply it measuring in normal conditions (except for the following three cases). Calibrate the gain of the sidestream CO₂ module when the following three conditions occur.

- After the CO₂ module is used for half a year and one year;
- Clinicians doubt the accuracy of readings;
- After the latest calibration, atmospheric pressure or height above sea level varies evidently.



CAUTION:

- Recommend that users carry out calibration operations under the guidance of technical service personnel authorized by the manufacturer. Incorrect calibration procedures may lead to incorrect readings.
- Masimo IRMA CO₂ and Nomoline ISA CO₂ have been permanently calibrated at the factory and do not need to be calibrated by the user.

15.10. Exhaust Emission



WARNING:

■ When using the sidestream/microflow CO₂ measurement on patients who are receiving or have 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 outlet connector of sensor.

15.11. Announcements



WARNING:

- 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.
- Before using, please check whether the airway adapter is damaged. Do not use if damage is found.
- If excessive secretions are found on the airway adapter, replace them immediately.
- When monitoring CO₂ waveform, if changes or abnormal phenomena are found, please check the airway adapter or sampling tube. If necessary, please replace it immediately.
- Note whether the baseline of CO₂ waveform is too high. Sensor or patient problems will cause the baseline to be too high.
- Regularly check CO₂ sensor and pipeline for excessive moisture or secretion accumulation.
- Do not operate the CO₂ module when it is wet or has exterior condensation.
- Do not use microflow CO₂ sensors for patients who cannot tolerate the withdrawal of 50 mL/min±10 mL/min from the airway or patients that cannot tolerate the added dead space to the airway.
- Do not connect the exhaust tube of sidestream/microflow CO₂ module to the ventilator circuit.



CAUTION:

- Use only accessories provided by manufacturer.
- Do not sterilize or immerse the CO₂ sensor in liquids.
- Clean the CO₂ sensor and accessories as directed in this manual.
- Do not apply excessive tension to the CO₂ sensor cable.
- When aerosol drugs are present, please keep the airway adapter away from the breathing circuit. The viscous substance of the aerosol drug can pollute the window of the airway adapter, and the airway adapter needs to be cleaned or replaced in advance.
- For further information on the use of Masimo IRMA CO₂ mainstream module and Nomoline ISA CO₂ module, please refer to the user's manual included with the module.



NOTE:

- This product and its accessories are latex free.
- After the life cycles of the CO₂ 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 CO₂ module.
- Do not place the airway adapter between the ET tube and the elbow, as this may allow patient secretions to block the adapter windows.
- Position the airway adapter with its windows in a vertical and not a horizontal position, this helps keep patient secretions from pooling on the windows.

15.12. CO₂ Troubleshooting

This chapter describes the problems that may be encountered during the use of the monitor. You can first refer to the following table to eliminate them. If the problem persists, please contact the maintenance personnel.



NOTE:

■ For the physiological and technical alarm messages, see *D Alarm Message*.

15.12.1. The Sidestream/Microflow CO₂ module Troubleshooting

Problem	Solution
EtCO ₂ measurement value	1. Judge whether the CO ₂ concentration in the use
too low	environment is too high. If the environmental
	concentration is too high, the measured value is too
	low. If it is more serious, zero will fail. Please pay
	attention to the ventilation of the environment at
	this time.
	2. Check the sampling tube and connectors for
	leakage.
	3. Check the patient status.

15.12.2. The Mainstream CO₂ Module Troubleshooting

Problem	Solution
Elevated baseline	1. Check the patient status.
	2. Check the sensor.

Chapter 16 Anesthetic Gas (AG)

16.1. Introduction

The measuring principle is that anesthetic gas can absorb infrared light. Gases that can be measured by AG module are able to absorb infrared light. Besides, each gas has its own absorption characteristic. First the gas is driven into a sample cell. Then the optic infrared filter selects the infrared light with special wavelength to penetrate this gas. For a given volume, the higher the gas concentration is, the more infrared light is absorbed. We may measure the quantity of the infrared light that have penetrated the gas and then calculate the gas concentration via specialized formula.

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

- 1. Mainstream: Uses an AG sensor attached to an airway adapter directly inserted into the patient's breathing system.
- 2. 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.

AG measurement is applicable for adults, pediatric and neonate.



NOTE: This chapter describes the operation of IRMA gas sensor and ISA

gas module, if you use the IRMA CO₂ or ISA CO₂ sensor, please refer to this chapter.

16.2. Safety information



WARNING:

- To avoid explosion hazard, do not use flammable anesthetic agent such as ether and cyclopropane.
- Using high-frequency electrosurgical units may increase the risk of burning. In this case, do not use antistatic or conductive respiratory tubing.
- Carefully route the sampling line to reduce the risk of patient entanglement or strangulation.



CAUTION:

- Perform the measurement in a well-ventilated environment.
- EtCO₂ values measured from the AG module may differ from that of from the blood gas analysis.
- The AG module has an automatic alarm suppression function. The AG module can perform physiological alarms only after detecting the respiratory wave. When using the AG module to monitor, please ensure that the connection between device and patient is correct.

16.3. AG Measurement Limitation

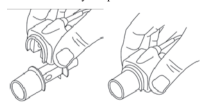
Factors which can affect the accuracy of AG measurement, as following:

- Leaks or internal venting of sampled gas;
- Mechanical shock:
- ◆ Cyclic pressure up to 10kPa (100cmH₂O);
- ◆ Other sources of interference, if any

16.4. AG Monitoring Procedure

16.4.1.1. Mainstream AG module

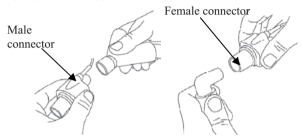
- 1. Plug the AG sensor connector into the AG connector on the monitor.
- 2. Attach AG sensor on the AG airway adapter. Shown as follows:



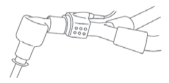
3. A green LED indicates that the AG sensor is ready for use. A blue LED indicates that may measurement of anesthetic gases.



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.



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

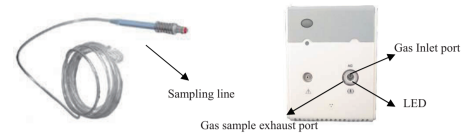


- 6. Pre-use Check
- (1) Before connecting the airway adapter to the breathing circuit, check if the readings on the monitor are correct.
- (2) Before connecting the airway adapter to the patient circuit, verify the gas reading and waveform through the display on the monitor.
- (3) After installing the mainstream CO₂ sensor on the airway adapter, please check the tightness of the patient circuit.
- ♦ Status indicated by LED on AG gas sensor:

LED	The indicated status
Steady green light	System OK
Blinking green light	Zeroing in progress
Steady blue light	Anesthetic in the system
Steady red light	Sensor error
Blinking red light	Check adapter

16.4.2. Sidestream AG module

1. Connect a Nomoline sampling line to the inlet port of the AG module.



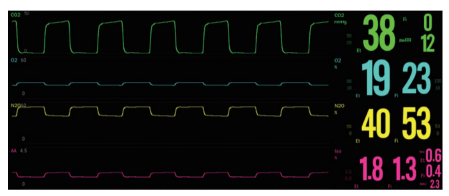
- 2. Connect the gas sample exhaust port to a scavenging system or return the gas to the patient circuit.
- 3. Power up the patient monitor.
- 4. A green LED indicates that the AG module is ready for use.
- 5. Perform a pre-use check. (Before connecting the Nomoline sampling line to the breathing circuit, shall 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 O₂ option fitted: Check that the O₂ 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.
- (7) If applicable: Perform a tightness check of the patient circuit with the sampling line attached.

The state of the LED on the AG module:

LED	The indicated status
Steady green light	System OK
Blinking green light	Zeroing in progress

Steady blue light	Anesthetic in the system
Steady red light	Sensor error
Blinking red light	Check adapter

16.5. AG Display



AG module can send waves and numerics for all measured gases for display on the monitor screen. Including:

- 1. Waveform of CO₂, O₂, N₂O and AA;
- 2. Et and Fi Value of CO₂, O₂, N₂O and AA;
- 3. awRR: airway respiratory rate
- MAC: MAC is defined as the minimum alveolar concentration at steady-state that prevents reaction to a standard surgical stimulus (skin incision) in 50% of patients at 1 atmosphere (i.e. sea level);
- 5. Gas Unit;
- 6. Alarm limit of gas;

AA means a kind of anesthetic agent among the Desflurane (Des), Isoflurane (Iso), Enflurane (Enf), Sevoflurane (Sev) and Halothane (Hal).

When there is only one anesthetic gas, the AA waveform is that of the anesthetic gas. When there are multiple anesthetic gases, the AA waveform shows the main anesthetic gas waveform.

16.6. 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 = 0.77%, Enf = 1.7%, Iso = 1.15%, Sev = 2.1%, Des = 6.0%



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 and ISA AX+/OR+ sensors.

16.7. Setting AG

16.7.1. Setting the AG alarm

You can set the AG alarm through following steps:

- 1. Select the AG parameter area or waveform area to enter the **[Gas]** menu.
- 2. Select the desired gas submenu.
- 3. Select [Alarm] submenu.

If the alarm setting is protected by password, enter the password. For detail, please refer to 7.7.2 Changing Alarm Setup Protection Mode.

4. Set alarms as needed.

16.7.2. Setting Apnea Alarm Duration

You can set apnea alarm duration through following steps. When patient's apnea time exceeds the set time, the monitor will trigger an alarm.

- 1. Select the AG parameter area or waveform area to enter the **[Gas]** menu.
- 2. Select **[CO₂]** submenu.
- 3. Select [Alarm] submenu.
- 4. Set [Apnea Delay].



WARNING:

■ It is cannot judge the cause of respiratory apnea through CO₂ monitoring. If the monitor cannot detect the patient's breath again from the moment of the last breath to the preset time, the monitor only provides the alarm of respiratory apnea. Therefore, the alarm of respiratory apnea should not be used for the diagnosis of the patient.

16.7.3. Setting the Gas Unit

For N_2O and Anesthetic Gas AA, the unit of measurement gas is fixed as %. The unit of CO_2 and O_2 can be set by the following steps below:

- 1. Select the AG parameter area or waveform area to enter the **[Gas]** menu.
- 2. Select $[CO_2]$ or $[O_2]$ submenu.
- 3. Select **Setup** submenu.
- 4. Set $\{Unit\}$ of CO_2 or O_2 .

16.7.4. Setting the AG Waveform

You can set the AG Waveform by the following steps below:

- 1. Select the AG parameter area or waveform area to enter the **Gas** menu.
- 2. Select the desired gas submenu.
- 3. Select **Setup** submenu.
- 4. Set **[Wave Speed]** and **[Scale]**. For CO₂, you can also set waveform mode.

16.7.5. Setting the Operation Mode

You can select operation mode for AG module according to the actual situation, the steps are as follows:

- 1. Select the AG parameter area or waveform area to enter the **[Gas]** menu.
- 2. Select **Setup** submenu.
- 3. Set **[Operation Mode]**.
 - Measure: Select measurement mode is required when measuring with the AG module.
 - Sleep: When not using AG for monitoring, it is recommended to set the AG module to sleep mode to increase the life of the AG module.

16.7.6. Setting the Display Switch of MAC value

You can choose whether to display the MAC value in the parameter area as follows:

- 1. Select the AG parameter area or waveform area to enter the **Gas** menu.
- 2. Select AA submenu.
- 3. Select **[Setup]** submenu.
- 4. Turn on or turn off MAC switch.

16.7.7. Entering Intubation Status

When performing intubation during general anesthesia, you can enter the intubation mode in order to reduce unnecessary alarms. To enter the intubation mode, follow this procedure:

- 1. Select the AG parameter area or waveform area to enter the **【AG】** menu.
- 2. Select [Intubation Status] button.

For the details of the intubation mode, see 7.12 Intubation Status.

16.8. Announcements

16.8.1. Mainstream AG module



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.
- 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.
- 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.
- 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.
- 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.
- If the AG airway adapter is detached from the sensor, or there is something wrong with the sensor, the prompting message may pop up on one of above conditions.

16.8.2. Sidestream AG module



WARNING:

- 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.
- Do not use external ambient cooling of the ISA device.
- Do not apply negative pressure to the Nomoline (i.e. by a syringe) to remove condensed water.
- 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.
- Always use a bacteria filter on the evac side if sampled gas is intended to be re-breathed.
- Do not place the sidestream AG module in any position that might cause it to fall on the patient.



CAUTION:

■ Do not operate the sidestream AG module outside the specified operating temperature environment.

16.9. AG Maintenance

16.9.1. Zeroing

♦ Mainstream AG module:

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 have error;
- ---- A"Zero Required" alarm message is displayed;
- ---- Airway adapter is replaced.

As following procedures:

- 1. 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. Wait for the sensor to warm up:
 - ---- For IRMA CO₂ sensor: Allow 10s for warm up of the sensor after power on and after changing the IRMA airway adapter;
 - ---- For IRMA AX+ sensor: Allow 30s for warm up of the sensor after power on

and after changing the IRMA airway adapter.

- 3. Use any of the following methods for zero:
 - Select the AG parameter area or waveform area to enter 【Gas】 menu→ select 【CO₂】 submenu→select 【Zero】.
 - > Select 【Zero】 quick key→select 【CO₂ Zero】 submenu→select 【Zero】
- 4. In the process of zero calibration, the screen prompts 【CO₂ Zero In Progress...】, and the indicator light on the sensor flashes in green. After the indicator light returns to steady green light, the carbon dioxide display value on the screen is "0". After finishing zeroing, the prompt information disappears, and the measurement can be started.



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.

♦ Sidestream AG module:

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 3s for CO₂ module and less than 10s 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.

16.9.2. Preventive maintenance

♦ Maintenance for sidestream AG module

The sidestream AG 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 contact manufacturer or the professional person authorized by manufacturer.

16.10. AG Troubleshooting

When the AG airway is occluded, the message **【AG Airway Occluded】** appears. In this case, check for the follows until the message disappears:

- 1. Check the airway adapter for occlusion and replace if necessary.
- 2. Check the sample line for occlusion or kinking and replace if necessary.
- 3. Check the gas outlet and the exhaust tube for any occlusion. If the message of occlusion does not disappear, it is probably the module fault.



NOTE:

■ For the physiological and technical alarm messages, see *D Alarm Message*.

Chapter 17 Drip Monitor (DM)

17.1. Introduction

DM (Drip Monitor) module uses photoelectric non-contact principle to detect the dropping of medical drops in the infusion tube set, trigger the circuit to work, count the dropping frequency of medicine drops, and thus obtains the drip rate of infusion drops. After the completion of infusion is detected, the infusion pipeline is clamped, the infusion is blocked and a signal is sent to the monitor, and the monitor generates an infusion completion alarm message according to the signal to prompt medical personnel to change the liquid medicine or perform needle pulling operation.

DM is applicable for adult, pediatric and neonatal patients.



CAUTION:

■ DM only measures the number of drops in the set infusion tube assembly and does not participate in drop control.

17.2. DM Safety information



WARNING:

- During measurement, the liquid level in the drip chamber should be kept below the liquid level indicator line of the infusion monitoring module.
- Please make sure that the outside of the drip chamber is not sticky with water, otherwise the dripping rate measurement may be inaccurate.
- The operator should pay attention to the length of the infusion tube and use the extension tube when necessary to avoid accidents caused by pulling infusion tube due to the patients turning over.
- The Drip monitor (DM) measurement function isn't intended to measure the drips rate in the infusion process of analgesic, chemotherapy medicine and insulinum.



CAUTION:

- In order to ensure the accuracy of dripping rate measurement, the drip monitor module should be vertically installed or naturally hung on the infusion stand using matching bracket.
- This function is an assistive technology implementation method designed for the high-quality infusion nursing services, and cannot replace manual monitoring and speed control operations during infusion.
- This function is suitable for working under relatively static conditions. Therefore, avoid using it in a moving state, and avoid shaking and tilting at a large angle. When the water mist and small water drops in the dropper are seriously hung on the wall, it may interfere with the detection. If necessary, you can flick the wall of the dropper with your finger to shake off the small water drops.
- The DM module uses infrared sensing to detect, so it should be avoided in strong light environment.

17.3. DM measurement

17.3.1. Start infusion

17.3.1.1. Connect DM module cable

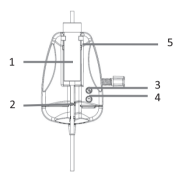
Insert the DM module cable into the DM socket on the monitor, and the drip monitor interface will be displayed on the monitor.

17.3.1.2. Pre-perfusion infusion tube

Close the flow clamp tightly and connect the infusion tube set with the infusion container, then squeeze the drip chamber and pour the liquid medicine to the 1/2 position of the drip chamber. Open the flow clamp, fill the liquid medicine to the tip needle, and then close the flow clamp tightly.

17.3.1.3. Install infusion tube set into DM module

Push the drip chamber into the slot of the drip chamber of the DM module, and clamp the pipeline which is connected to the lower part of the drip chamber into the clamping groove of DM module, as shown in the figure. The DM module is fixed to a suitable position by means of supporting brackets or hanging ropes. Then, exhaust the pipeline to ensure that the gas in the pipe set is exhausted and close the flow clamp tightly.



- (1) The slot of the drip chamber
- (2) Liquid stop clamp
- (3) Start/Stop DM button
- (4) Liquid stop clamp reset button
- (5) Liquid level indicator line of DM module



WARNING:

■ Ventilating operation can only be performed when the infusion is not performed and the infusion tube is not connected to the patient.

17.3.1.4. Configure related parameters

If the unit of mL/h needs to be adopted, can switch "Drops/min" to "mL/h" in the menu of the monitor and set the conversion parameter between the number of drops and mL.



CAUTION:

■ The drip rate corresponding to the 1 mL infusion volume must be entered

according to the relevant statement of the infusion set used. For example, the Double-Dove tube declares that 20 drops of distilled water are equivalent to 1 mL \pm 0.1 mL, so enter: 20 in the Drops/mL parameter setup.

17.3.1.5. Start DM measurement and adjust drip rate

Connect the infusion tube to the patient, start drip monitor (DM) measurement through the "Start/Stop" button on the DM module, and adjust to the desired drip rate via the flow clamp. The DM module indicator light switches from yellow to green and blinks synchronously with the dropping of liquid drops.

17.3.2. Stop infusion

During the infusion or after the infusion is completed, press the "Start/Stop" button of the DM module, and the indicator light of the module will switch to a yellow state. At this time, the monitor will exit the DM function and will no longer perform drip monitor measurement.



WARNING:

■ In the non-infusion drip monitor state, when infusion is completed, the monitor will not stop the infusion and send out an infusion completion alarm. Just quit DM function, but the infusion is still continuing. If the infusion needs to be stopped, the liquid stop clamp of the tube set needs to be operated to stop the infusion.

17.3.3. Infusion completion

When infusion is completed, the indicator light of DM module is switched to red flashing state, the liquid stop clamp is automatically closed, block the pipeline, stop infusion, and the monitor generates infusion completion alarm.

After recognizing the alarm, the medical staff confirms the alarm, separates the infusion tube from the patient, presses the "Reset" button on the DM module, opens the liquid stop clamp, takes out the infusion tube set, and finishes a drip monitor.

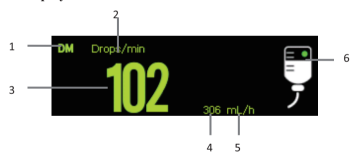
If you need to replace the liquid medicine container, please follow the steps as below:

- When the liquid stop clamp of the DM module is closed, remove the liquid medicine container from the infusion pipeline;
- 2) Connect the infusion pipeline with a new liquid medicine container;
- 3) Open the liquid stop clamp through the "Reset" button on the DM module, and then press the "Start/Stop" button to continue drip monitoring.

17.4. DM module indicator

Status	Indicator
Drip monitoring	The green light is always on and flickers with
	drops of liquid.
Suspension or stop of infusion	Yellow is always bright.
Infusion completed and stopped	Red light flashing (2Hz)

17.5. DM display



- (1) Parameter Label
- (2) Drip rate unit
- (3) The value of drip rate
- (4) The value of flow rate
- (5) Flow rate unit
- (6) Working state diagram

The green dot blinks during drip monitoring, and when drip monitor is stopped, the yellow dot light without flashing; when the infusion is completed, the whole symbol light white and red alternately.

17.6. Setting DM

17.6.1. Setting main unit

You can set the display of the DM main unit by the following steps as below:

- 1. Select the DM parameter area or waveform area to enter **[DM]** menu.
- 2. Set **[Unit]** of DM. The selected unit is displayed in the DM parameter area in the form of a main unit.

17.6.2. Setting unit conversion parameters

In order to ensure the accuracy of the flow rate, you need to set the conversion parameter between the number of drops and mL. The steps are as below:

- 1. Select the DM parameter area or waveform area to enter **[DM]** menu.
- 2. Set **[Drop Per Milliliter]** of DM. The default value is 20.

17.7. DM examination

The DM module has already been calibrated before leaving factory. Generally, the user can directly measure it. Please measure and calibrate the DM module when the following two situations occur:

- ——After the DM module is used for half a year to one year;
- ——Clinicians doubt the accuracy of readings;

DM calibration must be completed by maintenance personnel.

Chapter 18 Bispectral index (BIS)

18.1. Introduction

Bispectral index (BIS) monitoring is used to monitor the level of consciousness state of patient with general anesthesia or sedative treatment in the operating room and intensive care unit. BIS monitoring involves putting a BIS sensor in patient's forehead to receive, filtrate, and process the patient's EEG signals through the BIS module. Through bispectral index and power spectrum analysis, a BIS value which can represent the consciousness level of the patient can then be obtained. Furthermore, by using this method, patient's level of consciousness state can be comprehensively evaluated and depth of anesthesia can be determined based on EEG signals.

BIS measurement is applicable for adult and pediatric.

18.2. Safety information



WARNING:

- BIS monitoring is not intended for neonatal patients.
- The conductive parts of electrodes, sensors, and connectors can't contact other conductive parts, including the ground.
- To reduce patients' risk of ecrasement, the monitor must be placed carefully and the patient interface cable must be immobilized.
- When a defibrillator is used for patients, the BIS sensor can't be placed between the pads of the defibrillator.
- In high-frequency surgery, when the electrodes are connected, the BIS sensor cannot be put between the surgery units and the return electrode of electrical surgical equipment in case of burns.
- When brain stimulation components are used, to reduce the risk of combustion, the stimulation electrode should be placed far away from the BIS sensor. Meanwhile, the sensor should be placed according to the instructions for use.
- The clinical effect, risk, benefit and application of the BIS measurement have not undergone full evaluation in the pediatric population.
- Due to limited clinical experience, for patients with neurological disorders,

- patients taking psychoactive medication, and pediatric under one year old, BIS values should be interpreted cautionsly.
- BIS monitoring can only be used as an auxiliary means of clinical judgment and training. The interpretation of BIS must be combined with other clinical signs. It is not recommended to rely solely on BIS value for anesthesia management during operation.
- Misinterpretation of BIS measurement values may result in incorrect administration of anesthetic agents and/or other potential complications of anesthesia or sedation.
- When using some anesthetic combinations, you should be cautious when interpreting BIS values, such as when relying mainly on ketamine or nitrous oxide/chloroethane for anesthesia.



- Make sure that the BIS sensor does not contact the patient's skin for a long time because the equipment will generate heat, causing discomfort.
- Do not use the BIS sensor if the sensor gel is dry. To avoid dryout, do not open the pack until you are ready to use the sensor.
- When using electro-convulsive therapy (ECT) equipment during BIS monitoring, place ECG electrodes as far as possible from the BIS sensor to minimize the effect of interference. Certain ECT equipment may interfere with the normal function of the BIS monitoring. Check for compatibility of equipment.
- The BIS measurement based on measuring the EEG signal is inherently very sensitive. Do not use electrical radiating equipment close to the BIS equipment.
- Abnormal or excessive electronic interference or myoelectric activity, such as shivering, muscle activity or rigideity, sustained eye movements, head and body motion and other interference may lead to inaccurate BIS value. Also, improper sensor placement and poor skin contact (high impedance) may cause artifact, thus interfering with BIS measurement.
- External radiating devices may affect the BIS measurement.
- Poor signal quality may lead to inaccurate BIS values.

18.3. BIS parameter

Single side BIS monitoring provides the following parameters:

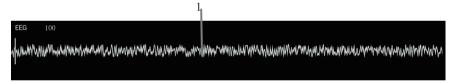
Parameter	Description		
	Reflect the patient's level of consciousness. During the		
	operation, the BIS value of patients under general anesthesia is		
	generally within 40 - 60.		
	100: the patient is wide awake.		
DIG (D' 1	70: the patient is under mild hypnosis and the possibility of being		
BIS (Bispectral	awakened is low.		
Index)	60: the patient is under general anesthesia, loses consciousness,		
	and is under moderate hypnosis.		
	40: the patient is in deep hypnotic state and loses consciousness.		
	0: EEG waveform is a flat line, the patient has no electrical brain		
	activity.		
	Reflect the signal quality and provide information about the		
	reliability of the BIS, SEF, TP and SR values during the last		
	minute, and its range is 0-100%. The greater the SQI value, the		
SQI (Signal Quality	better signal quality. When the signal quality is too low and the		
Index)	BIS value cannot be calculated, the BIS value or other parameter		
maex)	values may not displayed.		
	0%~15%: Cannot deduce value.		
	15%~50%: Cannot deduce value reliably.		
	50%~100%: The value is reliable.		
	Reflect the electrical power of muscle activity and high		
EMG	frequency artifacts. Low EMG indicates that EMG activity is		
(Electromyography)	low. The minimum EMG may be about 25 dB.		
	EMG<55 dB: This is an acceptable EMG;		
	EMG≤30 dB: This is the optimal EMG.		
SR (Suppression	SR numeric is the percentage of time over the last 63s period		
Ratio)	during which the EEG is considered to be in a suppressed state.		

Parameter	Description		
SEF (Spectral Edge	The SEF is a frequency below which 95% of the total power is		
Frequency)	measured.		
	Indicate the status of brain, which indicated the absolute total		
TP (Total Power)	power in the frequency ranging from 0.5-30Hz. The useful range		
	is from 30-100 dB.		
	BC values are used to quantitate inhibition by measuring the		
BC (Burst Count)	number of EEG burst pulse within each minute. A "burst" is		
	defined as a short EEG active phase, preceded and followed by a		
	resting phase respectively.		

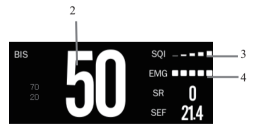
18.4. BIS Display

The BIS parameter area displays the value of a primary parameter and up to 4 secondary parameters. You can select the parameter for display in the **[Select Parameter]** page of the **[BIS]** menu. See *18.6.5 Selecting the Displayed BIS Parameters* for details.

♦ Waveform Display



♦ Parameter Display



(1) BIS waveform area

The display of BIS waveform area depends on the setting of 【Display】 from the 【Setup】 of BIS menu. For more information, see 18.6.5 Setting the Displayed of BIS Waveform Area.

- (2) BIS value
- (3) SQI indicator
 - Empty: SQI<15%, unable to calculate the values of BIS and secondary parameter. BIS and secondary parameter values are displayed as "---".
 - 1-2 bars: SQI 15%-49%, cannot calculate the parameter values reliably.
 - 3-5 bars: SQI 50%-100%, values are reliable.
- (4) EMG indicator
 - Empty: EMG<30dB, this is optimal measurement status.
 - 1-4 bars: EMG 30dB-55dB, this is an acceptable EMG.
 - 5 bars: EMG>55dB, this is an unacceptable EMG.

18.5. BIS Monitoring Procedure

To perform BIS monitoring, follow this procedure:

- 1. Connect the BIS cable to the BIS module.
- 2. Fix the BIS cable in the convenient location near the patient, taking care not to be higher than the patient's head.
- Attach the BIS sensor to the patient's forehead following the instructions supplied with the sensor.
- 4. Connect the patient cable to the BIS cable.
- Connect the BIS sensor to patient cable. Once a valid sensor is detected, the device
 will automatically measure the impedance of all electrodes and display the
 measured results in the BIS window.



WARNING:

■ To minimize the risk of patient strangulation, the patient cable must be carefully placed and secured.



- Ensure that the BIS cable does not come into prolonged contact with your patient's skin, as it may generate heat and cause discomfort.
- Please make sure that the patient's skin is dry because a wet sensor or salt bridge

may cause wrong BIS and resistance values.

- After the BIS cable and BIS module are connected, make sure that the connection of BIS cable will not happen again. When the BIS cable need be disconnected from the BIS module, carefully hold the connector and then pull it out, rather than pull and drag the cable.
- Do not use the BIS sensor if the sensor gel is dry. To avoid dryout, do not open the pack until you are ready to use the sensor.
- Due to intimate skin contact, reuse may pose risk of infection. If skin rash or other unusual symptom develops, stop using and remove.

18.6. Setting BIS

18.6.1. Setting the BIS Alarm

You can set the BIS alarm through the following steps:

- 1. Select the BIS parameter area or waveform area to enter the **[BIS]** menu.
- 2. Select [Alarm] submenu.
 - If the alarm setting is protected by password, enter the password. For detail, please refer to 7.7.2 Changing Alarm Setup Protection Mode.
- Set alarms as needed.

18.6.2. Setting the BIS average time

The average time is the time interval for averaging the BIS data. The shorter the average time, the more sensitive the monitor will respond to changes in the patient's state; the longer the average time, the smoother the BIS trend, the smaller the variation, and the less likely to be interfered by artifacts.

The steps of setting the average time are as follows:

- 1. Select the BIS parameter area or waveform area to enter the **[BIS]** menu.
- 2. Select **[Setup]** submenu.
- 3. Set [Smoothing Rate].

18.6.3. Setting the Display of BIS Waveform Area

You can set the display of BIS waveform area by the following steps below:

- 1. Select the BIS parameter area or waveform area to enter the **[BIS]** menu.
- 2. Select **[Setup]** submenu.
- 3. Set [Display].
 - ◆ EEG waveform (such as 【EEG】): set 【Scale】 and 【Speed】 of EEG waveform
 - ♦ BIS parameter trend: set 【Trend Time】.

18.6.4. Switching Off the Filter

The filter can filter EEG interference. It is switched on by default. To disable the filter, follow this procedure:

- 1. Select the BIS parameter area or waveform area to enter the **[BIS]** menu.
- 2. Select **[Setup]** submenu.
- 3. Switch off [Filter].

18.6.5. Setting the Displayed BIS Parameters

Besides the BIS value, you can also display up to four secondary parameters in the BIS numeric area. To select the displayed parameters, follow this procedure:

- 1. Select the BIS parameter area or waveform area to enter the **【BIS】** menu.
- 2. Select [Select Parameter] submenu.
- 3. From the 【BIS Parameter Area】, select a secondary parameter block, and then select a secondary parameter from 【Parameters】 area.

18.7. Sensor Check

Impedance Inspection refers to the impedance inspection of signal electrodes and reference electrodes. Within the effective rage, this inspection will not lead to prompts.

 Auto: Automatic impedance inspection of ground electrodes is operated every 10 minutes. This process can cause artifacts to EEG waveforms. In the case that the inspection of ground electrode fails, impedance inspection will be restarted. This operation continues until ground electrodes pass the inspection.

- Manual: Open the 【BIS】 menu. After select 【Sensor Check】, each time of
 press on the 【Start Check】 button results in an inspection of ground
 electrodes.
- Cycle Impedance Check: For measuring the exact impedance on each electrode. Once Sensor is connected, Cycle Impedance Inspection will start automatically. When electrode impedance is too high, such information will be displayed on the screen.

18.7.1. Automatic Sensor Check

Once the sensor is connected, a cycle impedance check starts to check the sensor type, status and impedance of all the electrodes, including the signal electrodes, the reference electrodes and the ground electrode.

After the initial sensor check, the monitor performs automatic impedance check during BIS monitoring. The items are as below:

- Automatically and continuously check the impedance of the signal electrode and reference electrode. This check does not affect the EEG waveform, as long as the impedance is within the valid range, no prompt information and check result will be given.
- Check the impedance of the ground electrode every 10 minutes, and each time it takes about 4s. The ground electrode impedance inspection will cause artifacts in the EEG waveform. If the ground electrode check fails, the monitor will start the ground electrode impedance check again, and this will continue until the ground electrode impedance check passes.

18.7.2. Manual Sensor Check

You can start sensor check manually when you need. The steps are as below:

- 1. Select **[Sensor Check]** submenu on the **[BIS]** menu.
- 2. Select **[Start Check]** to start sensor check.

The sensor check window displays the following items:

• The status of each electrode

- Sensor Type
- Expiration time or usable cycles.

18.7.3. BIS Sensor Status

The color of each electrode indicates its status:

Color	Status	Description	Action
Red	Lead	Electrode falls off	Press down on the edge of the sensor to ensure
	off	and has no skin	that the sensor is in good contact with the skin.
		contact	Check the connection of sensor. If the problem
			still exists, remove the sensors and clean the skin
			thoroughly, and then place the sensor again or
			replace with a new sensor after the skin is dry.
Grey	Noise	The EEG signal is	Check the contact between the sensor and the
		too noisy.	skin. Press down on the edge of the sensor to
		Impedance cannot	ensure that the sensor is in good contact with the
		be measured	skin.
Yello	High	The impedance is	Check the contact between the sensor and the
w	impeda	above the limit	skin. Press down on the edge of the sensor to
	nce		ensure that the sensor is in good contact with the
			skin.
Green	Pass	The impedance is	No action necessary.
		within valid range	

Although BIS may still be measured when the electrode status is **[Noise]** (Grey) or **[High Impedance]** (Yellow), for best performance, all electrodes should be in **[Pass]** status.

The sensor check may fail for the following reasons:

- Sensor impedance too high
- Incorrect sensor application
- Poor sensor connection
- Sensor or patient cable failure

If the sensor check fails, you can take the following measures:

- Check the sensor
- Reapply the sensor according to instructions
- Check sensor connection
- Replace sensor or patient cable

18.8. BIS Troubleshooting

This chapter describes the problems that may be encountered during the use of the monitor. You can first refer to the following table to eliminate them. If the problem persists, please contact the maintenance personnel.



NOTE: For the physiological and technical alarm messages, see D Alarm

Message.

Problem	Solution	
Unable to start measurement	1. Check if the application site of sensor is correct and if	
	the contact with the patient's skin is good.	
	2. Check if the sensor type is correct.	
	3. Check whether the patient cable and module are	
	connected or not.	

Chapter 19 Electroencephalograph (EEG)

19.1. Introduction

There are frequently electrical activities in human brain cortex. Brain cortex neuron has spontaneous electrical activities too. When the brain is working, the bioelectricity produced by the outer cortex cells will change with time and space. Using the electrodes placed on the scalp surface to detect the electric potential difference over time, and then the EEG waves recorded can be called EEG. Through analysis of primitive brain waves and compressed spectrum array (CSA), the EEG module can diagnose brain abnormal circumstances and brain lesions timely.

EEG measurement is applicable for adult, pediatric and neonate patients.

19.2. Safety Information



WARNING:

- Do not touch the patient, or table, or instruments during defibrillation.
- When connecting electrodes and/or patient cables, ensure that the EEG leads and connectors do not come into contact with other conductive parts or earth.
- When using high-frequency surgical equipment, do not place EEG electrodes between the surgical site and the negative plate of the electrosurgical equipment to avoid burns.
- The EEG electrode must not be located between defibrillator pads when a defibrillator is used on a patient under monitoring.
- To ensure proper defibrillator protection, use only recommended cables and leadwires.
- EEG monitoring can only be used as an auxiliary means of clinical judgment and training.



- Interference from a non-grounded instrument near the patient and electrosurgery interference can cause problems with the waveforms and the CSA.
- Radiated field strengths above 1 V/m and patient signals $\leq 50 \mu V$ may cause noise on the EEG waves at various frequencies.
- Only use accessories specified in this manual. Follow the instructions for use and adhere to all warnings and cautions.
- Implants (such as pacemakers), other devices attached to the patient, and other devices nearby (such as high-frequency equipments) can interfere with the display of waveforms, values, and compressed spectrum.
- External radiating may disturb the measurement. It is recommended to avoid the use of electrical radiating equipment in close proximity to the monitor.
- Interference from ECG can be eliminated by adjusting the filter settings.

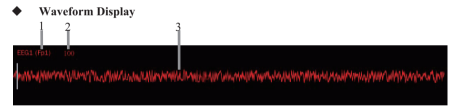
19.3. EEG Parameters

EEG monitoring provides these parameters: SEF, MDF, PPF, TP, Delta, Theta, Alpha, Beta. The parameter area of each channel can display up to 6 parameters.

Parameter	Description	
SEF (Spectral Edge	The SEF is the frequency below which a defined percentage of	
Frequency)	the Total Power lies.	
MDF (Mean Dominant	The MDF is the mean value of the frequency which dominates	
Frequency)	the measured EEG.	
PPF (Peak Power		
Frequency)	The PPF is the frequency with the highest measured amplitude.	
	The TP numeric indicates the power in the measured frequency	
TP (Total Power)	band.	

Parameter	Description
	EEG is traditionally divided into four frequency bands: Delta,
	Theta, Alpha, Beta. Relative Power is the percentage of total
	power falling in corresponding band. For example, Delta% =
	Power in Delta band/Total power.
	Delta: 0.5~4 Hz. Appearing on the condition of Sleep, deep
Delta, Theta, Alpha,	anesthesia, hypoxia or brain organic lesions.
Beta (Relative Power)	Theta: 4~8 Hz. Appearing on the condition of drowsy and
	central nervous system inhibition.
	Alpha: 8~13Hz. It is the most obvious wave of rhythmic
	waves, and can be produced by the whole cortex.
	Beta: 13~30Hz. It is a kind of fast wave. Its emergence means
	the brain is a little exciting.

19.4. EEG Display



Parameter Display



- (1) Lead label
- (2) EEG waveform scale. For more information, see 19.7.1 Setting the waveform

scale.

(3) EEG waveform

You can configure the displayed EEG waveform. A maximum of 4 EEG waveforms can be displayed.

(4) EEG parameters

You can configure the displayed EEG parameters. A maximum of 6 EEG parameters can be displayed. For more information, see 19.7.6 Setting Displayed EEG Parameters.

19.5. EEG Monitoring Procedure

To monitor EEG, follow this procedure:

- 1. Connect EEG module, patient cable and lead wires.
- Select lead combination. You can select a predefined lead combination and you
 can also customize a lead combination. See 19.5.1.2 Customizing a Lead
 Combination for details.
- 3. Mark the electrode site on the patient's head according to the lead combination you have chosen.
- 4. Prepare the skin of the electrode application sites.
- Apply the electrodes according to the site indicated on the screen. See 19.5.2
 Applying Electrode for details.
- 6. Perform sensor check and observe the results. See 19.6 EEG Sensor Check.
- When the electrode connection status of each channel turns green, the measurement can be started.

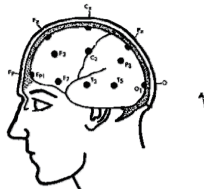
19.5.1. Select the lead combination

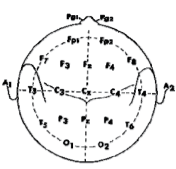
The monitor provides a predefined lead combination, you can modify the predefined lead combination, and set it as a customized lead combination. For more information, see 19.5.1.2 Customizing a Lead Combination.

19.5.1.1. EEG Electrodes Sites

The electrode locations are labeled according to the international 10-20 electrode placement system.

The measurement in longitudinal direction of the head is based on the midline connected from nasion to occipital protuberance. According to the corresponding distance from the midline, ten points are defined: left forehead point FP1, right forehead point FP2, left brow point F3, right brow point F4, left central point C3, right central point C4, left apex point P3, right apex point P4, left occipital point O1 and right occipital point O2. Relative pictures are as follows:





- Odd electrode: placed on the left.
- Even electrode: placed on the right.
 - ◆ F: Frontal
 - ◆ T: Temporal
 - ◆ C: Central
 - P: Parietal
 - O: occipital
 - ◆ Z: Midline electrodes

19.5.1.2. Customizing a Lead Combination

Set a lead combination through the following steps:

- 1. Select the EEG parameter area or waveform area to enter the **【EEG】** menu.
- 2. Select **[Sensor Check]** submenu.
- 3. Set the electrodes sites of each channels in 【Sensor Check】 menu: Select an electrode from the channel area on the left, and then select the electrode site from the electrode map on the right. Repeat this operation until you have defined all the electrodes as desired.

19.5.2. Applying electrode

The steps of applying electrodes are as following below:

- 1. Mark the electrode site on the patient's head according to the lead combination you have chosen.
- Comb or cut the hair away from the spots and rub the skin with the abrasive paste to remove oil and grease.
- 3. Wear EEG cap for the patient, and stick the tape fixed in patient's chin firmly.
- 4. Place the pillar- type electrodes (CH1、CH2、CH3、CH4 and GND) on the patient's head according to the selected "lead setup". Clip the electrodes (A1、A2) on the ears' lobes respectively.
- 5. Pouring moderate coupling agent into the pillar- type electrodes holes in turn. When the contact area of electrodes and scalp is overflowing the coupling agent, the coupling is done.



■ Make sure that you have attached GND electrode.

19.6. EEG Sensor Check

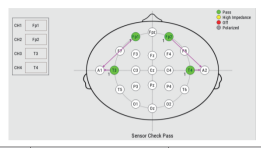
The monitor has an EEG sensor check function. The **【Sensor Check】** submenu displays the status of each electrode and result of sensor check.

19.6.1. Displaying/Hiding Impedance Value

You can display the impedance value on the electrode map of **[Sensor Check]** menu by clicking **[Display Imped. Values (k\Omega)]** key, or hide the value by clicking **[Hide Imped. Values (k\Omega)]** key.

19.6.2. EEG Electrodes status

The status of the electrode is marked by color. The following table lists all electrode status.



Color	status	Description	Action
Red	Lead off	Electrode falls off and has	Reconnect the electrode marked in
		no skin contact.	red.
Grey	Polarization	Electrode polarization or	Check the connection of electrode,
		poor contact	and reconnect the electrode.
Yellow	High	The impedance is above	Check the contact between the
	impedance	the limit	electrode and skin. Reconnect the
			electrode if necessary.
Green	Pass	The impedance is within	No action is required.
		valid range	

For each channel, to get reliable results, all electrodes for this channel should be in **[Pass]** status (Green).

19.7. Setting the EEG

19.7.1. Setting the Waveform Scale

You can set the waveform scale through the following steps:

- 1. Select the EEG parameter area or waveform area to enter the EEG menu.
- 2. Select [Setup] submenu.

3. Set [Scale].

19.7.2. Setting the Waveform Speed

You can set the waveform speed through the following steps:

- 1. Select the EEG parameter area or waveform area to enter the EEG menu.
- 2. Select **[Setup]** submenu.
- 3. Set **[Wave Speed]**. The larger the value, the faster the scanning speed.

19.7.3. Setting the Highpass/Lowpass Filter

The low and high filters can screen out undesirable interference which may come from respiration, movement, etc. The current EEG high and low filter settings are shown at the top of DSA and CSA view.

To set the filter settings, follow this procedure:

- 1. Select the EEG parameter area or waveform area to enter the EEG menu.
- 2. Select **[Setup]** submenu.
- 3. Set [Lowpass Filter] and [Highpass Filter].

19.7.4. Setting Notch Filter

The notch filter can screen out 50Hz/60Hz power line noise. The notch filter is enabled by default. Follow the steps below to turn the notch switch on or off:

- 1. Select the EEG parameter area or waveform area to enter the EEG menu.
- 2. Select **[Setup]** submenu.
- Switch on or off [Notch Filter].

19.7.5. Setting the SEF Threshold

To set the SEF threshold, follow these procedures:

- 1. Select the EEG parameter area or waveform area to enter the EEG menu.
- 2. Select **[Setup]** submenu.
- 3. Set **[SEF Threshold]**.

19.7.6. Setting the Displayed EEG Parameters

You can select the displayed EEG parameters, follow this procedure:

- 1. Select the EEG parameter area or waveform area to enter the EEG menu.
- Select [Select Parameter] submenu.
- 3. Select the EEG parameters to be displayed.

You can select up to 6 EEG parameters. And the selected parameters are applied to all EEG channels.

19.8. Entering the EEG Expand Screen

You can enter the EEG expand screen, follow this procedure:

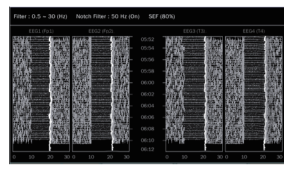
- 1. Select the EEG parameter area or waveform area to enter the EEG menu.
- 2. Select **[EEG Expand]** to enter the EEG expand screen.
- 3. Select the desired submenu to enter corresponding screen:
 - Select 【EEG】 submenu→select 【EEG Channels】, 【Scale】 and 【Speed】 to view corresponding EEG waveform.
 - Select 【Parameter】 submenu, and view the parameter values of each EEG channels.
 - Select 【Trend】 submenu→select 【EEG Channels】, 【Parameters】 and 【Trend Length】 to view corresponding EEG channel and the trend of parameters.
 - Select 【CSA】 submenu to enter CSA screen. See 19.8.1 Entering the EEG CSA screen for details.
 - Select 【DSA】 submenu to enter DSA screen. See 19.8.1 Entering the EEG DSA screen for details.

19.8.1. Entering the EEG CSA screen

The continuous EEG signal is sampled periodically and this value is stored in a frame. Each frame is processed using Fast Fourier Transformation (FFT) to provide a frequency spectrum displayed as a compressed spectral array (CSA). CSA screen shows the EEG value change situation over time. The steps of entering the CSA screen are as

following below:

- 1. Select the EEG parameter area or waveform area to enter the EEG menu.
- 2. Select **[EEG Expand]** to enter the EEG expand screen.
- 3. Select **[CSA]** submenu.



The CSA screen provides CSA of up to 4 channels, which including the following information:

Displayed items	Description
Status bar	The first line displays the current filter range, notch frequency settings, SEF percentage, and trend line labels and color codes. The second line displays the EEC channel labels and lead labels.
Frequency Scale	It is the horizontal axis. The scale range is 0~30Hz.
Spectral lines	The energy at each frequency is computed and displayed as a spectral line.
Trend lines	EEG values are sampled at configured time intervals and displayed as color-coded trendlines. Trendlines are available for up to three frequency numerics (SEF, MF, and PPF). SEF trendline is white, MF trendline is purple, and PPF trendline is green.

From the CSA screen you can select the following items:

- **■ 【EEG Channels**】
- **■** 【Parameters】
- 【Trend Length】
- Power Scale

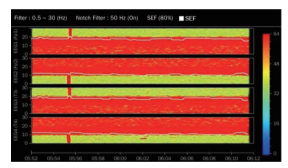
■ 【CSA Clipping】

- Changing [Power Scale] can adjust the amplitude of spectral lines. The wider the power range, the greater amplitude of the spectral lines is.
- Switch on 【CSA Clipping】, the latest spectral line is displayed normally, and other spectral line that cross the latest spectral line area are cleared. Switch off 【CSA Clipping】, all spectral lines will be displayed normally.

19.8.2. Entering the EEG DSA screen

The Density Spectral Array (DSA) is to show changes in the power spectrum distribution over time. To display the DSA screen, follow this procedure:

- 1. Select the EEG parameter area or waveform area to enter the EEG menu.
- 2. Select **[EEG Expand]** to enter the EEG expand screen.
- 3. Select **[DSA]** submenu.



The DSA screen provides DSA of up to 4 channels, which including the following informations:

Displayed items	Description
Status bar	The current filter range, notch frequency settings, SEF percentage, and trendline labels and color codes.
Color bar	It is located at the right of the DSA screen. The color bar color codes the power. Red indicates maximum power, and blue indicates minimum power. You can change the setting of 【Power Range】 to adjust the color of corresponding power.

Frequency Scale	It is the horizontal axis. The scale range is 0~30Hz.
Trend lines	Use colored lines to display the parameter trend within the set trend time. Trendlines are available for up to three frequency numerics (SEF, MDF, and PPF). SEF trendline is white, MDF trendline is purple, and PPF trendline is green.

From the DSA screen you can select the following items:

- **■ 【EEG Channels 】**
- 【Parameters】
- **■** 【Trend Length】
- 【Power Scale】

19.9. EEG Troubleshooting

This chapter describes the problems that may be encountered during the use of the monitor. You can first refer to the following table to eliminate them. If the problem persists, please contact the maintenance personnel.



NOTE: For the physiological and technical alarm messages, see D Alarm

Message.

Problem	Solution	
EEG signal is noisy	1.	Check that the electrodes are properly connected and not
		dried out.
	2.	Check that the electrodes properly contact with skin.
	3.	Check electrodes status, and make sure that the electrodes
		impedance is normal.
	4.	Calm the patient since frontal muscle activity can cause
		artifact.
	5.	Remove sources of external electrical noise (for example,
		the lamps) from the vicinity of the patient's head.
	6.	ECG monitoring may cause artifact; change electrode

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	sites.		
EEG cable and	1. The number of channels in the lead combination is smaller		
electrodes properly	than the number of channels connected to patient. Check the		
connected, but no	number of channels.		
EEG waveforms.	2. Check screen setup and make sure that you have selected		
	the EEG parameter.		
The EEG numerics	The patient has high muscle activity in the head area, or noise		
area displays ""	from some interfering equipment is coupling to electrode cables.		
	Relax the patient and remove the source of noise.		
EEG waveform	Check patient, sweating may cause variations in the		
baseline fluctuates	electrode impedance.		
	2. If the fluctuation if disturbing, prepare the skin and replace		
	the electrodes if necessary.		
The electrode	The ground electrode is poorly connected to the patient. Check		
impedances show	the electrode and cable. If the impedance of the electrode is too		
"" and there is a	high, the measurement fails even if the electrode is properly		
message prompting	attached. Use better electrodes or prepare the skin better.		
to check the ground			
electrode			

Chapter 20 Respiratory Mechanics (RM)

20.1. Introduction

RM (respiratory mechanics) monitoring adopts a flow sensor connected between Y piece of ventilator and patient circuit to measure and display respiratory volume, respiratory flow and airway pressure in airway. From these three parameters, other parameters such as RR, I:E, Compl, etc. are derived. The ventilator settings and respiratory status can be displayed in real time by the RM monitoring, which give a reference for understanding patient respiratory status and correctly using ventilator.

RM monitoring is applicable for adult, pediatric and neonate.

20.2. Safety information



WARNING:

- RM monitoring is for mechanically ventilated patients only.
- The RM module cannot be used with high frequency ventilators.

20.3. RM Parameters

RM monitoring displays the following waveforms and loops:

- Flow waveform
- Paw waveform
- Vol waveform
- FV (flow-volume) loop
- PV (paw-volume) loop
- PF (paw-flow) loop

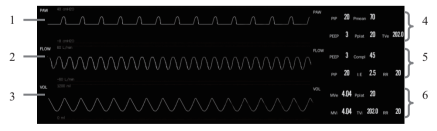
RM monitoring provides 13 measurement values of parameters. According to the nature of the parameters, there are four categories:

Parameter Label	Description	Unit
Paw parameters		
PIP	peak inspiratory pressure	cmH ₂ O

Pplat	plateau pressure	cmH ₂ O		
PEEP	positive end expiratory	cmH ₂ O		
	pressure			
Pmean	mean pressure	cmH ₂ O		
Vol parameters				
TVi	inspiratory tidal volume	ml		
TVe	expiratory tidal volume	ml		
MVi	inspiratory minute volume	L/min		
MVe	expiratory minute volume	L/min		
Other parameters				
RR	respiration rate	rpm		
I : E	inspiratory-expiratory ratio	/		
Compl	compliance	ml/cmH ₂ O		
TV1	First second expiratory rate	%		
Resi	Airway resistance	cmH ₂ O/L/s		

20.4. RM Display

Paw waveform, flow waveform, Vol waveform and relevant parameters can be display in the waveform area. You can select the parameter and waveform for display in the **[Select Parameter]** page of the **[RM]** menu. For more information, see 20.8.6 Setting Parameter for Display.



(1) Paw waveform

(4) Paw parameter area

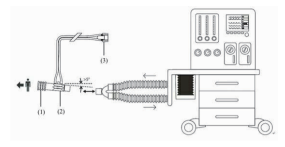
(2) Flow waveform

(5) Flow parameter area

(3) Vol waveform

(6) Vol parameter area

20.5. RM Equipment Connection



- (1) To patient
- (2) RM flow sensor
- (3) Connect to RM plug-in module

20.6. RM Monitoring Procedure

You can prepare for RM monitoring, following these steps:

- 1. Select an appropriate flow sensor based on the patient's type.
- 2. Connect the flow sensor to the RM connector of the module.
- 3. Connect the end of the flow sensor marked **†** to the patient tracheal tube.
- Connect the other end of the flow sensor to the Y-tube of a ventilator or anesthesia machine.
- 5. Check that the connections are tight.



- Before using RM module, must set the barometric pressure properly. Improper settings will result in erroneous reading.
- Check for leaks in the breathing circuit system, as they may significantly affect respiratory mechanics readings.
- Select the appropriate sensor based on the patient type. Improper sensor selection may produce excessive ventilation resistance or introduce excessive airway dead space.
- Support the sensor and airway adapter to prevent the sensor or the airway adapter from being pressed onto the tracheal tube.

■ Carefully route the sensor tubing to avoid the risk of patient entanglement or strangulation.



NOTE:

- In order to avoid the influence of excessive moisture in the measurement air circuit, when installing the flow sensor in the patient air circuit, please place the tube upward, and place the end of the flow sensor connected to the ventilator or anesthesia machine at a certain angle to the horizontal position.
- Do not place the airway adapter between the endotracheal tube and an elbow connector as this may cause patient secretions to block the adapter windows.
- Measurement values provided by a ventilator or an anesthesia machine may differ significantly from the values provided by the RM module, due to different locations of the flow sensor.
- For best measurement result, a heat moisture exchanger should be put between the tracheal tube and the flow sensor. Otherwise, it is necessary to periodically check the flow sensor and catheter for excessive water or secretion condensation, and remove it if necessary.
- During RM monitoring, should avoid placing the patient's breathing circuit close to condensing equipment, such as fans, air conditioners, etc.

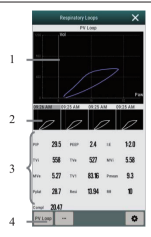
20.7. Viewing the Respiratory Loops

Respiratory loops reflect patient lungs function and ventilation condition, such as the lung's compliance, over-inflation, breathing system leakage and airway blockage.

The monitor provides 3 types of respiratory loops: PV (pressure-volume) loop, FV (flow-volume) loop, and PF (flow-pressure) loop. The three types of loops come from pressure, flow and volume waveforms data.

To view the respiratory loops, choose any of the following ways:

- Select the 【Loops】 quick key.
- Select parameter area or waveform area of Paw, Flow or Vol to enter [RM]
 menu, select [Respiratory Loops].



- (1) Respiratory loop
- (2) Reference loop
- (3) RM parameters
- (4) Button area

20.7.1. Changing the Loop Type

At the same time, the monitor can only display one type of loop. You can select the respiratory loop to be displayed through the following steps:

- Select parameter area or waveform area of Paw, Flow or Vol to enter menu.
- 2. Select [Loops].
- 3. Select the type of loops in the **【Loops】** window.

20.7.2. Saving the reference loop

You can save the real time loops as reference loops. To save the loops, follow this procedure:

- Select parameter area or waveform area of Paw, Flow or Vol to enter menu.
- 2. Select [Loops].
- 3. Select [...] to enter more settings menu.

4. Select [Save Reference Loops].

The reference loops and the time at which the reference loops are saved displayed in the **【Loops】** window simultaneously. Up to 4 groups of loops can be saved as reference loops. If the fifth group of loops needs to be saved as reference, the monitor will prompt that an older group of reference loops should be replaced by the fifth group.

20.7.3. Displaying the Reference Loops

The reference loop and real time loop can overlap and be displayed in the same area of the **【Loops】** window. In this case, the reference loop is drawn in white. To display the reference loop, follow this procedure:

- Select parameter area or waveform area of Paw, Flow or Vol to enter RM menu.
- 2. Select [Loops].
- 3. Select the reference loop to be displayed.
- 4. Select [...] to enter more settings menu.
- 5. Select [Display Reference Loops].

To hide the reference loop, select **【Hide Reference Loops】** button in the more settings menu.

20.7.4. Adjusting the Scale

The scales of the loops and the corresponding waveforms are same. You can adjust the scales of loops, see *20.8.3 Setting the RM waveform scale* for details.

20.8. Setting RM

20.8.1. Setting the RM Alarm

You can set the RM alarm through following steps:

- Select parameter area or waveform area of Paw, Flow or Vol to enter RM menu.
- 2. Select [Alarm] submenu.

If the alarm setting is protected by password, enter the password. For detail,

please refer to 7.7.2 Changing Alarm Setup Protection Mode.

Set alarms as needed.

20.8.2. Setting the RM waveform speed

You can set the wave speed of Paw, Flow and Vol waveform through the following steps:

- Select parameter area or waveform area of Paw, Flow or Vol to enter RM menu.
- 2. Select [Setup] submenu.
- 3. Set [Wave Speed].

20.8.3. Setting the RM waveform scale

You can set the wave speed of Paw, Flow and Vol waveform through the following steps:

- Select parameter area or waveform area of Paw, Flow or Vol to enter RM menu.
- 2. Select **Setup** submenu.
- 3. Set [Paw Scale], [Flow Scale] or [Vol Scale].

20.8.4. Setting the Ambient Temperature

To set the ambient temperature, follow these steps:

- Select parameter area or waveform area of Paw, Flow or Vol to enter RM menu.
- 2. Select **[Setup]** submenu.
- 3. Set **【Atmosphere Temp】**.

20.8.5. Setting the Relative Humidity

To set the relative humidity, follow these steps:

Select parameter area or waveform area of Paw, Flow or Vol to enter RM menu.

- 2. Select **Setup** submenu.
- 3. Set [Relative Humidity].

20.8.6. Setting Parameter for Display

Each parameter areas can display up to 6 parameters. To set the parameters for display, follow these procedure:

- Select parameter area or waveform area of Paw, Flow or Vol to enter RM menu.
- 2. Select **Setup Parameters** submenu.
- 3. Select the parameters for display of **[Paw]**, **[Flow]** and **[Vol]**.

20.8.7. Entering Intubation Status

When performing intubation during general anesthesia, you can enter the intubation mode in order to reduce unnecessary alarms. To enter the intubation mode, follow this procedure:

- Select parameter area or waveform area of Paw, Flow or Vol to enter RM menu.
- 2. Select [Intubation Status] button.

For the details of the intubation status, see 7.12 Intubation Status.

20.8.8. Setting the Barometric Pressure

To set the barometric pressure, follow these procedures:

- Select 【Main Menu】 quick key→from the 【System】 column to select
 【Maintenance】 button→input password→enter.
- 2. Select [Other] submenu.
- 3. Select **[Barometric Pressure]**, then input the value of barometric pressure to which the patient monitor is exposed to.



WARNING:

■ Be sure to set the barometric pressure properly before using the RM module. Improper settings will result in erroneous RM reading.

20.9. RM Troubleshooting

This section describes problems you may encounter during using. You can refer to the following table for troubleshooting. If the problem persists, please contact maintenance staff.



NOTE: See D Alarm Message for physiological alarms and technical alarm

information.

Problem	Solution	
Do not see parameter area	1. Check whether the display of Paw, Flow or Vol is	
and waveform area of Paw,	set in the 【Screen Layout】 menu or not. For	
Flow or Vol.	details, see 3.7.1 Setting Parameters.	
	2. Check whether the RM parameter switch is turned	
	on or not. For details, see 3.7.1 Setting	
	Parameters.	
	3. Check the flow sensors are connected or not.	
Erroneous values	1. Check that the tube connectors and their	
Values seems unstable	connections are tight and not leaking.	
	2. Check that the sensor type is correct.	
	3. Disconnect the flow sensor, and remove the water	
	from the flow sensor.	
Strong vibrations in the	1. Check the patient status.	
loop	2. Check the breathing system for water or	
	secretions.	

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Imcomplete or interrupted	Check the breathing system for leakage.
loops	

Chapter 21 Regional oxygen saturation (rSO₂)

21.1. Introduction

The rSO_2 monitoring provides noninvasive and continuous information of changes in regional oxygen saturation of blood. The measurement takes place in real time, providing an immediate indication of a change in the critical balance of regional oxygen delivery and oxygen consumption. The monitor supports 2 channels of rSO_2 measurement.

 rSO_2 measurement uses near-infrared spectroscopy: the harmless near-infrared light generated by the light-emitting diode (LED) on the sensor passes through the scalp and bone tissue under the sensor. After entering the body, they are either absorbed or scattered back to the shallow and deep detectors of the sensor. Red hemoglobin molecules in red blood cells absorb the most infrared light, and the solid red shadow of each hemoglobin molecule indicates the amount of oxygen it carries. The type and quantity of infrared light absorption data returned to the detector reflect the relative quantity of deoxygenated hemoglobin and total hemoglobin, and the value of a specific area under the sensor can be obtained through calculation.

 rSO_2 measurement is used clinically and can be used for auxiliary monitoring of blood oxygen saturation in the head and forehead of adult, pediatric and neonate, and other parts of body.

The monitor support Masimo rSO₂ plug-in module and Nonin rSO₂ plug-in module.

21.2. Safety Information



- Do not use the rSO₂ measurement value as the only basis for diagnosis and treatment. The rSO₂ rending reflects the oxygenation status of the local tissue under the sensor, and may not reflect the oxygenation changes occurring in other parts.
- Use only recommended or provided accessories. Use any other sensor will compromise accuracy.

■ Use of an electrosurgical instrument in the vicinity of the monitor may interfere with the signal and result in inaccurate rSO₂ measurement.



NOTE:

■ Environments with excessive ambient light such as bright sunlight or strong operating room lighting may require loosely covering the area of the sensor with an opaque drape.

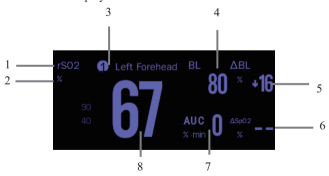
21.3. rSO₂ Measurement Limitation

Factors which can affect the accuracy of AG measurement, as following:

- Cardiogreen, indigo carmine or other intravascular dyes
- Carboxyhemoglobin or other dyshemoglobins
- Hemoglobinopathies
- Conjugated hyperbilirubinemia (direct)
- Myoglobin (Mb) in muscle tissues

21.4. rSO₂ Display

Each parameter area displays one measurement channel.





- (1) parameter label
- (2) rSO₂ unit
- (3) channel number
- (4) Baseline value (BL)
- (5) Change percentage of the rSO₂ value (Δ BL) from baseline: Δ BL = $\frac{rSO_2 BL}{BL}$ * 100%, when the rSO₂ value is greater than or equal to the baseline, the arrowhead is upward; When the rSO₂ value is smaller than the baseline, the arrowhead is downward.
- (6) Difference between SpO₂ and rSO₂: Δ SpO₂ = $|SpO_2 rSO_2|$
- (7) Area Under the Curve (AUC)
- (8) Realtime rSO₂ value
- (9) rSO₂ average of latest 60 minutes

21.5. rSO₂ Monitoring Procedure

- Select the application sites of rSO₂ sensor. See 21.5.1 Selecting the application site of rSO₂ sensor for details.
- 2. Prepare the patient's skin. See *21.5.2 Preparing the skin* for details.
- 3. Apply the rSO₂ sensor. See 21.5.3 Applying the rSO₂ sensor for details.
- 4. Connect the rSO₂ parts. See *21.5.4 Connecting the rSO₂ cable* for details.

21.5.1. Selecting the application site of rSO₂ sensor

The rSO₂ sensor can be placed in a suitable position on the forehead or torso.

Cerebral site selection: select sensor site on the right and/or left side of forehead. Placement of the sensor in other cerebral locations, or over hair, may cause inaccurate readings. Somatic site selection: select sensor site over tissue area of interest (site selection will determine which body region is monitored)



- For cerebral site selection, do not place the sensor over nevi, sinus cavities, the superior sagittal sinus, subdural or epidural hematomas, injured skin or other anomalies such as arteriovenous malformations, as this may cause readings that are not reflective of brain tissue or no readings at all.
- For somatic site selection, avoid placing the sensor over thick fatty deposits, hair or bony protuberances. Do not place the sensor over nevi, hematomas or broken skin, as this may cause readings that are not reflective of somatic tissue or no readings at all.



NOTE:

■ For the specific application site selection of sensors, see the instruction for use of corresponding sensor for detailed information.

21.5.2. Preparing the skin

To achieve the optimal measurement result, the sensor application site should be clean and dry. To properly prepare the skin, follow these steps:

- 1. Shave hair from selected areas
- 2. Gently rub skin surface at sites to remove dead skin cells.
- 3. Thoroughly cleanse the site with a mild soap and water solution.
- 4. Dry the skin completely.

21.5.3. Applying the rSO₂ sensor

To apply the rSO₂ sensor:

1. Remove the protective backing label from the adhesive side of the sensor and

apply the sensor to the skin.

2. Continue applying the sensor by smoothing it to the skin from the center outward. Ensure edges of the sensor are sealed to prevent light from entering.



- rSO₂ sensor is disposable, do not reuse. Reuse the disposable sensor may cause inaccurate reading, unstable reading or no readings at all. Also, reuse may cause an increased risk of cross-contamination among patients.
- Do not autoclave or gas sterilize the sensor.
- Do not place the sensor on a site with severe tissue edema to reduce the possibility of skin damage.
- Please be careful when placing or removing the sensor. Do not put on broken or fragile skin.



NOTE:

- If the patient is monitored for rSO₂ for a long time, the sensor should be replaced every 24 hours or when the adhesive between the sensor and the skin fails.
- The sensor is for external use only and cannot be used in the body under any circumstances.
- To avoid bedsores or pressure sores, do not apply external pressure to the sensor (for example, use hair bands, bandages, tape, etc.).

21.5.4. Connecting the rSO₂ cable

To connect the rSO₂ cable, follow these steps:

- 1. Insert rSO₂ sensor into rSO₂ cable.
- 2. Insert rSO₂ cable into rSO₂ module.



NOTE:

■ Different sensors (adult, pediatric and infant/neonatal) cannot be used simultaneously on the same monitor. Cerebral sensors can be used with somatic sensors on the same monitor.

21.6. Setting rSO₂

21.6.1. Setting the rSO₂ Alarm

To set the alarm, follow these procedures:

- 1. Select the rSO₂ parameter area to enter the **[rSO₂]** menu.
- 2. Select [Alarm] submenu.

If the alarm setting is protected by password, enter the password. For detail, please refer to 7.7.2 Changing Alarm Setup Protection Mode.

Set alarms as needed.

21.6.2. Setting the rSO₂ Auto Low Limit Switch

To set the rSO₂ Auto Low Limit, follow these steps:

- 1. Select the rSO_2 parameter area to enter the $[rSO_2]$ menu.
- 2. Select [Alarm] submenu.
- 3. Set [Auto Low Limit] switch.
 - ◆ When 【Auto Low Limit】 is switched on, 【rSO₂-1%BL】 and 【rSO₂-2%BL】 are activated to allow you to set the percentage of rSO₂ low limits below the baseline. Then the monitor calculated the rSO₂ low limits automatically based on the setting.
 - ◆ When 【Auto Low Limit】 is switched off, the rSO₂ low limits should be set manually.

21.6.3. Setting the rSO₂ Sensor Site

To set the locations of each channel sensors, follow these steps:

- 1. Select the rSO_2 parameter area to enter the $[rSO_2]$ menu.
- 2. Select **[Setup]** submenu.
- 3. Set [rSO₂-1 Sensor Site] and [rSO₂-2 Sensor Site].

21.6.4. Setting the AUC Mode (Only for Nonin rSO₂)

To set the AUC mode, follow these procedures:

- 1. Select the rSO_2 parameter area to enter the $[rSO_2]$ menu.
- 2. Select **[Setup]** submenu.
- Set 【AUC Mode】.
 - ♦ 【Fixed】: the item 【Fixed Threshold】 is activated. In this case, AUC is calculated according to the set fixed threshold.
 - ♦ 【Below Base Percentage】: the item 【Percentage Below Baseline】 is activated. In this case, AUC is calculated according to the set percentage below baseline.

21.6.5. Clear AUC

To clear the current AUC value, follow these steps:

■ Select 【Zero AUC】 button on the 【rSO₂】 menu.

21.6.6. Setting the rSO₂ Baseline

To set the rSO₂ baseline of single channel, follow these steps:

- 1. Select the rSO₂ parameter area to enter the **[rSO₂]** menu.
- 2. Select [Setup] submenu.
- 3. Set $[rSO_2-1]$ Baseline and $[rSO_2-2]$ Baseline . The monitor will set the current rSO_2 value as the baseline.

You can set the baseline of all rSO₂ channels through the following shortcuts:

■ Select **[Set Baselines]** button on the **[rSO₂]** menu. The monitor will set all current rSO₂ value of each channel as the baseline.



NOTE:

■ Set the rSO₂ value measured when patient is sober and eupraxic as baseline. The baseline will be set automatically if it is not set within 5 to 6 minutes and current rSO₂ value is effective.

21.6.7. Setting the Averaging time (Only for Masimo rSO₂)

The shorter the averaging time, the quick the monitor responds to changes in the patient's status. To set the averaging time, you can follow these steps:

- 1. Select the rSO₂ parameter area to enter the **[rSO₂]** menu.
- 2. Select **[Setup]** submenu.
- 3. Set [Averaging Time].

21.6.8. Selecting the rSO₂ Parameters

In the rSO_2 parameter area, rSO_2 is a fixed display parameter, and other parameters can be selected for display. You can also select the displayed parameters according to the following steps:

- 1. Select the rSO₂ parameter area to enter the **[rSO₂]** menu.
- 2. Select [Select parameter] submenu.
- 3. Select parameters to be displayed.

21.7. rSO₂ Troubleshooting

For more information, see D Alarm Message.

Chapter 22 Neuromuscular Transmission (NMT)

22.1. Introduction

NMT measurement uses three-axis acceleration sensor technology, which measures the acceleration of muscle movement by applying controllable electrical stimulation to specific motor nerves, calculates muscle contraction force, and completes the quantitative assessment of muscle relaxation under neuromuscular block.

NMT measurement is applicable for adult and pediatric patients.

22.2. Safety Information



WARNING:

- Use of cables or accessories other than those supplied with the manufacturer may result in serious injury.
- Maintenance on this device may only be performed by the manufacturer or persons explicitly authorized by the manufacturer.
- Do not use the device in close proximity to equipment that produces strong electromagnetic fields, such as high frequency surgical equipment. The cable leads could act as antennae and dangerous currents could be induced as a result.
- Do not apply the device to patients with implanted electrical devices, such as cardiac pacemakers, without first consulting with an appropriate medical specialist.
- The patient should avoid contact with metallic objects that are grounded, produce an electrical conductive connection with other equipment and/or enable capacitive coupling.
- The cables should be positioned in such a way that they do not contact either the patient or other cables.
- Simultaneous connection of a patient to high frequency surgical ME equipment may result in burns and possible damage to the stimulator.
- Operation in close proximity (e.g. 1m) to a shortwave or microwave therapy ME equipment may produce instability in the stimulator output.

- Application of electrodes near the thorax may increase the risk of cardiac fibrillation.
- The NMT stimulation should not be applied across or through the head, directly on the eyes, covering the mouth, on the front of the neck, especially the carotid sinus, or from electrodes placed on the chest and the upper back or cross over the heart.



- Prior to changing the batteries, be sure to switch off the NMT Sub-machine and remove all the cables.
- Remove elements which may adversely affect the connection between the electrodes and the skin, e.g., dirt, hair, oil.
- Ensure that ECG electrodes are not damaged or dried out.
- Large current densities associated with failing ECG electrodes may cause superficial burns.
- Compatible with standard ECG electrodes, but for large currents, it is recommended to use dedicated electrodes.
- Electrodes that have current densities exceeding 2mA/cm2 may require special attention of the operator.
- All the accessories have been certified latex free.
- Inspect all parts for any damage or manipulation. Never use any damaged or manipulated part!
- If an electrically conductive surface of the NMT Sub-machine or its cables are exposed, such electrically conductive surface may shock an operator. Do not use such a device or accessory, please contact the manufacturer for repair.
- The refractory period delay is set at a default value to prevent the user from repeating stimulation while the nerve synapse is recovering from effects of the previous stimulation. A refractory period of less than 12 seconds in TOF mode is not advisable as measurements might not represent the effect of blocking agents on the neuromuscular junction.

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22.3. Stimulation Mode

NMT monitoring provides 7 stimulation modes.

22.3.1. Train-of-Four (TOF) Mode

The TOF stimulation comprises four, square waves with a pulse width of 200

microseconds, 500 milliseconds apart.

The relative contraction strength caused by each stimulus is indicated

graphically in the monitor screen as shown in the picture.

 \triangleright In the case that all four contractions could be measured, the percentage of

measured contraction strength of the fourth stimulus compared to the first

stimulus will be displayed in the monitor screen.

If less than four contractions were measurable, the number of contractions that

could be identified by the accelerometer will be displayed, e.g., 2/4.

22.3.2. Double Burst (DB) Mode

The DB stimulation comprises a burst of three square waves of 200 microseconds

pulse width, 20 milliseconds apart, followed by another burst of three square waves, 750

milliseconds later.

The relative contraction strength caused by each stimulus is indicated

graphically in the monitor screen as shown in the picture.

The percentage of the measured contraction strength of the second contraction

compared to the first contraction will be displayed in the monitor screen.

22.3.3. Post Tetanic Count (PTC) Mode

Tetanus: 50Hz for 5 seconds

Delay: 3 seconds

Twitch: 20 twitches at 1Hz

The PTC stimulation comprises a tetanus stimulation followed by a delay and a

number of twitches. (Default settings are as shown above)

Each counted twitch is indicated graphically in the monitor screen as shown in

the picture.

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The numbers of twitches counted are displayed in the diagnostic screen.

22.3.4. Supra Maximal Current (SMC) Mode

The SMC mode is used to determine the current at which the response from the patient no longer increases by stimulating at increasing intensities (5mA to 80mA). The SMC stimulation comprises of sixteen 1 Hz square wave pulses with a pulse width of 200 microseconds.

> Each counted twitch is indicated graphically in the monitor screen as shown in the picture.

22.3.5. Auto Mode

In auto mode the module automatically changes the stimulation mode depending on the depth of the block. In this mode, display one of six depth-of-block zones according to the captured accelerometer response.

- Recovered: A TOF ratio of larger than 90%.
- Minimal: A TOF ratio between 40% and 90%.
- Shallow: A TOF ratio less than 40%, but all pulses present.
- Moderate: A TOF count of 1 to 3.
- Deep: A PTC of one or more.
- Profound: A PTC of zero.
- > The relative contraction strength caused by each stimulus is indicated graphically in the monitor screen.

22.3.6. Twitch (TWI) Mode

The Twitch stimulation comprises a 200 ms square wave pulse. The optional frequency is 1Hz, 2Hz and 5Hz, and the default repetition frequency is 2Hz.

22.3.7. Tetanus (TET) Mode

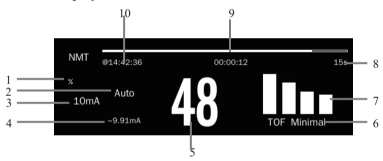
The Tetanus stimulation comprises series of 200 ms square wave pulses repeated at a 50Hz or 100Hz repetition rate or frequency.

22.4. NMT Parameters

The following	table lists th	e NMT	parameters in	n different	stimulation mod	des.

Stimulation Modes	Unit of parameters	Number of bars
TOF	%	4
DB	%	2
PTC	/	20
SMC	/	16
Auto	% or /	20
TWI	/	/
TET	/	/

22.5. NMT Display



- (1) Unit
- (2) Stimulation mode
- (3) Stimulation current
- (4) Average of actual current delivered
- (5) Parameter value: Calculated percentage in TOF and DB mode. The number of contractions in TOF mode is less than 4, and the number of recognizable contractions is displayed / 4, and the number of twitch stimuli counted in PTC mode is displayed.
- (6) The depth of the block in the Auto Mode: Display six depth-of-block zones, see **22.3.5** Auto Mode for details.
- (7) Bar graph: indicates the amplitude of response to the stimulation.
- (8) Measurement interval time: In three modes of TOF, DB and PTC, display the set

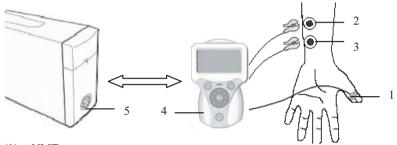
value of the repeat timer during period measurement, and display the set value of refractory period timer during single measurement.

- (9) Countdown timer: display the time until the next measurement.
- (10) The completion time of the last measurement

22.6. Preparing for NMT Monitoring

22.6.1. NMT device Connection

The following picture shows NMT cable and patient connection.



- (1) NMT sensor
- (2) Proximal electrode (red)
- (3) Distal electrode (black)
- (4) NMT Sub-machine
- (5) NMT plug-in module

22.6.2. Preparing the Patient Skin

Good electrode-to-skin contact is important for good signal quality. Before applying the electrodes, clean the application site of oil and dirt and avoid placing the electrodes over excessive body hair or lesions. Insufficient cleaning of the skin can cause high skin impedance which could cause the stimulation to stop.

- 1. Select sites with intact skin, without lesion of any kind.
- Shave hair from skin at chosen sties.
- 3. Gently rub skin surface at sites to remove dead skin cells.
- 4. Clean the skin where the electrode is placed with soapy water.
- 5. Dry the skin completely before applying the electrodes.

22.6.3. Placing the Electrodes and Sensor

22.6.3.1. Applying the Electrodes

Anatomical stimulation sites are chosen based on:

- their accessibility during surgery
- the ability to observe the neuromuscular response
- the nerve should be a suitable distance from the responding muscle to prevent direct muscle stimulation

Anatomically ideal stimulation sites:

Targeted Nerve	Affected Muscle	Contracting Appendage
Ulnar nerve	adductor pollicis muscle	Thumb
Posterior tibial nerve	flexor halluces brevis muscle	Big toe
Facial nerve (Zygomatic Branch)	orbicularis oculi muscle	Eye lid
Facial nerve (Temporal Branch)	corrugator supercili muscle	Eye brow

Electrode placement relies on the distal electrode (black electrode clip) to be as close to the targeted nerve as possible in order to effectively depolarize the nerve. The proximal electrode (red electrode clip) should be away from the targeted nerve.

22.6.3.2. Applying the Sensor

The tri-axial accelerometer should be attached to the contracting appendage of the patient, to measure the strength of the contraction resulting from the applied electrical stimulus. The accelerometer is only used in the Train-of-Four (TOF), Double Burst (DB), Post Tetanic Count (PTC), Supra Maximal Current (SMC) and Auto Modes, to facilitate monitoring of the efficacy of the Neuromuscular Blocking Agent.



- When placing the electrodes, make sure that they do not touch each other.
- If the electrodes are placed incorrectly, wrong nerves are stimulated and this causes wrong muscle response.
- When multiple nerves are stimulated, the measured response may be affected by activity of other muscles.

- If the stimulation electrodes are placed very close to the palm of the hand, the muscles are stimulated directly by the stimulation pulses.
- If the current is too strong, it may stimulate the muscles too much.
- Moving or touching the patient during measurement may cause incorrect results.
- Make sure that the NMT cables do not contact external pacemaker or catheter wires.
- To avoid unintentional electrical shocks always make sure that the NMT stimulation has been stopped before touching the electrodes.
- Take care to handle the NMT sensor, avoiding forcefully striking the sensor.
- After repositioning the patient, check the NMT sensor application site and ensure that the sensor is still properly applied and the thumb can move freely.



NOTE:

- Correct positioning of the electrodes is important. Small displacements may result in considerable changes in stimulation current requirements. Furthermore, the electrodes must be positioned in such a way to avoid direct stimulation of the muscle.
- It has been found that slight pressure on the electrodes may improve the stimulation considerably. Therefore, taping the electrodes to the skin may be advisable.
- The more distal the sensor is placed on the thumb, the stronger the acceleration signal. This effect can be used to adjust the signal strength.
- During NMT measurement, the arm applied with NMT electrodes and sensor should be kept immobile during the whole procedure.

22.7. Starting NMT monitoring

The NMT Sub-machine uses TOF mode to measure by default. You can start NMT monitoring with the following steps:

 Plug the NMT cable into the NMT Sub-machine and NMT plug-in module, and enter NMT measurement mode.

- Select [menu/MODE] button on the NMT Sub-machine, select stimulation mode.
- 3. Short press ► II button on the NMT Sub-machine, start a single measurement manually. If long press ► II button, it will start period measurement automatically.



NOTE:

- Stop NMT measurements if you need to change the NMT settings.
- When removing the sensor after finishing measurement, do not pull the sensor cable.

22.8. Stopping NMT Period Monitoring

Long press ▶ II button on the NMT Sub-machine to stop NMT measurement.

22.9. Setting NMT

22.9.1. Setting the Stimulation Current

To set the stimulation current, follow these steps:

- Sliding the adjustment wheel of the NMT Sub-machine can adjust the current, and the current setting will flash and start to adjust accordingly.
- 2. Press **[enter/Hz]** on the NMT Sub-machine to confirm the current change within 2 seconds.

22.9.2. Setting the Stimulation Mode

Press [menu/MODE] button on the NMT Sub-machine to toggle between different Stimulation Modes, and select a stimulation mode you need.

22.9.3. Setting the Stimulation Frequency

In TWI or TET stimulation mode, you can follow steps below to set the stimulation

frequency. In TWI mode, the default stimulation frequency is 2Hz. In TET mode, the default stimulation frequency is 50Hz.

- 1. Press **[enter/Hz]** button on the NMT Sub-machine to switch between different stimulation frequency.
- 2. Set the stimulation frequency as required.

22.9.4. Setting the Repeat Timer

In the three modes of TOF, DB and PTC, you can set the repeat timer during period measurement. The setting steps are as follows:

- Press and hold the [menu/MODE] button on the NMT Sub-machine and wait for the setup menu to appear.
- 2. Sliding the adjustment wheel of the NMT Sub-machine to **【NMT Settings】**, and press **【enter/Hz】** to enter the NMT setup menu.
- 3. Sliding the adjustment wheel of the NMT Sub-machine again to 【Repeat Timers】, and press 【enter/Hz】 to enter the Repeat Timer setup menu.
- 4. Set the repeat timers in TOF, DB and PTC modes respectively as needed.

22.9.5. Setting the Refractory Timer

The three modes of TOF, DB and PTC will be affected by the delay of the refractory period in a single measurement. NMT Sub-machine provides a safe period to prevent the user from repeatedly applying stimulation while the nerve synapse recovers from the previous stimulation.

The default refractory delays for the three modes are as follows:

• TOF: 15s

DB: 1 min

• PTC: 2 min

You can set the refractory timer as needed. The setting steps are as follows:

- Press and hold the 【menu/MODE】 button on the NMT Sub-machine and wait for the setup menu to appear.
- Sliding the adjustment wheel of the NMT Sub-machine to 【NMT Settings】,
 and press 【enter/Hz】 to enter the NMT setup menu.

- Sliding the adjustment wheel of the NMT Sub-machine again to 【Refractory Timers】, and press 【enter/Hz】 to enter the Refractory Timer setup menu.
- 4. Set the refractory timers in TOF, DB and PTC modes respectively as needed.

22.10. NMT Troubleshooting

This section lists the problems that might occur. If you encounter the problems when using the equipment or accessories, check the table below before requesting for services. If the problem persists, contact your service personnel.



NOTE: For the physiological and technical alarm messages, see *D Alarm Message*.

Problem	Corrective Actions	
Do not see NMT parameter	1. Check that the NMT is set to display in the 【Screen	
area on the screen	Layout I menu. For details, see 3.7.2 Setting Display	
	Screen.	
	2. Check whether the NMT parameter switch is on or	
	not. For details, see 3.7.1 Setting Parameters.	
NMT measurement fail	Check that the quality of electrodes or if the	
	electrodes are properly applied.	
	2. Replacing the electrodes if necessary.	
NMT measurement is	1. Do not touch the arm where electrodes are applied.	
disturbed	2. Check that the electrodes and sensors are properly	
	applied.	
Unable to search for	Check that the electrodes are properly applied. Also,	
supramaximal current	supramaximal current may not be found if the patient is	
	already relaxed.	

Chapter 23 Review

23.1. Introduction

You can know how the patient's condition is developing through reviewing interface to check the trend data, events, waveforms, and so on. You can also view the trend data through the OxyCRG screen to know the changes in the patient's condition.



■ Changing the date and time will affect the storage of trends and events and may result in data loss.

23.2. Reviewing Page

The review page contains graphic trends and tabular. The review page where each submenu is located displays patient trend data in different forms.

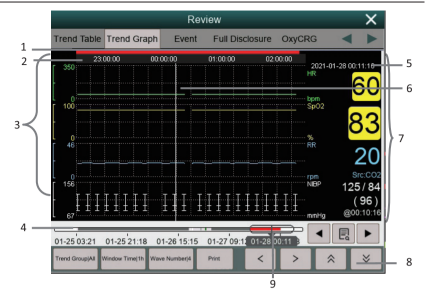
23.2.1. Accessing the Review Page

Choose one of the following methods to enter the review page

- > Select [Review] quick key.
- Select 【Main Menu】 quick key→from the 【Review】 column select the desired option.

23.2.2. The structure of review page

The review pages have similar structure. We take the graphic trends review page as an example. These contents will not be introduced in each review page.



- (1) Event type indicator: Different color blocks match different types of events.
 - Red: high priority alarm event
 - Yellow: medium priority alarm event
 - Cyan: low priority alarm event
- (2) Current window time line: indicates the time length of the current window.
- (3) Waveform area: display trend curves. The color of trend curves is consistent with the color of parameter labels.
- (4) Time line:
 - can be moved within this time length.
 - Different color blocks at the time line indicate alarm events of different types.

 The color of the color block is consistent with the color of the event identifier.
- (5) Cursor time
- (6) Cursor
- (7) Waveform area: displays the parameter value at the cursor time.
- (8) Button area
- (9) Slider: indicates the position of current window time in the entire time length. Dragging this button left or right enables you to locate the trend data at a specific time and also refreshes trend data in current window accordingly.

23.2.3. Symbols on Review Pages

The following table lists the symbols on review pages.

Symbol	Description
Ф	Slider: indicates the position of current window time in the entire time length. Dragging this button left or right enables you to locate the trend data at a specific time and also refreshes trend data in current window accordingly.
◀ or ▶	Goes to the previous or next event.
	Event list: displays events in a chronological order. The most recent event is displayed at the top. The number of asterisk symbols (*) before an event an event matches alarm priority.

23.2.4. Common Operations of Review Page

This section describes common operations for all review pages.

23.2.4.1. Browsing Trend Data

In review page, the user can browse trend data in one of the following ways:

- ♦ Move the slider □.
- Move the cursor.
- ♦ Slide page.

23.2.4.2. Viewing events

View the events in one of the following ways:

- ♦ Select to open the event list. You can select the event you want to view from the event list.
- ◆ Select or to view the previous or next events. In event list, events are displayed in a chronological order. The most recent event is displayed at the top. The number of asterisk symbols (*) before an event an event matches alarm priority as follow:
 - > ***: high priority alarm
 - ➤ **: medium priority alarm
 - *: low priority alarm

23.2.5. Tabular Trends Review Page

The trend table review page displays how the patient's physiological parameter trend is developing in a tabular manner.

23.2.5.1. Entering the Tabular Trends Review Page

Choose one of the following methods to enter the tabular trends review page:

- ◆ Select 【Review】 quick key→Select 【Trend Table】 submenu.
- ◆ Select [Main Menu] quick key→from [Review] column to select [Tabular Trends].

23.2.5.2. Selecting the Trend Group

The method for selecting trend groups is as follows:

- 1. Choose one of the following methods to enter the tabular trends review page:
 - ◆ Select 【Review】 quick key→Select 【Tabular Table】 submenu.
 - ◆ Select [Main Menu] quick key→from [Review] column to select [Tabular Trends].
- 2. Select 【Trend Group】 button→Select 【Select Trend Group】 submenu.
- 3. Select the displayed parameter combination as required.

23.2.5.3. Editing the Trend Group

The trend group defines the trend data displayed on the tabular trends review page. If you have selected a 【Trend Group】 other than 【All】 and 【Standard】, you can edit the trend group. To do so, follow this procedure:

- 1. Enter the tabular trends review page by either of the following ways:
 - ◆ Select 【Review】 quick key→Select 【Trend Table】 submenu.
 - ◆ Select [Main Menu] quick key→from [Review] column to select [Tabular Trends].
- 2. Select **Trend Group** button.
- 3. Select the trend group submenu to edit.
 - ◆ Add parameters: select desired parameters from the 【Choices】

column on the left and select [Add].

- ◆ Delete parameters: select desired parameter from the 【Selected】 column on the right and then select 【Delete】.
- ◆ Move the position of parameters: select desired parameters from the 【Selected】 column on the right and select 【Move Up】, 【Move Down】, 【Move To Top】 or 【Move To Bottom】.

Selecting **[Default Config]** will resume the trend group setting to factory defaults.



CAUTION:

- When 【Trend Group】 is set to 【All】 or 【Standard】, you cannot edit the trend group.
- ECG parameter and waveform are always displayed in the first row on the trend page. It cannot be deleted or moved.

23.2.5.4. Changing the Resolution of Trend Data

The resolution of tabular trends defines the interval of displaying trend data. High resolution is especially suited for neonatal monitoring, where the clinical situation may change very quickly. In adult monitoring, where the patient's status typically changes more gradually, a low resolution may be more informative.

To change resolution, follow this procedure:

- 1. Enter the tabular trends review page.
- 2. Select [Sample Rate].
 - > **[5s or 30s]**: The trend of parameters in the last 6 hours was observed at intervals of 5 seconds or 30 seconds.
 - > [1 min, 5 min, 10min, 15min, 30min, 1h, 2h, 3 h]: According to the selected time interval, observe the parameter trend of the last 180 hours.
 - NIBP: The tabular trends show the values of each parameter at the measurement time of NIBP parameters.

23.2.6. Graphics Trends Review Page

The graphic trends review page displays trend data in a graphic form.

23.2.6.1. Entering the Graphic Trends Review Page

Choose one of the following methods to enter the graphic trends review page:

- ◆ Select 【Review】 quick key→Select 【Trend Graph】 submenu.
- ◆ Select [Main Menu] quick key→from [Review] column to select [Graphic Trends].

23.2.6.2. Selecting the Trend Group

For more information, see 23.2.5.2 Selecting the Trend Group.

23.2.6.3. Editing the Trend Group

For more information, see 23.2.5.3 Editing the Trend Group.

23.2.6.4. Changing the Window Time

To set the length of time for each screen to display data as follows:

- 1. Enter the graphic trends review page.
- 2. Select [Window Time].
 - ♦ 【8min、30min】: Each screen displays trend data for the set time, and you can observe the trend in the last 6 hours.
 - ◆ 【1h、2h、4h】: Each screen displays trend data for the set time, and you can observe the trend in the last 180 hours.

23.2.6.5. Setting the Number of Waveforms

Follow these steps to select the number of waveforms to display in the graphic trends:

- 1. Enter the graphic trends review page.
- 2. Select [Wave Number].

23.2.7. Events Review Page

The monitor stores alarm events and system events in real time. Alarm event types include physiological alarm event. When an alarm event occurs, the monitor will store the values of relevant parameters at the time of occurrence and the relevant waveforms for 16 seconds before and after the time of occurrence.



CAUTION:

■ A sudden loss of power has no impact on the events stored.

23.2.7.1. Entering the Events Review Page

Choose one of the following methods to enter the events review page:

- ◆ Select 【Review】 quick key→Select 【Event】 submenu.
- ◆ Select [Main Menu] quick key→from [Review] column to select [Events].

The event review page displays a list of events in the order in which they occurred. The most recent event is displayed at the top. The number of asterisk symbols (*) before an event an event matches alarm priority.

The event identifier on the left side of the alarm event displays different types of events with different color blocks:

- Red: high priority alarm event
- ♦ Yellow: medium priority alarm event
- Cyan: low priority alarm event

The number of currently events and the total number of filtered events are displayed at the top right corner of the event list. For example, 3/10 indicates that there are a total of 10 selected events, and currently there are 3 events.

23.2.7.2. Configuring the Filter

You can filter events by time, alarm priority, parameter category or event type. To configure the filter, follow this procedure:

- 1. Enter events review page to switch on **[Filter]**.
- 2. Select **[Filter Setup]** and set the desired filter criterion. Events after filtering will be displayed in the event list.



CAUTION:

■ If 【Filter】 is not switch on, the relevant setting in 【Filter Setup】 will not take effect.

23.2.7.3. Viewing Event Details

To view waveforms and parameter values at the selected event time, follow this procedure:

- 1. Enter the event review page.
- 2. Select **[Detail]** . You can perform the following operations on this page:
 - ◆ Set [Wave Speed], [Record] and [ECG Gain].
 - ◆ Select 【Overview】 to return to the compressed waveform page.



CAUTION:

■ Please ensure that the best ECG lead with largest waveform amplitude and the highest signal-to-noise ration is selected. Choosing the best ECG lead is very important to recognize cardiac beat, classify cardiac beat and recognize ventricular fibrillation.

23.2.8. Full Disclosure Review Page

On the Full Disclosure review page, you can review waveform data 72 hours. You can view compressed waveforms, full waveforms and numeric values.

23.2.8.1. Entering the Full Disclosure Review Page

Choose one of the following methods to enter the Full Disclosure review page:

◆ Select 【Review 】 quick key→Select 【Full Disclosure 】 submenu.

◆ Select [Main Menu] quick key→ from [Review] column select [Full Disclosure].

23.2.8.2. Selecting Compressed waveforms

To review compressed waveforms, you must first select which parameters to store and display. Follow these steps:

- 1. Enter the Full Disclosure review page.
- 2. Select [Setup] submenu to enter [Full Disclosure Setup] page.
- 3. Select **[Storage]** submenu and select the desired waveforms to be stored.
- 4. Select **[Display (Maximum: 3)]** submenu and select the desired waveform to be displayed from the stored waveforms.



CAUTION:

■ If more waveforms are selected in the 【Storage】 column, the storage time of these waveforms will be shortened due to the limitation of memory size. The waveforms may not be stored for 72 hours. Please exert caution when selecting waveforms.

When an alarm occurs, the band on the compressed waveform at the alarm time will use different shading to indicate different alarm levels.

- Red: high priority alarm
- ♦ Yellow: med priority alarm
- Cyan: low alarm priority

23.2.8.3. Setting Gain and Duration

To set the length of time each compressed waveform is displayed and the ECG waveform height. Follow these steps:

- 1. Enter the holographic waveform review page.
- 2. Select **[Duration]** to set the length of time for each compressed waveform display.

3. Select **[Gain]** to set ECG waveform gain.

23.2.8.4. Viewing Details of Compressed Waveforms

To view the full waveforms and numeric values of compressed waveforms, follow this procedure:

- 1. Enter the holographic waveform review page.
- 2. Select **[Details]** . You can perform the following operations on this page:
 - ◆ Set [Wave Speed], [Record] and [Gain].
 - Select **[Overview]** to return to the compressed waveform page.

23.2.9. OxyCRG Review Page

You can review up to 48 hours' trend curves and compressed waveforms on the OxyCRG review page.

23.2.9.1. Entering the OxyCRG Review Page

Choose one of the following methods to enter the OxyCRG Review Page:

- ◆ Select 【Review】 button on the OxyCRG Review Page.
- ◆ Select [Review] quick key→Select [OxvCRG] submenu.
- ◆ Select [Main Menu] quick key→enter [Review] column →Select [OxyCRG].

23.2.9.2. Changing the Resolution of Trend Curves

To set the resolution of trend curves, follow this procedure:

- Enter the OxyCRG review page.
- 2. Set **[Zoom]**.

23.2.9.3. Setting the Compressed waveform

To set the compressed waveform, follow this procedure:

- 1. Enter the OxyCRG review page.
- 2. Set [Waveform].

Chapter 24 Clinical Assistive Applications (CAA)

Clinical Assistive Applications (CAA) is the comprehensive analysis and centralized presentation of the existing measurement results of the monitor, and it is the electronic application of common clinical guidelines and tools.

The main purpose of clinical assistive applications is to improve the working efficiency of doctors. It is not used for diagnosis and cannot replace medical staff to make decisions.

24.1. Early Warning Score (EWS)

The Early Warning Score can help identify early signs of deterioration in patients and is an early warning indicator for critical or potentially critical illness. The Early Warning Score System obtains corresponding scores by monitoring and observing patients' vital signs and states, and gives corresponding suggestions on solutions according to the scoring results.

The monitor provides the following scoring system:

- National early warning score (NEWS);
- Modified early warning score (MEWS).

NEWS and MEWS are aggregate scoring systems that score each parameter selected and then calculate an aggregate score. The grading of each parameter is color-coded to indicate the corresponding critical level. Provide action when the total score exceeds the range. NEWS and MEWS scoring systems are applicable for adult only.



WARNING:

- The results of the Early Warning Scores and the action measures provided are for reference only and cannot be directly used as a basis for clinical treatment.
- Early warning scores cannot be an indicator for predicting patient development and overall prognosis; it is not a tool for comprehensive clinical judgment and cannot completely replace clinicians' assessment of patients.
- Early warning scores are not available for pregnant women, those with COPD (chronic obstructive pulmonary disease), and those under 16 years of age.

24.1.1. Parameters participating in the scoring

The following table lists parameters used for evaluation by each scoring system:

National early warning score (NEWS)	Modified early warning score (MEWS)
RR, SpO ₂ , Oxygen Supply, TEMP, NIBP-S (BP-S), HR/PR, Consciousness (AVPU)	RR, TEMP, NIBP-S (BP-S), HR/PR, Consciousness (AVPU)

24.1.2. Display EWS Parameter Area

The steps to display the EWS parameter area are as follows:

- 1. Enter the **[Screen layout]** page in one of the following ways:
 - Select 【Screen Setup】 quick key → select 【Screen layout】 submenu.
 - ➤ Select [Main Menu] quick key → select [Screen layout] from [Display] column.
- 2. Select the parameter area where you want to display EWS, and select **[EWS]** from the pop-up parameter list. The following figure shows an example of EWS parameter area. The display of EWS parameter area will vary according to the settings:



- (1) Name of scoring system
- (2) Total score, the color of the circle indicates the current score level.
- (3) This scoring time
- (4) Indicator of scoring grade: increasing from top to bottom according to the degree of early warning danger. The current level is shown in the box.
- (5) Score interval

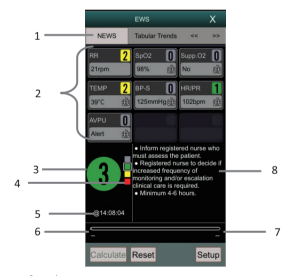
(6) Countdown to next scoring

24.1.3. Entering EWS Screen

In addition to the EWS parameter area, this monitor also provides an independent EWS interface. Select one of the following methods to enter the EWS interface:

- > Select **[EWS]** parameter area.
- Select 【Screen Setup】 quick key → select 【Screen Select】 submenu
 → select 【EWS screen】.
- Select 【Main Menu】 quick key → select 【Screen Select】 from 【Display】 column → select 【EWS screen】.

Take NEWS as an example, the EWS screen displays as follows. The actual interface display will vary according to the selected scoring system and settings.



- (1) Name of scoring system
- (2) Parameter area: displays the parameter value and score of a single parameter. The keyboard icon indicates that the parameter value comes from manual input.
- (3) Total score. The color of the circle indicates the current score level.
- (4) Indicator of scoring grade: increasing from top to bottom according to the degree of early warning danger. The current level is shown in the box.

- (5) This scoring time
- (6) Countdown to next scoring
- (7) Scoring interval
- (8) Recommended action measures

24.1.4. Calculation of Scores

Calculate the score as follows:

- Select [Reset] to clear the last scoring result, and refresh the parameter values automatically obtained from the monitor and the scoring of corresponding parameters.
- 2. Measure or manually enter parameter values of other parameters.
- 3. Select **[Calculate]** to obtain the scoring results.



CAUTION:

- Before each scoring, please press the 【Reset】 key to clear the last scoring result.
- The keyboard symbol to the right of the parameter value indicates that the parameter value is manually entered.
- You can calculate the score only when the parameter values of all the parameters involved in the calculation are valid.

24.1.5. Automatic Scoring

Set up the automatic scoring method as follows:

- 1. Select the **[Setup]** button from the scoring interface of EWS.
- 2. In the **[Auto Scoring]** area, select as required:
 - ◆ 【Interval Mode】: The monitor automatically calculates the score according to the set time interval.
 - ♦ 【NIBP】: The monitor automatically calculates the score after each NIBP measurement is completed.
 - ◆ 【Alarm】: Scores are automatically calculated after physiological alarm occurs to scoring parameters.

24.1.6. Setting EWS

24.1.6.1. Select a scoring system

The monitor is equipped with a default scoring system. You can select other scoring systems as required, as follows:

- 1. Select the **Setup** button in the scoring interface of EWS.
- 2. Set [Score].

24.1.6.2. Set scoring interval time

When the **[Interval Mode]** in the **[Auto Scoring]** area is selected, the measurement interval for automatic scoring can be set as follows:

- 1. Select the **[Setup]** button from the scoring interface of EWS.
- 2. Set [Interval].

24.1.6.3. Set the invalid time of the parameter

For manually entered parameter values, you can set the invalid time of the parameter values as follows:

- 1. Select the **Setup** button from the scoring interface of EWS.
- 2. Set [Manual Data Timeout].

24.1.7. EWS score review

In EWS screen, select **【Tabular Trends】** submenu or **【Graphic】** submenu to view the parameter values and scores of all measurement parameters and input parameters.

24.2. Glasgow Coma Scale (GCS)

The Glasgow Coma Scale (GCS) is based on the content of the Glasgow Coma Index (1974_Lancet_ Teasdale Assessment of Coma and Impaired Consciousness-A Practical Scale). GCS can be used for coma patients caused by various causes to objectively express the state of consciousness of patients. The GCS score includes three aspects: eye opening, verbal response and motor response. The scores of the three

aspects are the coma index.

The GCS score is applicable for adults and pediatric.



CAUTION:

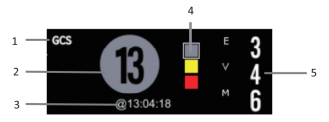
- The results of the GCS score are for reference only, please use other clinical evidence for diagnosis.
- GCS is not applicable for patients with sedation or muscle relaxants, artificial airways, drunkenness, and status epilepticus.
- GCS is not applicable for language impediments, deaf people, and people with mental disorders.
- When applied to children younger than 5 years old or elder people who are slow, the GCS score might be low.

24.2.1. Display GCS parameter area

The monitor can display GCS parameters and status in the parameter area. The steps to display the GCS parameter area are as follows:

- 1. Use one of the following methods to enter the **[Screen layout]** page:
 - Select 【Screen Setup】 quick key → select 【Screen layout】
 submenu.
 - ➤ Select [Main Menu] quick key → select [Screen layout] from [Display] column.
- Select the parameter area where you want to display the GCS and select
 【GCS】 from the pop-up parameter list.

The figure below shows an example of the GCS parameter area. The display of the GCS parameter area varies depending on the setting.



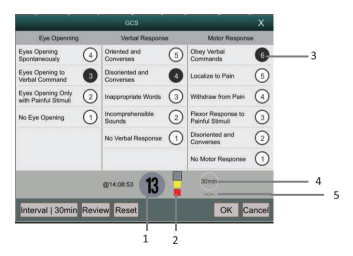
- (1) Scoring system name
- (2) The total score (coma index), the color of the circle indicates the current scoring level.
- (3) This scoring time
- (4) Indicator of scoring grade: increasing from top to bottom according to the degree of early warning danger. The current level is shown in the box.
- (5) Subscores:
 - Eye opening
 - Verbal response
 - Motor response

24.2.2. Entering GCS screen

Choose one of the following methods to enter the GCS interface:

- > Select **[GCS]** parameter area.
- ➤ Select [Main Menu]quick key → select [GCS] from [CAA]column.

The GCS interface is shown below. The actual interface display will vary depending on the settings.



- (1) The total score, the color of the circle indicates the current scoring level.
- (2) Indicator of scoring grade: increasing from top to bottom according to the degree of early warning danger. The current level is shown in the box.
- (3) Subscores

- (4) Total score invalid time
- (5) Score invalidation countdown

24.2.3. Performing GCS Scoring

Follow these steps to perform a GCS score:

- In the 【GCS】 menu, select an option corresponding to the patient status from the three areas of 【Eye opening】, 【Verbal Response】, and 【Motor Response】.
- 2. Select **[OK]** to confirm the score.

The table below lists the default score ranges and colors for each scoring level:

Level	Score	Background	Description
	range	color	
Mild	12-15	Gray	The brain function is normal or
			mildly damaged.
Moderate	5-11	Yellow	The brain function is suffered
			from moderate to severe damage.
Severe	3-4	Red	Can be brain death or remain
			vegetative.

24.2.4. Set GCS score invalid time

Select the 【Invalid Time】 button in the GCS menu to set the invalid time of the GCS score. If the set score interval is reached without re-scoring, the original score is invalidated and displayed in a hollow word.

24.2.5. Set the GCS score threshold

The coma score threshold was set as follows:

- Select 【Main Menu】 quick key → select 【Maintenance】 from 【System】 column→input maintenance password→enter.
- 2. Select 【CAA】 → 【GCS】 submenu.
- 3. Set the high and low limits of **[Mild]**, **[Moderate]**, and **[Severe]** respectively.

24.2.6. GCS score review

In the GCS menu, select the **Review** button to enter the **Review** menu, and view the GCS score trend from the **Trend Table** page.

24.3. BoA View

According to the characteristics of different stages of surgery, the Balance of Anesthesia (BoA) View provides corresponding auxiliary functions in the preoperative induction, intraoperative maintenance and postoperative resuscitation phases, which helps the doctors to monitor the patient's status during intubation, anesthetic induction, intraoperative anesthesia maintenance, and postoperative recovery.

24.3.1. Entering the BoA View

Choose one of the following methods to enter the BoA View interface:

- ➤ Select 【Screen Setup】 quick key → select 【Screen Select】 submenu
 - → select 【BoA View】.
- Select 【Main Menu】 quick key → Select 【BoA View】 from 【CAA】
 →column.

BoA View contains the following three pages:

- Induction page
- Anesthesia Maintenance page
- Recovery page

24.3.2. Induction

In BoA View page, select **【Induction】** submenu to enter **【Induction】** page. You can check parameter short trends and apnea time from the **【Induction】** page.

Select **[Induction]** to start apnea detection, mark the induction event, start an NIBP continuous measurement, and set reference value.

The following figure shows the 【Induction】 page:

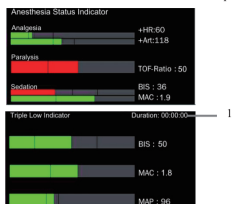


- (1) Cursor: indicates the current apnea time.
- (2) Apnea indicator: apnea time scale.
- (3) Parameter short trends area: you can select the parameters you want to view. See *24.3.5.1 Selecting the short trend parameters* for details.
- (4) Timer: displays the time elapsed since the timer was started.

24.3.3. Anesthesia Maintenance

You can check the parameter trends, anesthesia status and triple low status of patients in Anesthesia Maintenance Page. Select 【Anesthesia Status】 or 【Triple Low】 to choose the type of indicator.

- The anesthesia status indicator reflects the patient's status in terms of pain, consciousness, and neuromuscular blockage.
- The triple low status displays BIS, MAC and mean pressure of patients. The simultaneous occurrence of low MAC and low mean pressure during surgery is a risk factor for increased postoperative mortality and prolonged hospital stay. When low BIS occurs at the same time, the risk of postoperative mortality and prolonged hospital stay will further increase.



The following figures show the anesthesia status indicator and triple low indicator:

- (1) Accumulated duration of triple low states: when the measured values of three parameters are lower than the threshold, it can be judged as triple low. Refer to 24.3.5.5 Setting the Thresholds for Triple Low Status Parameters for the method of setting the threshold of the triple low status parameters.
 - The three parameter bars of the anesthesia status indicator respectively indicate pain, consciousness and neuromuscular blockage. You can refer to 24.3.5.5 Setting the Thresholds for Triple Low Status Parameters to select parameters for the anesthesia status indicator.
 - The three parameter bars of the anesthesia status indicator respectively indicate BIS, MAC and mean blood pressure.
 - The filling length of the parameter bar indicates the parameter value.
 - The color of the parameter bar indicates parameter status: green indicates that parameter value is within normal range. Red indicates that parameter value is beyond normal range. Gray indicates that parameter value is unavailable or invalid. The black lines on the parameter bars indicate the normal range of parameter.

24.3.4. Recovery

You can view parameter short trend from the recovery page.

Select **[Aldrete Score]** can re-scoring the patient's condition, and then close **[Aldrete Score]** to view the new scoring result to understand the latest status of others.



WARNING:

■ The Alderte score and recommendation is for reference only. Clinicians must make the decision of discharging the patient from recovery according to the patient's actual situation.

24.3.5. Setting the BoA View

You can select short trend parameter and set that anesthesia status indicator and triple low indicator.

24.3.5.1. Selecting the short trend parameter

You can view the short trends of up to 5 parameters from the induction page and recovery page. Choose one of following method to select the parameters you want.

- Select 🌣
 - Select **[Induction]** or **[Recovery]** submenu, select the parameter you want to display in induction page or recovery page.
 - > Select **[Defaults]** to resume the default setting.
- Select a parameter on the screen, and set which parameter you want to display in this position.

24.3.5.2. Selecting the Anesthesia Status Indicator Parameters

In the Anesthesia maintenance stage, you can evaluate the pain status of patients by HR and/or blood pressure. You can evaluate the consciousness status of patients by BIS or MAC values. The muscle relaxant status is always evaluated by NMT. You can select parameters for anesthesia status indicator by following procedures:

- 1. Select .
- 2. Select [Maintenance] submenu.
- 3. Set [Analgesia] and [Sedation] in the [Parameter] area.

24.3.5.3. Setting the reference values of HR and systolic blood pressure

The current HR and systolic blood pressure reference values are displayed as white lines in short trends area. You can set these two parameter reference values manually through these procedures:

- 1. Select [Set Reference].
- 2. Select **[OK]** to save the current measurement values as reference values.

You can also manually set the reference values for heart rate and systolic blood pressure using the following methods:

- 1. Select .
- 2. Select [Maintenance] submenu.
- 3. Set [HR] and [BP-S] in the [Reference] area.

24.3.5.4. Setting the Thresholds for Anesthesia Status Parameters

To set the thresholds for the anesthesia status parameter, follow this procedure:

- 1. Select .
- 2. Select [Maintenance] submenu.
- 3. Set the high limit and low limit of every parameter in **【Threshold】** area.

24.3.5.5. Setting the Thresholds for Triple Low Status Parameters

To set the thresholds for the triple low status parameter, follow this procedure:

- 1. Select 🗖.
- 2. Select 【Maintenance】 submenu → 【Triple Low】 submenu.
- 3. Set the thresholds of BIS, MAC and MAP.

24.4. Sepsis View

The Sepsis View function is based on Sepsis-3 (Third International Consensus Definitions for Sepsis and Septic Shock) and Surviving Sepsis Campaign: International Guidelines for Management of Severe Sepsis and Septic Shock: 2012 (SSC Guideline 2012). It assists you in recognizing the early signs and symptoms of sepsis by comparing the state of patient to the defined criteria and then guides you through the recommended treatment protocol.



NOTE:

- Due to the space limitations on the screen, the recommendations of the Sepsis View function cannot be as detailed as a complete procedure.
- Sepsis View is not a diagnostic or treatment tool, and cannot replace the professional judgment of a physician.
- The Sepsis View function is only applicable for adult patients with sepsis and suspected sepsis.

24.4.1. Entering the Sepsis View menu

Choose one of the following methods to enter the menu:

Select 【Main Menu】 quick key → select 【Sepsis View】 from 【CAA】
column

24.4.2. Screening

According to the definition of Sepsis-3, Sepsis View supports the following two types of scoring: qSOFA scoring (quick SOFA scoring) and SOFA scoring. qSOFA score is used for rapid screening of sepsis; SOFA score is used for further screening of patients with positive qSOFA screening.

24.4.2.1. Entering the qSOFA scoring

The qSOFA score is based on the patient's respiration rate, systolic blood pressure and changes in mental state. If you are monitoring the patient's breathing and blood pressure, the monitor can automatically obtain the RR value and BP-S value for scoring, or you can manually enter the parameter values. Manually select the mental state of the patient. The monitor will automatically calculate the qSOFA score based on these parameters.

Click the RR and BP-S parameter and manually enter the parameter value. The keyboard symbol on the right side of the parameter value indicates that the parameter is input manually. If sepsis is suspected or the total score is greater than or equal to 2,

select **SOFA>>** for SOFA scoring.

Select **[Reset]** to clear the current score.



NOTE:

■ When the score value is displayed "?", it means that more parameter values are needed for scoring.

24.4.2.2. Performing the SOFA

SOFA score is systemic infection-related organ failure score or sequential organ failure assessment score. Manually select each scoring item option or enter the parameter value, the monitor will automatically calculate the total score.

To clear the current scores, select **[Reset]**. If Sepsis criteria is met, make a further check on the clinical features

24.4.3. Therapy

According to the SSC Guideilne 2012, Sepsis View provides SSC Bundles and Supportive Therapy. SSC Bundles provides the situation of the resuscitation in the first 6 hours and recommended treatments to be completed within 3 hours and 6 hours. Supportive Therapy provides the completion status of the supportive treatment project.

24.4.3.1. SSC Bundles Therapy

Select **【SSC Bundles 】** menu on Sepsis View menu to enter corresponding page. You can perform the following operations on SSC Bundles Therapy page.

- For the completed treatment items, tick to mark. The marked time is automatically recorded and displayed. You can click this time to manually change it.
- Select "..." on the right side of each item to view specific guidelines. The five-pointed star symbol "★" indicates the recommendation level of this treatment item: two five-pointed stars indicate key recommendations, and one five-pointed star indicates general recommendations.

■ Select **Reset** to clear records.

24.4.3.2. Supportive Therapy

Select **[Supportive Therapy]** submenu on Sepsis View menu to enter corresponding page. You can perform the following operations on this page:

- For the completed treatment items, tick to mark. The marked time is automatically recorded and displayed. You can click this time to manually change it.
- Select "..." on the right side of each item to view specific guidelines. The five-pointed star symbol "★" indicates the recommendation level of this treatment item: two five-pointed stars indicate key recommendations, and one five-pointed star indicates general recommendations.
- Select 【Reset】 to clear records.

24.4.4. Viewing the parameter trend

Select **【Graphic Trends】** submenu on Sepsis View menu to view the change trend of parameter of resuscitation within 8 hours.

After manually checking items on the **【SSC Bundles】** page and the **【Supportive**Therapy **】** page, corresponding event markers will be generated on the **【Graphic**Trends **】** page. Vertical lines in different colors indicate different types of event markers:

■ White: inspection

■ Blue: medication

■ Green: goal achieved

■ Purple: other

24.4.5. Setting the SSC

From the Sepsis View menu select the **[Setup]** button, you can change the following settings:

■ In the 【Screening】 area, set the positive threshold of RR and BP-S for

qSOFA.

- In the 【Unit】 area, set the unit of Bilirubin and Creatinine.
- In the 【Goal of Initial Resuscitation】 area, set the goals of the first 6 hours of resuscitation.

24.5. Rescue Mode

When rescuing a patient, you can set the monitor to rescue mode.

The rescue mode provides CPR assistant, which is convenient for you to record the medication and operation during the rescue. You can output rescue reports. The rescue mode is applicable for adult and pediatric patients.



WARNING:

- The rescue mode is not applicable for neonatal patient.
- After entering the rescue mode, all physiological and some technical alarms will be turned off.



NOTE:

- The CPR assistant function requires license support.
- Rescue records shall be operated by a special nurse and shall not affect rescue.

24.5.1. Entering the Rescue Mode

You can enter the rescue mode in any of the following ways:

- > Select **[Rescue Mode]** quick key.
- Select [Main Menu] quick key → select [Rescue Mode] from the [Alarm].

24.5.2. CPR assistant

The CPR assistant helps you record the medication and operation during the rescue.

You can record the following on the monitor:

- Start and stop rescue
- Use records of adrenalline, amiodarone, and other drugs
- Chest compressions, defibrillation and other rescue operations, such as tracheal intubation, establishing venous access, etc.

24.5.2.1. Enabling the CPR assistant

The first time you enter the rescue mode, the CPR assistant will be turned on automatically. When the CPR assistant is turned off, if you need to reopen it, select [Main Menu] quick key -> select [CPR Assistant] from the [CAA].

24.5.2.2. Rescue record

The method of recording the patient's rescue process in the CPR assistant page is as follows:

- Record the start time of rescue: select 【Start Rescue】. When entering the rescue mode, the monitor will automatically record the start time of the rescue.
- Record the drugs and doses used: select [Adrenaline], [Amiodarone] and/or [Other Drugs] according to the actual use of the patient during rescue.
- Record the rescue operation: select [Start/Pause Compression], [Defibrillation] and/or [Other Treatments] according to the actual treatment performed during the rescue of the patient.
- Record the end time of rescue: select **End Rescue**.

24.5.2.3. Saving the Rescue record

Select **Save** in the CPR assistant page to save the rescue records.

24.5.2.4. Exporting the Rescue record

You can use a USB disk to export the rescue records, as follows:

1. Insert the USB disk into the monitor's USB port.

2. Select **[Export]** to export the rescue records.

24.5.2.5. Customizing the Drugs and Operations

In addition to sodium bicarbonate, vasopressin and atropine, you can also customize five other drugs; in addition to tracheal intubation and establishing venous access, you can also customize two other operations. The methods for customizing drugs and operations are as follows:

- Select 【Main Menu]quick key → select 【Maintenance] from 【System]
 column→input maintenance password→enter.
- 2. Select 【CAA】 submenu → 【CPR】 submenu.
- Select 【Customized Drugs】 to customize other drugs commonly used in rescue.
- 4. Select **Customized Treatments** to customize other operations commonly used in rescue.

24.5.2.6. Turning off the CRP assistant

When exiting the rescue mode, 【CPR Assistant】 will be turned off automatically. In rescue mode, if need to turn off 【CPR Assistant】, choose one of the following options:

- Select the exit button in the upper right corner of the **CPR Assistant** page.
- Select 【Main Menu】 quick key → select 【Exit CPR assistant】 from the 【Alarm】.

24.6. CCHD Screening

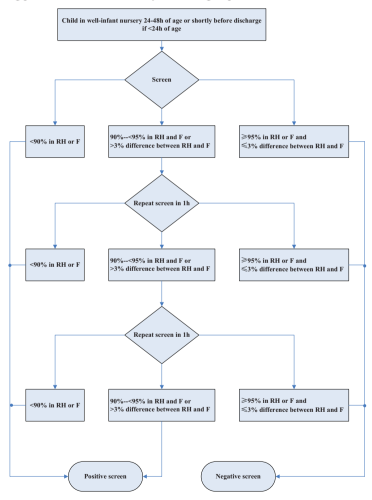
A work group (hereinafter referred to as work group) was convened with members selected by the Secretary's Advisory Committee on Heritable Disorders in Newborns and Children (SACHDNC), American Academy of Pediatrics (AAP), American College of Cardiology Foundation (ACCF) and American Heart Association (AHA). The work group recommended that screening not begin until 24 hours of life, or as late as possible if earlier discharge is planned, and be completed on the second day of life.

CCHD screening is performed by detecting the SpO_2 value of the newborn's right hand and any foot, and comparing the difference in SpO_2 value (ΔSpO_2).

CCHD screening is only applicable to neonatal patients.

24.6.1. CCHD Screening Protocol

The CCHD screening protocol for the monitor was derived from the neonatal screening procedure recommended by the work group. The flow chart is as follows:



24.6.2. Entering the CCHD Screening Page

Choose one of the following methods to enter the CCHD Screening Page:

- > Select 【CCHD】 quick key.
- Select 【Main Menu】 quick key → select 【CCHD】 from 【CAA】 column.

24.6.3. Performing CCHD Screening

You need to measure the neonate's right hand and then measure SpO_2 of one foot to complete the CCHD screening. CCHD screening procedures are as follows:

- 1. Entering CCHD interface, see **24.6.2** Entering the CCHD Screening Page for details.
- 2. Operate according to the interface prompts.

SpO₂ measurement values and screening results are displayed above the CCHD interface:

- Negative: Screening is finished.
- Positive: Failure to pass CCHD screening, echocardiography is recommended.
- Re-measurement after 1 hour: It is temporarily impossible to determine
 whether it is negative or positive. You need to wait 1 hour and then
 re-measurement.



CAUTION:

■ Please keep the patient quiet during SpO₂ measurement.

24.6.4. Re-performing CCHD Screening

If you question the results of the screening, you can re-perform the CCHD screening. Re-performing the CCHD screening will clear the current measurement data and screening results.

Select **[New CCHD]**, and select **[OK]** in the pop-up dialog box to re-measure.

Chapter 25 Calculations

25.1. Introduction

The monitor provides calculation functions. The calculated values, which are not directly measured, are computed based on the data and measurement values you enter. The calculation is independent of other monitoring functions and the object of calculation may not be the patient monitored by this monitor. The calculation operation will not affect the patients being monitored.

The following calculations can be performed on this monitor:

- Drug calculation
- ♦ Hemodynamic calculation
- Oxygenation calculation
- ◆ Ventilation calculation
- ♦ Nephridium Calculation

25.2. Safety information



WARNING:

- The dosage of drugs must be decided by the physician in charge.
- During calculation, check that the entered values are correct and the calculated values are appropriate. We assume no responsibility for any consequences caused by wrong entries and improper operations.

25.3. Drug Calculation

The monitor provides the drug calculation function.

25.3.1. Calculation Step

The Drug calculation steps are as follows:

- 1. Access drug calculation page by either of the following ways:
 - ◆ Select 【Calculations】 quick key.

- ◆ Select [Main Menu] quick key→from [Calculations] column to select [Drug] .
- Set 【Drug Name】 and 【Patient Type】. If the selected drug is affected
 by weight, switch on 【Weight Participation】 and enter the patient's
 weight.
- 3. Enter the drug-related information such as total amount, volume and dose of drugs.
- 4. Select **[Calculate]** button to calculate. Red arrow marks are displayed before the calculation results.



CAUTION:

■ If available, the patient category and weight from the patient demographics menu are automatically entered when you first access drug calculation. You can change the patient category and weight. This will not change the patient category and weight stored in the patient demographic information.

25.3.2. Checking the Titration Table

The Titration Table shows informations on the currently used drugs. You can check the dose received by the patient at different infusion rate through the Titration Table. The procedure for viewing the titration table is as follows:

- 1. Access drug calculation page by either of the following ways:
 - ♦ Select 【Calculations 】 quick key.
 - ◆ Select [Main Menu] quick key→from [Calculations] column to select [Drug].
- 2. Select **[Titration]** sub menu.
- 3. Select **[Dose Type]** at the bottom of the interface to set the unit type of drug dose in the titration table.
- 4. Select **[Step]** to set the interval between two adjacent titration table item.

You can choose the sorting method of titration table:

♦ 【Dose】: The titration table is listed in the sequence of increased drug

dose.

♦ 【INF Rate】: The titration table is listed in the sequence of increased infusion rate.

25.3.3. Drug calculation Formula

Description	Unit	Formula
Drug Amount	g series: mcg, mg, g; unit series: unit, kU, MU; mEq series: mEq;	Dose Amount = Liquid Volume × drug concentration
Liquid Volume	ml	Manual input required
Drug concentration	mcg/ml, mg/ml, g/ml, Unit/ml, kU/ml, MU/ml, mEq/ml	Drug concentration = Dose Amount / Liquid Volume
Drop Size	GTT/ml	Manual input required
Dose/hr	g series: mcg, mg, g; unit series: unit, kU, MU; mEq series: mEq;	Dose/h=Dose/min×60
Dose/min	g series: mcg, mg, g; unit series: unit, kU, MU; mEq series: mEq;	DoseMin = DoseMin
Dose/kg/hr (weight based)	g series: mcg, mg, g; unit series: unit, kU, MU; mEq series: mEq;	Dose/h= Dose/h/weight
Dose/kg/min (weight based)	g series: mcg, mg, g; unit series: unit, kU, MU; mEq series: mEq;	Dose/min= Dose/min /weight
INF rate	ml/h	Infusion rate = Dose/h/drug concentration

Description	Unit	Formula
Drip rate	GTT/min	Drip rate = infusion rate*volume per drop/60
Duration	h	Duration = drug amount/dose/h

25.4. Hemodynamic Calculations

The monitor provides the hemodynamic calculation function. The monitor can save the results of up to 20 calculations, which are displayed in groups.

25.4.1. Calculation Step

To perform hemodynamic calculation, follow this procedure:

- 1. Access hemodynamic calculation page by either of the following ways:
 - ◆ Select 【Calculations】 quick key→ 【Hemodynamics】 submenu.
 - ◆ Select [Main Menu] quick key→from [Calculations] column to select [Hemodynamics].
- Enter the correct value for each parameter. For a patient who is being monitored, the monitor takes the current measurements as input values for some parameters, and the height and weight are derived from the patient information entered.
- 3. Select 【Calculate】 to calculate the value of each output parameter. The calculated value is greater than the normal upper limit is indicated by an up arrow "↑"; the calculated value is lower than the normal lower limit is indicated by a down arrow "↓".
 - ◆ Select 【Range】 to show the normal range of each parameter.
 - Select **[Unit]** to show the unit of each parameter.

25.4.2. Input Parameters

Abbreviation	Unit	Full Name
C.O.	L/min	cardiac output

Abbreviation	Unit	Full Name
HR	bpm	heart rate
PAWP	mmHg	pulmonary artery wedge pressure
MAP	mmHg	artery mean pressure
MPAP	mmHg	mean pulmonary artery pressure
CVP	mmHg	central venous pressure
EDV	mL	end-diastolic volume
Height	cm	height
Weight	kg	weight

25.4.3. Output Parameters and calculation formula

Output Parameter	Unit	Full Name	Formula
C.I.	mL/min/m ²	cardiac index	C. I. = C. O. / BSA
BSA	m ²	body surface	BSA=HT ^{0.725} × WT ^{0.425} ×
		area	0.007184
SV	mL	stroke volume	$SV = 1000 \times C.O./HR$
SVI	mL/m ²	stroke index	SVI= SV/BSA
SVR	dyn*s/cm ⁵	systemic vascular resistance	$SVR = 79.96 \times \frac{MAP - CVP}{C. O.}$
SVRI	dyn*s*m²/c m ⁵	systemic vascular resistance index	SVRI = SVI /BSA
PVR	dyn*s /cm ⁵	pulmonary vascular resistance	$= 79.96 \times \frac{\text{MPAP} - \text{PAWP}}{\text{C. O.}}$
PVRI	dyn*s*m²/c m ⁵	pulmonary vascular	PVRI= PVR×BSA

Output Parameter	Unit	Full Name	Formula
		resistance index	
LCW	kg*m	left cardiac work	$LCW = 0.0136 \times PAMAP \times C.O.$
LCWI	kg*m/m²	left cardiac work index	LCWI = RCW×BSA
LVSW	g*m	left ventricularstrok e work	LVSW = 0.0136 ×MAP×SV
LVSWI	g*m/m²	left ventricular stroke work index	LVSWI = LVSW/BSA
RCW	kg*m	right cardiac work	$RCW = 0.0136 \times PMAP \times C. O.$
RCWI	kg*m/m²	right cardiac work index	RCWI= RCW/BSA
RVSW	g*m	right ventricular stroke work	$RVSW = 0.0136 \times MPAP \times SV$
RVSWI	g*m/m²	right ventricular stroke work index	R VSWI= RVSW /BSA
EF	%	ejection fraction	$EF = 100 \times SV / EDV$

25.5. Oxygenation Calculation

The monitor provides the oxygenation calculation function. The monitor can save the results of up to 20 calculations, which are displayed in groups:

25.5.1. Calculation Step

The oxygenation calculation steps are as follows:

- 1. Access oxygenation calculation page by either of the following ways:
 - ◆ Select [Calculations] quick key→ [Oxygenation] submenu.
 - ◆ Select [Main Menu] quick key→from [Calculations] column to

select [Oxygenation].

- Enter the correct value for each parameter. For a patient who is being monitored, the monitor takes the current measurements as input values for some parameters, and the height and weight are derived from the patient information entered.
- 3. Select 【Calculate】 to calculate the value of each output parameter. The calculated value is greater than the normal upper limit is indicated by an up arrow "↑"; the calculated value is lower than the normal lower limit is indicated by a down arrow "↓".

In the Oxygenation page, you can also perform the following operations:

- ◆ Select 【Oxygen Unit】, 【Hb Unit】 and 【Pressure Unit】, then corresponding parameter values will be automatically converted and updated accordingly.
- ◆ Select 【Range】 to show the normal range of each parameter.
- Select **[Unit]** to show the unit of each parameter.

25.5.2. Input Parameters

Input Parameter	Unit	Full Name
C.O.	L/min	cardiac output
FiO ₂	bpm	percentage fraction of inspired oxygen
PaO ₂	mmHg, kPa	partial pressure of oxygen in the arteries
PaCO ₂	mmHg, kPa	partial pressure of carbon dioxide in the arteries
SaO ₂	%	arterial oxygen saturation
PvO ₂	mmHg, kPa	partial pressure of oxygen in venous blood
SvO ₂	%	venous oxygen saturation
НЬ	g/L, g/dL, mmol/L	hemoglobin

Input Parameter	Unit	Full Name
CaO ₂	mL/dL,	arterial oxygen content
CaO2	mL/L	archar oxygen content
CvO ₂	mL/dL,	venous oxygen content
C V O 2	mL/L	, ,
VO ₂	mL/min	oxygen consumption
RQ		Respiratory quotient
ATMP	mmHg, kPa	atmospheric pressure
Height	cm, inch	height
Weight	kg, lb	weight

25.5.3. Output Parameters and calculation formula

Output Parameters	Unit	Full Name	Formula
BSA	m ²	body surface area	BSA=HT ^{0.725} × WT ^{0.425} × 0.007184
C(a-v)O ₂	mL/L, mL/dL	arteriovenous oxygen content difference	C(a-v)O ₂ =CaO ₂ ×CvO ₂
O ₂ ER	%	oxygen extraction ratio	$O_2ER = (CaO_2 - CvO_2) / CaO_2$
DO_2	mL/min	oxygen transport	DO ₂ =C.O.×CaO ₂
PAO ₂	mmHg, kPa	partial pressure of oxygen in the alveolar	PAO ₂ = T FiO ₂ ×(ATMP—water pressure) 1 —(PaCO ₂ ×1.25) Wherein the water pressure is selected to be 47mmHg (6.3kPa)

Output Parameters	Unit	Full Name	Formula
AaDO ₂	mmHg, kPa	alveolar-arterial oxygen difference	AaDO ₂ =PAO ₂ —PaO ₂
CcO ₂	mL/L, mL/dL	capillary oxygen content	$CcO_2 = Hb \times 1.34$ $+0.031 \times PAO_2$
Qs/Qt	%	venousad mixture	$Qs/Qt = (CcO_2 - CaO_2)/(CcO_2 - CvO_2)$

25.6. Ventilation Calculations

The monitor provides the ventilation calculation function. The monitor can save the results of up to 20 calculations, which are displayed in groups.

25.6.1. Calculation Step

The ventilation calculation steps are as follows:

- 1. Access ventilation calculation page by either of the following ways:
 - ◆ Select 【Calculations】 quick key→ 【Ventilation】 submenu.
 - ◆ Select [Main Menu] quick key→from [Calculations] column to select [Ventilation].
- Enter the correct value for each parameter. For a patient who is being monitored, the monitor takes the current measurements as input values for some parameters.
- 3. Select 【Calculate】 to calculate the value of each output parameter. The calculated value is greater than the normal upper limit is indicated by an up arrow "↑"; the calculated value is lower than the normal lower limit is indicated by a down arrow "↓".

On the ventilation page, you can also perform the following operations:

- ◆ Select 【Pressure Unit】, then corresponding parameter values will be automatically converted and updated accordingly.
- ◆ Select 【Range】 to show the normal range of each parameter.
- Select **[Unit]** to show the unit of each parameter.

25.6.2. Input Parameter

Input Parameter	Unit	Full Name
FiO ₂	%	percentage fraction of inspired oxygen
RR	rpm	respiration rate
PeCO ₂	mmHg, kPa	partial pressure of mixed expiratory CO ₂
PaCO ₂	mmHg, kPa	partial pressure of carbon dioxide inthearteries
PaO ₂	mmHg, kPa	partial pressure of oxygen in the arteries
TV	mL	tidal volume
RQ	/	respiratory quotient
ATMP	mmHg, kPa	atmospheric pressure

25.6.3. Output Parameter and calculation formula

Output Parameter	Unit	Full Name	Formula
PAO ₂	mmHg,kPa	partial pressure of oxygen in the alveolar	PAO ₂ = $\PFiO_2 \times (ATMP-water)$ pressure) $\PO(PaCO_2 \times 1.25)$ Wherein the water pressure is selected to be 47mmHg (6.3kPa)
AaDO ₂	mmHg,kPa	alveolar-arterial oxygen difference	AaDO ₂ =PAO ₂ —PaO ₂
Pa/FiO ₂	mmHg,kPa	oxygenation ratio	Pa/FiO ₂ = PaO ₂ /FiO ₂
a/AO ₂	%	arterial to alveolar oxygen ratio	$a/AO_2 = (100 \times PaO_2) /PAO_2$
MV	L/min	minute volume	MV=(TV/1000)×RR

Vd	mL	volume of physiological dead space	Vd=【(PaCO ₂ —PeCO ₂)×TV】 /PaCO ₂
Vd/Vt	%	physiologic dead space in percent of tidal volume	Vd/Vt= (PaCO ₂ —PeCO ₂)/ PaCO ₂ ×100%
VA	L/min	alveolar volume	VA=(TV-Vd) ×RR

25.7. Nephridium Calculation

The monitor provides the nephridium calculation function. The monitor can save the results of up to 20 calculations, which are displayed in groups.

25.7.1. Calculation Step

- 1. Access nephridium calculation page by either of the following ways:
 - ◆ Select [Calculations] quick key→ [Nephridium] submenu.
 - ◆ Select [Main Menu] quick key→from [Calculations] column to select [Nephridium].
- Enter the correct value for each parameter. For a patient who is being monitored, the height and weight are derived from the patient information entered.
- 3. Select 【Calculate】 to calculate the value of each output parameter. The calculated value is greater than the normal upper limit is indicated by an up arrow "↑"; the calculated value is lower than the normal lower limit is indicated by a down arrow "↓".
 - ◆ Select 【Range】 to show the normal range of each parameter.
 - Select **[Unit]** to show the unit of each parameter.

25.7.2. Input Parameter

Input Parameters	Unit	Full Name
URK	mmol/L	urine pstassium
URNa	mmol/L	urinary sodium
Urine	mL/24h	urine
Posm	mosm/kg	plasm osmolality
Uosm	mosm/kg	urineosmolality
SerNa	mmol/L	serumsodium
Cr	umol/L	creatinine
UCr	umol/L	urinecreatinine
BUN	mmol/L	blood urea nitrogen
Height	cm, inch	height
Weight	kg, lb	weight

25.7.3. Output Parameter and calculation formula

Output Parameter	Unit	Full Name	Formula
URNaEx	mmol/24h	urinesodium excretion	URNaEx= Urine × URNa/1000
Cosm	mL/min	osmolar clearance	Cosm=Uosm×(Urine/24/60)/ Posm
URKEx	mmol/24h	urine potassium excretion	URKEx= Urine × URK/1000

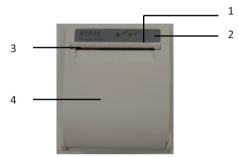
Output Parameter	Unit	Full Name	Formula
CH ₂ O	mL/h	free water	CH ₂ O =Urine/24×(1 —
C11 ₂ O		clearance	Uosm/Posm)
Na/K	0/0	sodium potassium	Na/K=URNaEx/URKEx
1144/12	/0	ratio	
U/Posm		urine to plasma	U/Posm =Uosm/Posm
0/1 00111		osmolality ratio	
BUN/Cr		blood ureanitrogen	BUN/Cr
		creatinine ratio	
CNa	mL/24h	clearance of	CNa(mL/24hrs)=URNa)×Ur
		sodium	ine/SerNa
	mL/min	creatinine clearance rate	$Ccr=(140-age) \times weight(kg)$
			/ [72×Scr(mg/dL)] or
			Ccr=[(140- age) ×
Cler			weight(kg)] / [0.818×Scr
			(umol/L)]
			Female: the calculation
			result × 0.85.
U/Cr		urine-serum	UCr/Cr
		creatinine ratio	
FENa	%	fractional excretion	FENa%=(URNa×Cr)/(SerNa
1 Liva		of sodium	×Ucr) ×100%
	l .		

^{*:} BUN/Cr is a ratio at mol unit system.

Chapter 26 Recording

26.1. Recorder

This monitor uses the thermal recorder which supports various record types. It can output the patient information, measurement data, review data and three waveforms at best.



- (1) Power indicator lamp
 - ◆ ON: The recorder works correctly.
 - OFF: The monitor is switched off.
- (2) Trouble indicator lamp
 - ON: There is something wrong with recorder, such as short of paper, door or the recorder not fasten up and something like that.
 - ◆ OFF: The recorder goes well.
- (3) Paper outlet
- (4) Recorder door

26.2. Recording Type

The records can be divided into the following types according to trigger modes:

- 1. Real-time record of manual startup;
- 2. The circular record of automatic strartup of the recording meter in line with the given time interval.
- 3. The alarm record triggered by out-of-limit parameter and so on.
- 4. Record started by manual operation and related to special function.

26.3. Starting Recordings

You can start recording by manual way through the following means:

- Press [Realtime Record] quick key below the monitor interface to start real-time recording.
- > Select **[Record]** button in the current window or above the menu to start the associated record of the special function.

The recorder can start recording automatically in the following situation:

- If the periodic recording has been started, the recorder will start recording in the set time interval. Refer to 26.6 Setting the Recorder for detailed instructions.
- When the [Alarm Switch] and [Alarm Output] of a parameter are both set to [On], once the parameter gives an alarm, the monitor will be triggered to start an alarm record.

26.4. Stopping Recordings

You can stop recording by manual way through the following means:

In the process of real-time recording, click the **[Realtime Record]** quick key.

The recorder will stop automatically in the following situation:

- The recorder has finished its task.
- The recorder is sort of paper.
- There is something wrong with the recorder

26.5. Recording Flags

When the printing of the record report is finished, there are the following flags:

- For automatically stopped recordings: Print "***END***" at the end of the report.
- For manually or abnormally stopped recordings: There is no flag printing at the end of the report.

26.6. Setting the Recorder

This section describes the definition of the main setting items. Users can refer to these definitions to select other similar setting items in the device according to their needs.

Select $[Main\ Menu] \to from [Report]$ column select $[Record\ Setup]$ to enter according corresponding menu.

26.6.1. Selecting the recorded waveform

The recorder can output up to 3 waveforms at a time. In the **【Record Setup】** menu, you can select **【Waveform 1】**, **【Waveform 2】**, **【Waveform 3】** in turn, and then select the name of the waveform in the pop-up list. Select **【Close】** to turn off the output of 1 waveform. These settings apply to real-time recording and periodic recording.

26.6.2. Setting the duration of real-time recording

When starting a real-time recording, the length of recording depends on your setting of recording duration.

- 1. Open the 【Record Setup】 menu.
- 2. Set [Record Duration] to:
 - ♦ 【8s】: Record the waveform of 4 seconds before and after the current time.
 - ◆ 【Continue】: Record the waveform 5 seconds before and after the current time until you manually stop recording.

26.6.3. Setting the interval for periodic recording

You can set a certain time interval, and the recorder automatically starts recording according to the set time interval.

- 1. Open the 【Record Setup】 menu.
- 2. Set [Cycle Record Interval].
- 3. After the setting is completed, the recorder starts each recording at the set interval.

26.6.4. Setting the duration of period recording

You can set the duration of every period recording in the following ways:

- 1. Open the **[Record Setup]** menu.
- 2. Set [Cycle Record Duration] to:
 - ♦ 【8s】: Record the waveform of 4 seconds before and after the current time.

26.6.5. Setting the recording speed

- 1. Open the 【Record Setup】 menu.
- 2. Set [Record Speed].

This setting is applicable to all recording tasks with waveforms.

26.6.6. Setting alarm recording duration

You can set how long the waveform needs to be recorded when an alarm occurs, as follows:

- 1. Open the 【Record Setup】 menu.
- 2. Set [Alarm Record Duration].
 - ◆ 【8s】: Record the waveform of 4 seconds before and after the alarm triggering time.

26.6.7. Setting NIBP Trigger Record

You can set to record the output NIBP measurement results when NIBP measurement is completed, as follows:

- 1. Open the 【Record Setup】 menu.
- 2. Set **[NIBP Trigger]** to **ON** or **OFF**.

26.7. Installing Recording Paper

If the record paper runs out, please install the record paper as the following step:

1. Press both sides of the recorder door with one hand and pull outwards to

open the recorder door;

- 2. Put the recording paper into the recorder with the thermal side which is smoother up.
- Close the door of the recorder, and pull some recording paper outside of the paper out port.
- 4. Check the position of the recording paper to ensure that the recording paper is aligned with the paper outlet.



CAUTION:

- Must use the thermo-sensitive paper that meets requirements; otherwise, it will lead to recording failure, bad-quality record or damage of thermo-sensitive printing head.
- Do not pull out the recording paper during recorder printing, otherwise the recording meter may be damaged.
- Unless for paper replacement or fault remedy, don't keep the recorder door open.

26.8. Clearing Jam Paper

While the sound of recorder operation or printing of recording meter is abnormal, please first check whether there is paper jam in the recording meter. If so, please clear it as per following steps:

- 1. Open the recorder door;
- 2. Pull out the recording paper, and cut off the wrinkle part;
- 3. Load recording paper once again and close the recording meter door.

26.9. Cleaning Recorder

After long-time service, some paper scrap and impurity will accumulate on the printing head, and affect printing quality as well as the service life of printing head and roll shaft. The recorder can be cleaned according to the following methods:

1. Before cleaning, the measures such as wearing anti-static wrist strap shall be adopted to avoid the damage to recording meter resulting from static;

- 2. Open the recorder door and pull out recording paper;
- 3. Use a tampon with some alcohol to sweep slightly the surface of thermo-sensitive parts of printing head;
- 4. After the alcohol entirely vaporizes, load recording paper once again and close the recorder's door.



CAUTION:

- Don't use any article that can damage the thermo-sensitive parts of recorder during cleaning.
- Don't heavily press the printing head of recorder.

Chapter 27 Other Functions

27.1. Analog Signal Output

The monitor has an auxiliary output port that can provide "analog signal output". Connect the monitor to equipment such as an oscillograph, and then do some associated setup, after that you can output the analog signal to the oscillograph through the port.

The setting ways of analog signal output are as below:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】→input maintenance password→Enter.
- 2. Select 【Module】 submenu→ 【Auxiliary Output】 submenu.
- 3. Select **【Analog Output】**, set the analog output signal as required.



CAUTION:

Analog output function is seldom used in clinic. If you need t know more detailed information, please contact the service personnel.

27.2. Network Settings

27.2.1. Setting the type of network

The steps for setting the network type are as follows:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】→input maintenance password→Enter.
- 2. Select 【Network Setup 】 submenu → 【Network Type 】.
- 3. Set to **[LAN]** or **[WLAN]** according to the network type used.

27.2.2. Setting the Wired Network

To set the wired network, follow this procedure:

Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance 】 →input maintenance password→Enter.

- 2. Select 【Network Setup】 submenu → 【LAN】 submenu.
- 3. Select how to get the IP address:
 - ♦ 【Obtain IP Address Automatically 】: The monitor automatically gets the IP address.
 - ◆ 【Use the Following Address】: you need to input the 【IP Address】,

 【Subnet mask】 and 【Gateway】.

27.2.3. Setting the Wireless network

To set the wireless network, follow this procedure:

- 1. Select [Network Setup] quick key.
- 2. The interface will show the surrounding wireless network, and you can choose to use the wireless network according to your needs.
- 3. If you need to manually add a wireless network, you can select the 【Add Net】 button at the bottom of the menu to set the 【SSID】, 【Security】, 【Password】 and 【DHCP】 of the network:
 - ◆ 【SSID】: Set name of the network.
 - ♦ 【Security 】: Set the encryption method.
 - **Password**: Set the password to enter the network.
 - ◆ 【DHCP】: Open 【DHCP】, and the monitor will automatically acquire the IP address; if close 【DHCP】, you need to manually enter the IP address, subnet mask and gateway.

27.2.4. Setting the wireless network frequency and antenna type

The steps for setting the wireless network frequency and antenna type are as follows:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance 】 →input maintenance password→Enter.
- 2. Select 【Network Setup】 submenu → 【WLAN】 submenu.
- 3. Set the **[Frequency]** and **[Antenna]** of the wireless network according to the usage.
 - ♦ **[Frequency]: [5G]** or **[2.4G]**.

- ♦ [Antenna]: [Build-in] or [External].
- 4. Restart the monitor

27.2.5. Connecting the Central Monitoring System (CMS)

The monitor can be connected to the central monitoring system via wired network or wireless network.

27.2.5.1. Setting the CMS IP Address

To set the IP address of CMS, follow this procedure:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】→input maintenance password→Enter.
- 2. Select 【Network Setup】 submenu → 【CMS】 submenu.
- Set the IP address of the CMS. The monitor can be received by the CMS of the IP address.

27.2.5.2. Setting the device number of the monitor

The device number of the networked monitor will be displayed when the central monitoring system and other beds are monitored. The steps for setting the device number of the monitor are as follows:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance 】 →input maintenance password→Enter.
- 2. Select 【Network Setup】 submenu → 【CMS】 submenu.
- 3. Set [Device No.] of the monitor.

Please refer to *the Central Monitoring System User's Manual* for detailed instructions.



NOTE: This monitor can only be connected to the central monitoring system provided by the manufacturer. Do not try to connect the monitor to other central monitoring system.

27.2.6. HL7 Settings

The real-time data, waveforms, and alarms of the monitor can be transmitted to the hospital's monitoring system through the HL7 protocol. The operating steps are as follows:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】→input maintenance password→Enter.
- 2. Select 【Network Setup】 submenu → 【HL7】 submenu.
- Select [Parameter], [Waveform] and [Alarm] sending function as required.
 - ◆ From 【Physiological data】 column to select monitor as 【Server】 or 【Client】. If select the monitor as 【Client】, set the 【IP】 and 【Port】 for the server receiving the real-time data and waveform. And can set 【Interval】 of data.
 - ◆ From 【Alarm Data】 column to select monitor as 【Server】 or 【Client】. If select the monitor as 【Client】, set the 【IP】 and 【Port】 for the server receiving the real-time data and waveform.

27.2.7. Connecting eGateway

The monitor can connect the eGateway server through wired and wireless networks to realize the interaction between the external devices and the monitor. The monitor has the following functions when connected to eGateway:

- ◆ Send parameters, waveforms, and events of this monitor to eGateway.
- Send data on external devices connected to the monitor to eGateway, including parameters, alarms, etc.
- Clock can be synchronized between the monitor and the eGateway.

27.2.8. Using the ADT Gateway

The ADT (admit-discharge-transfer) gateway is normally deployed in the eGateway. You can obtain patient information from the hospital ADT server through the ADT gateway.

The steps of setting the ADT gateway are as below:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】→input maintenance password→Enter.
- 2. Select 【Network Setup】 submenu → 【ADT】 submenu.
- 3. Set the IP address and port of the ADT gateway.

【ADT Query】 is switched on by default. You can load patient information to the monitor from the ADT server only when this function is enabled.

27.3. Network Printing

You can print the patient information and data through network printing.

27.3.1. Setting the Print Server

To set a network print server, please to do it follow below steps:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】→input maintenance password→Enter.
- 2. Select 【Print】 submenu → 【Printer】 submenu.
- 3. Set [Print Server IP].

27.3.2. Setting Patient Information

You can customize patient information that appears on the printed reports.

27.3.2.1. Setting Patient Information on ECG Reports

You can set the patient information on ECG reports by the following steps:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】→input maintenance password→Enter.
- 2. Select 【Print】 → 【ECG Report】.
- On the right of 【ECG Report】, select the desired patient information items.
 Patient ID, Patient name, Age and Gender are displayed on the ECG report by default.



CAUTION:

■ You can only set the patient information displays on the 【ECG Report】 from the ECG Report page. Patient information set in the 【Report Layout】 page is not displayed ECG reports.

27.3.2.2. Setting Patient Information on Other Reports

You can set the patient information on ECG reports by the following steps:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】→input maintenance password→Enter.
- 2. Select 【Print】 → 【Report Layout】.
- 3. Select desired items under **[Report Name]**. **[N/A]** indicates that the item is not displayed on the report.

27.3.3. Manually Printing Report

You can print a report manually.

27.3.3.1. Starting printing from the current page

Click 【Print】 button below the current page (such as the Review Page) to start printing.

27.3.3.2. Printing Realtime Report

You can select **[Realtime Report]** on **[Report Setup]** page to print. For more information, refer to 27.3.3.2 Printing Normal Report.

27.3.3.3. Printing Normal Report

The normal report contains the following types of reports:

- ECG Report
- Realtime Report

- Tabular Trends Report
- Graphic Trend Report

To print normal report, follow below steps:

- Select 【Main Menu 】 quick key→from 【Report】 column to select 【Report Setup】.
- 2. Select desired reports.
- Check the setting.
- 4. Select [Print].

27.3.4. Automatically Printing Report

You can start printing automatically when you set parameter alarm. To do so, follow this procedure:

- Select parameter alarm related menus such as 【Alarm】 submenu in one of the following ways:
 - > Select [Alarm Setup] quick key.
 - ➤ Select the parameter or waveform area of the desired parameters—select alarm related menus.
 - ➤ Select the 【 Parameters Setup 】 quick key→select desired parameters—select alarm related menus.
- 2. Switch on **[Switch]** and **[Alarm output]** of desired parameters. If an alarm of the parameter occurs, the printer will automatically start to print the measurement data of the parameter.

【Alarm Print】 is set to 【Recorder】 by default. If need to print automatically using printer when a parameter alarm occurs, 【Alarm Print】 need to set to 【Printer】, follow this procedure:

- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】→input maintenance password→Enter.
- 2. Select 【Alarm】 submenu→【Other】 submenu.
- 3. Set [Alarm Print] to be [Recorder].

27.3.5. Setting the Reports

This section described how to set ECG reports, realtime reports, tabular trends reports and graphic trend reports.

27.3.5.1. Setting ECG Reports

To set ECG report, follow this procedure:

- Select 【Main Menu】 quick key→from 【Report】 column to select 【Report Setup】.
- 2. Select **[ECG Report]**.
- 3. Set the desired items. The following table only lists some of the options.

3. Set the desired items. The following table only lists some of the options.			
Menu item	Function	Description	
12-Lead	Set the format of	【12×1】: displays 12-lead waveforms on	
Format	12-Lead waveforms on	one page in one column.	
	a printout.	【6×2】: displays 12-lead waveforms on one	
		page in two columns, with 6 lines in each	
		column.	
		【6×2+1】: displays 12-lead waveforms on	
		one page in two columns, with 6 lines in	
		each column, and one rhythm lead waveform	
		at the bottom.	
		【3×4+1】: displays 12-lead waveforms on	
		one page in 4 columns, with 3 lines in each	
		column, and on rhythm lead waveform at the	
		bottom.	
		【3×4+3】: displays 12-lead waveforms on	
		one page in 4 columns, with 3 lines in each	

		column, and three rhythm lead waveforms at the bottom.
Rhythm Lead1	Select the lead that will	I、II、III、aVR、aVL、aVF、V1、V2、
Rhythm Lead2	be used as Rhythm	V3、V4、V5、V6
Rhythm Lead3	Lead 1, 2, or 3.	
	Note: This setting is only relevant when $[6\times2+1]$, $[3\times4+1]$ or	
	【3×4+3】 is selected for 12-Lead Format.	



CAUTION:

■ When ECG lead is set 3 lead, ECG report cannot be printed.

27.3.5.2. Setting the Realtime Reports

To set realtime reports, follow this procedure:

- Select 【Main Menu 】 quick key→from 【Report 】 column to select 【Report Setup】.
- 2. Select 【Realtime Report】.
- 3. Set the desired items. The following table only lists some of the options.

Menu item	Function	Description
Select	Select the desired waveform	【Current Waveforms】: prints the
Waveform	to printout	realtime report for current waveforms.
		【Selected Waveforms】: prints the
		realtime report for the selected
		waveforms.

27.3.5.3. Setting Tabular Trends Reports

To set tabular trends reports, follow this procedure:

- Select [Main Menu] quick key→from [Report] column to select [Report Setup].
- 2. Select 【Tabular Trends Report】.
- 3. Set the desired items. The following table only lists some of the options.

Menu item	Function	Description
Period	Select the specified period	[Auto]: prints one page of tabular trends
	during which a tabular	report according to the selected resolution.
	trends report will be printed.	
Interval	Select the resolution of the	[NIBP] 、 [EWS] 、 [GCS] : at an
	tabular trends printed on a	interval of acquiring the values of selected
	report.	parameter.
		(Auto): using the (Interval) setting
		of the 【Tabular Trends】 review page.

27.3.5.4. Setting Graphic Trends Reports

- Select [Main Menu] quick key→from [Report] column to select [Report Setup].
- 2. Select [Graphic Trends Report].
- 3. Set the desired options.

27.3.5.5. Setting the Second Mark Switch

To set if a second mark displays on the printed reports, follow this procedure:

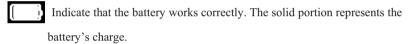
- Select 【Main Menu 】 quick key→from 【System 】 column to select
 【Maintenance】→input maintenance password→Enter.
- 2. Select 【Print】 submenu→【Other】 submenu.
- 3. Switch on or off [Second Mark (Print)] switch.

Chapter 28 Battery

28.1. Introduction

The monitor can be fitted with rechargeable battery to ensure the normal use of the monitor in case of intra-hospital patient transfer or whenever the power supply is interrupted. When the monitor is switched on with AC power, the battery can be charged regardless of whether the monitor is switched on or not. Since we do not provide external charging equipment, the battery can only be charged in the monitor. In case of sudden power failure, the system will automatically use battery to supply power to the monitor, thus not causing interruption of monitoring work.

On-screen battery symbols indicate the battery status as follows:



Indicate that the battery has low charge level and needs to be charged. In this case, the monitor sends out an alarm message.

Indicate that the battery is almost depleted and needs to be charged immediately. Otherwise, the monitor will be automatically shut down.

Indicate that no battery is installed.

The monitor is charging.

The power supply of battery can only function for a certain period. Excessively low voltage of battery will trigger a high priority technical alarm **【Battery Low】**. At this moment, the monitor shall immediately connect with alternating current power supply to charge the battery.

In case of long-term monitoring, a backup battery shall be installed and used after the AC power is plugged in. The AC power plug must be plugged into the special interface of the hospital.

28.2. Installing a Battery

The battery of this monitor must be installed and replaced by maintenance personnel trained and authorized by our company.

28.3. Battery Guidelines

The service life of the battery depends on the frequency and time of use. If lithium batteries are properly maintained and stored, their service life is about 3 years. If batteries are used improperly, their life may be shorter. We recommend replacing lithium batteries every 3 years.

In order to ensure the maximum capacity of the battery, please pay attention to the following instructions:

- ◆ The battery performance must be checked every two months. Before the monitor is repaired or when you suspect that the battery is the source of the fault, battery performance inspection is also required.
- When the battery is used or stored for three months or when the running time of the battery is significantly shortened, the battery performance is optimized once.
- If the monitor is not used for a long time, please optimize the battery performance every three months. Because not taking out the battery will shorten the battery life.
- ◆ If the lithium battery is put on hold when its charge is 50% of its full charge, the storage life of the lithium battery is about 6 months. After 6 months, the lithium battery must be used up before being charged to full capacity. The monitor is powered by the lithium battery, and the battery is taken out of the monitor and then put on hold when the battery is 50% of the full charge.



WARNING:

- Keep the battery out of the reach of children.
- Use only batteries specified in the manufacturer.
- If the battery shows signs of damage or signs of leakage, replace it immediately.

 Do not use a faulty battery in the monitor.

28.4. Battery Maintenance

28.4.1. Optimizing Battery Performance

A battery should be optimized before it is used for the first time. A battery optimizing cycle is one uninterrupted charge of the battery, followed by an uninterrupted battery discharge and charge. Batteries should be optimized regularly to maintain their lifetime.



A CAUTION:

Over time and with the use of batteries, the actual storage capacity of batteries will decrease. For old batteries, the full capacity icon does not mean that the battery storage capacity can still meet the manufacturer's specifications, nor does it mean that the battery power supply time can still meet the manufacturer's specifications. During optimization, if the battery power supply time is obviously shortened, please replace the battery.

To optimize a battery, follow this procedure:

- 1. Disconnect the monitor from the patient and stop all monitoring and measuring procedures.
- 2. Connect the monitor to the AC power supply and charge the battery continuously until the battery is full.
- 3. Remove the AC mains and allow the monitor to run from the battery until it shuts off.
- 4. Reconnect the monitor to AC power and recharge the battery.
- 5. The optimizing of the battery is over.

28.4.2. Checking Battery Performance

The performance of a battery may deteriorate over time. To check the performance of a battery, follow this procedure:

1. Disconnect the monitor from the patient and stop all monitoring and measuring procedures.

- 2. Connect the monitor to AC power and charge the battery continuously until the battery is full.
- Disconnect AC mains and allow the monitor to run on the battery until it shuts off.
- 4. The operating time of a battery reflects its performance directly. If the power supply time of the battery is obviously lower than the time stated in the specification, please consider replacing the battery or contact maintenance personnel.



CAUTION:

■ If the power supply time is too short after the battery is fully charged, the battery may have been damaged or malfunctioned. The power supply time of the battery depends on the equipment configuration and operation. For example, frequent NIBP measurement will also shorten the power supply time of the battery.

28.5. Battery Recycling

When a battery has visual signs of damage, or no longer holds a charge, it should be replaced. Removed the old battery from the monitor and recycle it properly. To dispose of the batteries, follow local laws for proper disposal.



WARNING:

■ Do not disassemble batteries, or dispose of them in fire, or cause them to short circuit. They may ignite, explode, or leak, causing personal injury.

Chapter 29 Maintenance and Cleaning

Use only the disinfectants, cleaners and methods listed in this section to clean or disinfect the monitor, plug-in modules, auxiliary module slots and some accessories. We do not provide any guarantee for damages or accidents caused by the use of other materials or methods.

Our company is not responsible for the effectiveness of the listed chemicals or methods as a means of controlling infection. Please consult the hospital's Infection Control Officer or Epidemiologist.

29.1. Introduction

Keep your monitor, plug-in modules, auxiliary module slots and accessories free of the dust and dirt. To avoid damage to the equipment, follow these rules:

- Please dilute the detergent and disinfectant according to the manufacturer's instructions, or use as low a concentration as possible.
- Do not immerse the monitor in liquid.
- Do not pour liquid on the monitor or accessories.
- ◆ Do not allow liquid to enter the cabinet.
- Abrasive materials (such as steel wool or silver polishing agent) and any strong solvent (such as acetone or detergent containing acetone) as well as liquids with strong conductivity (such as physiological saline) shall not be used.
- Please do not clean or disinfect the equipment when it is running or when it is exposed to direct sunlight.
- ◆ Ensure that all parts of the equipment are completely dry after cleaning and disinfection.



WARNING:

■ Disconnect the power cord from the socket before cleaning the monitor.



CAUTION:

- If you accidentally pour liquid on the monitor, plug-in modules, auxiliary module slots or accessories, please contact the maintenance personnel or our company immediately. Please do not use the equipment until it has been detected and confirmed that it can continue to be used.
- To clean or disinfect reusable accessories, please refer to the instructions provided with the accessories.

29.2. Cleaning of the monitor and other mounting accessories

The monitors, plug-in modules and auxiliary module slots should be cleaned regularly. If there is heavy pollution or lots of dust and sand in your place, the frequency of cleaning should be increased. Before cleaning the monitor, consult the hospital's regulations for cleaning the monitor.

Use a soft cloth that cannot bear balls, wet and clean it with an appropriate amount of water or alcohol-based detergent (such as 70% ethanol), and avoid interfaces and metal parts of equipment. Do not use strong solvents such as acetone or tichlorothylene. Be careful when cleaning the monitor's screen, which is more sensitive than the case. After cleaning, wipe the cleaner off the surface of the mainframe and other mounting accessories with a dry cloth, and place it in a ventilated and cool environment to dry.



CAUTION:

■ Interfaces and metal parts may be corroded after contacting with detergent.

29.3. Disinfecting the monitor and other mounting accessories

You can disinfect the monitors, plug-in modules and auxiliary module slots according to the hospital's disinfection procedures. The equipments should be cleaned before disinfection. The following table lists the recommended disinfectants:

Name	Туре	Manufacturer
Isopropyl alcohol, 70%	Liquid	-
Sodium hypochlorite, 0.5%	Liquid	-
Alcohol, 70%	Liquid	-
Hydrogen peroxide, 3%	Liquid	-

29.4. Cleaning and Sterilizing of Accessories

For cleaning, disinfection and sterilization methods of reusable accessories such as sensors, cables and lead wires, please refer to the instructions of relevant accessories. Please refer to this section if the attachment does not include instructions.

29.4.1. Safety information



CAUTION:

- Do not immerse accessories in water or disinfectant.
- Do not wet the pins of the accessories.
- Frequent disinfection of accessories can cause damage to them. It is suggested that according to hospital regulations, accessories should be disinfected only when necessary.
- When cleaning and disinfecting NIBP airpipe, liquid should be prevented from entering the airpipe.
- Use only the detergents and disinfectants specified in this manual.

29.4.2. Cleaning of the accessories

Use a soft cloth that cannot bear balls, wet and clean the accessories with an appropriate amount of water or alcohol-based detergent (such as 70% ethanol). After cleaning, place the accessories in a cool and ventilated environment to dry.

29.4.3. Disinfection of the accessories

You can disinfect the accessories of the monitor according to the disinfection procedures of the hospital. Recommended disinfectants include:

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Name	Туре	Manufacturer
Isopropyl alcohol, 70%	Liquid	-
Sodium hypochlorite, 10%	Liquid	-
Alcohol, 70%	Liquid	-
Hydrogen peroxide, 3%	Liquid	-
Glutaraldehyde solution, 2%	Liquid	-

29.5. Sterilization

Sterilization of this monitor, related products or accessories is not allowed unless otherwise stated in the accompanying instructions.

Chapter 30 Maintenance



WARNING:

- Hospitals or medical institutions that use monitors should establish perfect maintenance plans, otherwise may cause monitor failure and unpredictable consequences, and may endanger personal safety.
- The safety checks or maintenance involving any disassembly of the equipment should be performed by professional servicing personnel. Otherwise, undue equipment failure and possible health hazards could result.
- If necessary, please contact the manufacturer for product circuit diagrams, parts lists, calibration instructions or other equipment maintenance related information.
- If there is a problem with the monitor, please contact the maintenance personnel or us.

30.1. Inspection

Before use, after continuous use for 6-12 months, maintenance or upgrade, qualified maintenance personnel should conduct a comprehensive inspection to ensure the normal operation and work of the monitor.

Items to be inspected shall include:

- ◆ The environment and power supply meet the requirements.
- There is no mechanical damage to the monitor and accessories.
- ◆ The power cord has no abrasion and good insulation performance.
- Use the specified accessories.
- The alarm system functions normally.
- ◆ The recorder works normally and the recording paper meets the specified requirements.
- ◆ The performance of the battery.
- Various monitoring functions are in good working condition.
- Grounding impedance and leakage current meet the requirements.

If any damage or abnormal phenomenon is found, please do not use the monitor and immediately contact the medical engineer of the hospital or the maintenance personnel of the company.

30.2. Maintenance Schedule

The following tasks, except visual inspection, startup detection, touch screen calibration, battery inspection and recorder inspection, can only be completed by professional maintenance personnel. Please contact the maintenance personnel in time when the following maintenance is required. Before testing or maintenance, the equipment must be cleaned and disinfected.

Check / Maintenance item		Rec	Recommended frequency	
Preventative Maintenance Tests				
Visual inspe	ection	Wh	en first installed or reinstalled.	
NIBP test	Pressure check	1.	If you suspects that the measurement is	
	Leakage test		incorrect.	
CO ₂ and	Leakage test	2.	Following any repairs or replacement of	
AG test	Performance test		relevant module.	
	Module Calibration	3.	At least once a year.	
DM check	Module Check	1.	The DM module used for half a year to one	
			year.	
		2.	Clinicians doubt the accuracy of readings.	
Performano	ce Tests			
ECG test	Performance test			
	Module Calibration	1.	When you suspect that the measured value	
RESP Perfo	rmance test		is inaccurate.	
SpO ₂ test		2.	After the relevant modules are repaired or	
NIBP test	Pressure check		replaced.	
	Leakage test	3.	At least once every two years. NIBP, CO ₂	
TEMP test	TEMP test		and AG modules shall be provided at least	
IBP test	Performance test		once a year.	
IDI test	Pressure zero			

CO ₂ and	Leakage test			
AG test	Performance test			
	Module Calibration			
Nurse call f	unction test			
Analog outp	out performance test	When you suspect that the function is not normal.		
Defibrillation	on synchronization test			
Electrical S	Safety Tests			
Select test	items based on IEC	1. After repairing or replacing the power		
60601-1.		module.		
		2. Or after the monitor falls.		
		3. At least once every two years or as required.		
Other Test	s			
Power-on te	est	1. First installation, or after each reinstallation.		
		2. After each repair or replacement of mair		
		engine components.		
Print test		1. During the first installation.		
		2. After repairing or replacing the printer.		
Recorder check		After repairing or replacing the recorder.		
	Eunationality tost	1. During the first installation.		
Battery	Functionality test	2. After replacing the battery		
check	Performance test	Every two months or when the running time of		
	remainance test	the battery is significantly shortened.		

30.3. Checking the Version Information

You may be asked for version information of monitor and module during monitor maintenance.

Select【Main Menu]quick key→from【System]column to select【Version 】 to check the system software version informatiom.

You can also view more version information by following this procedure:

Select Main Menu quick key→from System column to select Maintenance
→input maintenance password→Enter.

2. Select **[Module Version]**. You can view module software and hardware version, and firmware version.

30.4. Disposing of the Monitor

After the equipment reaches its service life, please dispose of the monitor and its accessories according to local regulations.



WARNING:

■ For the disposal of parts and accessories, if there is no corresponding regulation, local regulations on disposal of hospital waste can be followed.

Chapter 31 Accessories

All accessories listed in this chapter meet the requirements of IEC 60601-1-2 when used with monitors. The accessory materials in contact with the patient passed the biocompatibility test and proved to meet the requirements of IEC 60601-1. For details of accessories, please refer to the relevant accessory instructions.



WARNING:

- Use only the accessories specified in this chapter. Use of other accessories may damage the monitor or fail to meet the specifications claimed in this manual.
- The accessories listed in this chapter must be used together with the monitoring equipment of our company. The user has the responsibility to read the operating instructions of the equipment (including accessories) or contact us for consultation to confirm the matching between the accessories and the equipment. Otherwise, it may cause injury to the patient.
- Disposable accessories can only be used once. Repeated use may cause performance degradation or cross infection.
- Do not open the disposable or sterilized accessory package too early, so as not to cause the accessory to fail or become contaminated.



CAUTION:

- If the use or storage environment of accessories exceeds the specified temperature or humidity range, the performance of accessories may not meet the claimed specifications. If the performance of accessories is degraded due to aging or environmental conditions, please contact customer service personnel.
- If there are signs of damage to the package of the accessory or the accessory itself, please do not use the accessory.
- Do not use the accessory if it expires.
- Disposable accessories must be handled in accordance with local regulations or hospital systems.



NOTE:

- For accessories with safe service life, see the package of accessories for service life.
- Please refer to the package of accessories for sterilization accessories. If the package of the accessories of the sterilization package is damaged, please do not use it.

31.1. Recommended Accessories

ECG cable

Accessories	Specification	Model / PN
	3-lead, IEC, Snap (12PIN)	15-031-0013
	3-lead, AHA, Snap (12PIN)	15-031-0014
	5-lead, IEC, Snap (12PIN)	15-031-0002
ECG cable	5-lead, AHA, Snap (12PIN)	15-031-0004
Legeable	6-lead, IEC, Snap (12PIN)	15-031-0051
	6-lead, AHA, Snap (12PIN)	15-031-0050
	12-lead, IEC, Snap (12PIN)	15-031-0001
	12-lead, AHA, Snap (12PIN)	15-031-0003

\triangleright SpO₂

BLT SpO₂

Accessories	Specification	Model / PN
	Reusable, adult finger	SRA-A11/15-100-0320
	Reusable, adult finger	SRA-A12/15-100-0321
	Reusable, pediatric finger	SRA-P11/15-100-0322
SpO ₂ sensor	Reusable, pediatric finger	SRA-P12/15-100-0323
	Reusable, neonatal	SRA-N13/15-100-0324
	Reusable, Y-type clip	SRA-N15/15-100-0353
	Disposable, adult/neonatal	SDA-N14/15-100-0326

Accessories	Specification	Model / PN
	Reusable, integrated, adult finger	SRA-A21/15-100-0358
	Reusable, integrated, adult finger	SRA-A22/15-100-0359
SpO ₂ Extension cable	Reusable	15-100-0357

The emission wavelength of the pulse oximeter probe is 600-1000nm, and the maximum optical output power is less than 18mW. Information on wavelength range and maximum optical output power is particularly useful to clinicians, for example, for photodynamic therapy.

Masimo SpO₂ (S5)

Accessories	Specification	Model / PN
	Disposable, Neonatal foot (2514)	M-LNCS Neo
	Disposable,Infant toe (2516)	M-LNCS Inf
	Reusable, Adult finger (2501)	M-LNCS DCI
	Reusable,Pediatric finger (2502)	M-LNCS DCIP
	Reusable,Infant foot/hand (2505)	M-LNCS YI
	Disposable, Neonatal (4003)	RD SET Neo
	Reusable, Adult finger (4050)	RD SET DCI
SpO ₂ sensor	Reusable,Pediatric finger (4051)	RD SET DCIP
	Reusable,Infant foot/hand (4054)	RD SET YI
	Disposable, Adult (4000)	RD SET Adt
	Disposable,Pediatric(4001)	RD SET Pdt
	Reusable, Adult finger (2696)	Rainbow DCI
	Reusable,Pediatric finger (2697)	Rainbow DCIP
	Reusable,Multi-site (3792)	Rainbow R2-25a,R2-25r,
	Reusable, Wulti-Site (3/92)	R2-20a,R2-20r
SpO ₂ Extension	RC-12 20pin, MLNC	Rainbow RC-12 / 2404
cable	MD20-12, RD	Rainbow SET
	111020 127 ND	MD20-12/4073

Nellcor SpO₂ (S2)

Accessories	Specification	Model / PN	
	Adult finger,reusable	DS-100A/15-100-0017	
	Neonatal foot,reusable	D-YS/15-100-0018	
	Adult finger (patient	MAX-A	
	size>30kg),disposable	WAX-A	
SpO ₂ sensor	Pediatric foot/hand (patient size	MAX-P	
Spo ₂ sensor	10-50kg),disposable		
	Infant foot/hand (patient size	MAX-I	
	3-20kg),disposable	1711 121	
	Adult finger or neonatal foot/hand	MAX-N	
	(patient size >40 kg or <3 kg),disposable		
SpO_2			
Extension	Nellcor SpO ₂ Extension cable	DOC-10/15-100-0144	
cable			

> TEMP

Accessories	Specification	Model / PN
TEMP Probe	Reusable, Surface	15-031-0005
	Reusable, Coelom	15-031-0012

> NIBP

BLT NIBP

Accessories	Specification	Model / PN
	Disposable, neonatal, 3-5.5cm	M5541-1#/15-100-0104
	Disposable, neonatal, 4-8cm	M5541-2#/15-100-0105
NIBP cuff	Disposable, neonatal, 6-11cm	M5541-3#/15-100-0106
THE CUIT	Disposable, neonatal, 7-13cm	M5541-4#/15-100-0107
	Reusable, neonatal, 6-11cm	M5121/15-100-0122
	Reusable, pediatric, 18-26cm	M5123/15-100-0121

Accessories	Specification	Model / PN
	Reusable, adult, 25-35cm	M5124/15-100-0118
	Reusable, large adult, 33-47cm	M5125/15-100-0120
	Reusable, adult thigh, 44-53cm	M5126/15-100-0142
NIBP air pipe	Reusable	15-031-0008

Suntech NIBP (N2)

Accessories	Specification	Model / PN
	Disposable, neonatal, 3-6cm	98-0400-99
	Disposable, neonatal, 4-8cm	98-0400-96
	Disposable, neonatal, 6-11cm	98-0400-97
	Disposable, neonatal, 7-13cm	98-0400-98
	Disposable, neonatal, 8-15cm	98-0400-90
NIBP cuff	Reusable, pediatric, 12-19cm	98-0600-E1
	Reusable, small adult, 17-25cm	98-0600-E3
	Reusable, adult, 23-33cm	98-0600-E5
	Reusable, large adult, 31-40cm	98-0600-E7
	Reusable, adult thigh, 38-50cm	98-0600-09
	Reusable, adult, 25-35cm	M5124/15-100-0118
NIBP air pipe	Reusable	15-031-0008

► CO₂

BLT Mainstream CO₂

Accessories	Model / PN
CO ₂ sensor	15-100-0199
CO ₂ sensor	16-100-0122

Accessories	Model / PN
Airway adapter (adult)	15-100-0212
Airway adapter (neonate)	15-100-0213
Airway adapter	16-100-0127

Masimo Mainstream CO₂

Specification	Model / PN
CO ₂ sensor	16-100-0017
Airway adapter (adult/pediatric)	16-100-0068
Airway adapter (neonate)	16-100-0067

BLT Sidestream/Microflow CO₂

Accessories	Model / PN
CO ₂ sensor	16-100-0121
CO ₂ water trap	15-100-0229
CO ₂ filter	15-100-0354
CO ₂ dehumidifying tube	16-100-0124
CO ₂ tube	15-100-0035
CO ₂ sampling tube	15-100-0187
CO ₂ L-type 3-way stopcock	15-100-0074
CO ₂ L-type 3-way stopcock	16-100-0126

Masimo Microflow CO₂

Accessories	Model / PN
HH, Airway Adapter Set (adult/pediatric)	3827
HH, Airway Adapter Set (adult/pediatric, 3m)	3828
HH, Airway Adapter Set (neonatal)	3829
LH, Nasal/Oral Cannula (adult)	3822
LH, Nasal/Oral Cannula (pediatric)	3823
HH, Nasal Cannula (adult)	3830
HH, Nasal Cannula (pediatric)	3831

Accessories	Model / PN
HH, Nasal Cannula (neonatal)	3832
HH, Nasal/Oral Cannula (adult)	3835
HH, Nasal/Oral Cannula (pediatric)	3836
HH, Nasal Cannula with O ₂ delivery (adult)	3833
HH, Nasal Cannula with O ₂ delivery (pediatric)	3834
HH, Nasal Cannula/Oral with O ₂ delivery (adult)	3837
HH, Nasal Cannula/Oral with O ₂ delivery (pediatric)	3838

CO₂ extension cable

Accessories	Model / PN
Reusable, CO ₂ extension cable	15-031-0010
Reusable, CO ₂ extension cable	15-031-0011

> DM module

Accessories	Model / PN
DM module	BLT-IA2/16-100-0113

> IBP

Accessories	Model / PN
IBP sensor	PT-1/15-100-0053
IBP cable	15-100-0029
IBP extension cable (4PIN to 6PIN)	15-031-0023

> EEG

Accessories	Model / PN
EEG Cable	15-100-0204

> BIS

Accessories	Model / PN
BIS Patient Cable	186-0195-SF
BIS Electrodes	186-0106

Accessories	Model / PN
BIS ICU Electrode	186-0160
BIS Pediatric Electrode	186-0200

> ICG

Accessories	Model / PN
ICG cable	N1301-4/15-100-0180
ICG electrode	N1201-5/15-100-0181

> AG

Mainstream AG

Accessories	Model / PN
AG module extension cable	15-031-0011
AG sensor	16-100-0019
Airway adapter (adult/pediatric)	16-100-0068
Airway adapter (neonatal)	16-100-0067

Sidestream AG

Accessories	Model / PN
HH, Airway Adapter Set (adult/pediatric)	3827
HH, Airway Adapter Set (adult/pediatric, 3m)	3828
HH, Airway Adapter Set (neonatal)	3829
LH, Nasal/Oral Cannula (adult)	3822
LH, Nasal/Oral Cannula (pediatric)	3823
HH, Nasal Cannula (adult)	3830
HH, Nasal Cannula (pediatric)	3831
HH, Nasal Cannula (neonatal)	3832
HH, Nasal/Oral Cannula (adult)	3835
HH, Nasal/Oral Cannula (pediatric)	3836
HH, Nasal Cannula with O ₂ delivery (adult)	3833
HH, Nasal Cannula with O ₂ delivery (pediatric)	3834
HH, Nasal Cannula/Oral with O ₂ delivery (adult)	3837

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Accessories	Model / PN
HH, Nasal Cannula/Oral with O ₂ delivery (pediatric)	3838

> RM

Accessories	Model / PN
Disposable, RM sensor (Adult/Pediatric)	040-001949-00/15-100-0327
Disposable, RM sensor (Neonate)	040-001950-00/15-100-0328

> NMT

Accessories	Model / PN
NMT Smart Data Cable(RS232)	XT-45100C-NMS/15-100-0415
NMT Sub-machine	NMS450X/15-100-0416
NMT NMBA Monitoring Cable	XT-45025/15-100-0417
NMT Nerve Mapping/Locating Cable	XT-41014/15-100-0418

➤ rSO₂

Masimo rSO₂ (R1)

Accessories	Model / PN
O3 cable	15-100-0404
Disposable, O3 Sensor rSO ₂ (Adult, Adhesive, ≥40Kg)	3756/15-100-0405
Disposable, O3 Sensor rSO ₂ (Pediatric, Adhesive, 5-<40Kg)	4235/15-100-0406

Nonin rSO₂ (R2)

Accessories	Model / PN
Nonin System Hub cable	X-100H/15-100-0407
Nonin blue Signal Processor	3500SP-1/15-100-0408
Nonin yellow Signal Processor	3500SP-2/15-100-0409
Nonin Intermediate Cable of 8204CA sensor	INT-100/15-100-0414
Disposable, Nonin adult/pediatric regional oximetry sensor, ≥40 kg	8204CA/15-100-0410

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Accessories	Model / PN
Disposable, Nonin adult/pediatric regional oximetry sensor, ≥40 kg	8004CA/15-100-0411
Disposable, Nonin neonatal/pediatric regional oximetry sensor, ≤40 kg	8004CB/15-100-0412
Disposable, Nonin neonatal/pediatric regional oximetry sensor, ≤40 kg, non-adhesive	8004CB-NA/15-100-0413

Other accessories

Accessories	Model / PN
Auxiliary Module Slots	15-091-0003
Auxiliary Module Slots cable	11-091-0092
Biolink cable (RS232)	11-100-0059

Appendix A Product Specifications

A.1 Safety Specifications

According to the MDD 93/42/EEC, the monitor is Type IIb equipment. Classified according to the IEC60601-1 is as follows:

Parts	Classifica tion of protectio n against electric shock	Degree of protectio n against electric shock	Degree of protectio n against ingress of liquid	Degree of protection against hazards of explosion	Recommen ded disinfection and sterilization methods	Mode of operation
Mainframe Fixed parameter (ECG, TEMP, RESP, NIBP, SpO ₂) IBP ICG EEG CO ₂ BIS AG RM DM rSO ₂	NA	Type CF defibrillati on proof Type BF defibrillati on proof	IP21	Not suitable	See Chapter 29 Maintenanc e and Cleaning of this manual for details.	Continuous
NMT		illation proof				

Note:

I: Class I, internally and externally powered equipment.

When you doubt about the protecting earth integrality or protecting earth lead of the equipment, you'd better change the equipment to internally powered equipment.

NA: Not applicable

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

A.2 Environmental Specifications

Components	Item	Operating condition	Storage condition
	Temperature (℃)	0~40	-20~+60
Mainframe and other modules	Relative humidity (non-condensing) (%)	15~95	10~95
	Barometric (kPa)	54.0~107.4	16.0~107.4
	Temperature (℃)	10~40	-40~+70
RM	Relative humidity (non-condensing) (%)	15~95	15~95
	Barometric (kPa)	70.0~107.4	70.0~107.4
	Temperature (°C)	10~40	0~+50
ICG	Relative humidity (non-condensing) (%)	15~95	15~95
	Barometric (kPa)	56.0~107.4	13.3~133.3
	Temperature (°C)	0~40	-40~+70
Microflow CO ₂ (C5)	Relative humidity (non-condensing) (%)	<4 kPa water (95%RH, 30°C)	5~100
	Barometric (kPa)	57.0~107.4	20~120
	Temperature (℃)	0~40	-40~+75
Mainstream CO ₂ (C2)	Relative humidity (non-condensing) (%)	10~95	5~100
	Barometric (kPa)	57.0~107.4	50~120
Mainstream CO2	Temperature (℃)	5~40	-40~+70
and Sidestream CO2 (C1, C11	Relative humidity (non-condensing) (%)	10~90	<90

and C6)	Barometric (kPa)	55.0~107.4	55.0~107.4
Mainstream AG	Temperature (°C)	10~40	-20~+75
	Relative humidity (non-condensing) (%)	10~95	5~100
	Barometric (kPa)	57.0~107.4	50~120
	Temperature (°C)	5~40	-40~+70
Sidestream AG	Relative humidity (non-condensing) (%)	<4 kPa water (95%RH, 30°C)	5~100
	Barometric (kPa)	57.0~107.4	20~120
Masimo rSO ₂	Temperature (℃)	0~40	-40~+70
	Relative humidity (non-condensing) (%)	10~95	10~95
	Barometric (kPa)	57~106	50~106
Nonin rSO ₂	Temperature (°C)	0~40	-40~+70
	Relative humidity (non-condensing) (%)	15~93	<93
	Barometric (kPa)	/ (altitude 0~4000m)	/



CAUTION:

The equipment must be used under the specified environmental specifications, otherwise it will not meet the technical specifications claimed in this manual and may lead to unexpected consequences such as equipment damage. If the performance of the equipment changes due to aging or environmental conditions, please contact the maintenance personnel.

A.3 Physical Specifications

Parts	Model	Weight	Size (L×H×D)	Remark
		P12: <5kg	P12: 323×281×157	
Main unit	P12/P15/	P15: <6kg	P15: 396 × 310 ×198	
Main unit	P18/P22	P18: <7kg	P18: 462 ×334×199	
		P22:<8kg	P22: 530×375 ×199	
P1	P1	<0.9kg	146 ×99 ×76	
MPS-P module	MPS-P	<0.6kg	136 ×106×81	
SpO ₂ module	S2	<0.3kg	136.6×102×40	Nellcor SpO ₂
SpO ₂ module	S5	<0.3kg	136.6×102×40	Masimo SpO ₂
Mainstream CO ₂ module	C2	<0.3kg	136.6×102×40	Masimo IRMA
Mainstream CO ₂ module	C6	<0.3kg	136.6×102×40	BLT CO ₂
Mainstream CO ₂ module	C8	<0.3kg	136.6×102×40	BLT CO ₂
Sidestream CO ₂ module	C1	<0.4kg	136.6×102×40	BLT CO ₂
Microflow CO ₂ module	C5	<0.3kg	136.6×102×40	Masimo ISA Capno
Microflow CO ₂ module	С9	<0.3kg	136.6×102×40	BLT CO ₂
Microflow CO ₂ module	C10	<0.3kg	136.6×102×40	BLT CO ₂
Microflow CO ₂ module	C11	<0.4kg	136.6×102×40	BLT CO ₂
NIBP module	N2	<0.3kg	136.6×102×40	SunTech NIBP
ICG module	ICG	<0.3kg	136.6×102×40	
BIS modulle	BIS	<0.3kg	136.6×102×40	
IBP module	IBP2	<0.3kg	136.6×102×40	
TEMP module	T2	<0.3kg	136.6×102×40	
EEG module	EEG	<0.3kg	136.6×102×40	
DM module	DM	<0.2kg	136.6×102×40	
AG module	AG1	<0.3kg	136.6×102×40	IRMA AX+

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AG module	AG4	<0.3kg	136.6×102×40	ISA AX+
AG module	AG5	<0.6kg	136.6×102×80.5	ISA OR+
RM module	RM	<0.3kg	136.6×102×40	
NMT module	NMT	<0.3kg	136.6×102×40	
rSO ₂ module	R1	<0.3kg	136.6×102×40	Masimo rSO ₂
rSO ₂ module	R2	<0.3kg	136.6×102×40	Nonin rSO ₂
BIOLINK module	BIOLIN K	<0.3kg	136.6×102×40	
Auxiliary Module Slots		<3kg	436×130×134	

A.4 Power Specifications

A.4.1 External power supply

≻ P12

Input voltage	AC (100-240) V (±10%)
Frequency	50Hz/60Hz
Input power	1.5A - 0.7A
Standard requirements	According to IEC 60601-1 and IEC 60601-1-2

> P15/P18/P22

7 1 13/1 10/1 22	
Input voltage	AC (100-240) V (±10%)
Frequency	50Hz/60Hz
Input power	2.0A - 0.9A
Standard requirements	According to IEC 60601-1 and IEC 60601-1-2

A.4.2 Battery

Battery (optional configuration)		
Type	Rechargeable lithium ion battery, 11.1 VDC, 5000mAh	
Operating time	In a new and fully charged battery at (25°C) ambient temperature,	
	typical configuration (screen brightness is level 1, connected to	

	SpO ₂ sensor, ECG cable, and NIBP works in an automatic
	measurement mode with a time interval of 15 min):
	P12 monitor: ≥4h
	P15/P18/P22 monitor: ≥3h
Charge time	The monitor is charged to 90% for less than 3 hours and 100% for
	less than 4 hours when it is turned off.
	When the monitor is turned on, it is charged to 90% for less than 5
	hours and 100% for less than 6 hours.
Turn off delay	5 min-15min (after the low battery alarm first occurs)

A.5 Hardware Specifications

A.5.1 Display

Host display	
Туре	Color TFT LCD
Size (diagonal)	P12: 12.1 inch
	P15: 15.6 inch
	P18: 18.5 inch
	P22: 22 inch
Resolution	P12: 1280×800 pixels
	P15/P18/P22: 1920×1080 pixels
External Display	
P12/P15/P18/P22	DVI Display, resolution above 1920×1080 pixels

A.5.2 Recorder

Type	BTR50S thermal dot array
Paper width	50 mm±1mm
Recording speed	12.5 mm/s, 25 mm/s, 50 mm/s
Recording waveform	Maximum 3 tracks

A.5.3 Mainframe LED

Alarm lamp	Cyan, yellow and red
Power indicating lamp	1 (Green/Orange)
	When powered with AC, it lights green while turn on and off
	the monitor.
	When powered with battery, the orange light is on when the
	battery is turned on, and no light is on when the battery is
	turned off.
Battery charging	1 (yellow), When charging, it is always on, and when it is fully
indicating lamp	charged, the light goes out.

A.5.4 Audio indicating

Speaker	Give alarm tone (45-85dB), QRS tones;
	Support PITCH TONE and multi-level tone modulation;
	Alarm tones meet the requirements of IEC 60601-1-8.

A.5.5 Input device

Keys	
Physical keys	1 power switch key
Touch screen	Support
Others	
Mouse input	Support (optional)
Keyboard input	Support (optional)
Barcode scanner	Support (optional)
Voice assistant	Support (optional)

A.5.6 Connectors

Power	1 AC power inlet with cable retainer
Wired network (standard	P12/P15: 1
RJ45 interfaces)	P18/P22: 2 (1 of them is reserved interface)

USB (standard USB 2.0	P12/P15: 4
sockets)	P18/P22: 8 (4 of them is reserved interface)
DVI connector	P12/P15: 1
	P18/P22: 2 (1 of them is reserved interface)
Equipotential grounding	1
point	
Nurse call interface	1
Auxiliary Modules Slots	1
connector	

A.5.7 Signal Output

Auxiliary output interface (optional)	
Standard	Meet the requirements of IEC 60601-1 for short-circuit
	protection and leakage current.
Output impedance	Rated 50Ω
ECG analog signals ou	tput
Output signal range	-10V~+10V
Maximum	25 ms
transmission delay	23 1115
Sensitivity	1V/mV±5%
PACE rejection /	Has PACE rejection function
strengthen	Thas I Med Tejection function
IBP analog signals out	put
Output signal range	-1V~+4V
Maximum	35 ms
transmission delay	33 1115
Sensitivity	1V/100mHg±5%
Nurse call output	
Output voltage range	High level: 3.5~5V, providing a maximum of 10 mA output
	current;

	Low level: < 0.5V, receiving a maximum of 5 mA input		
	current.		
Isolated voltage	1500 VAC		
ū			
Signal type	N.C., N.O., Pulse Output (optional);		
Rise and drop time	≤1ms		
Defibrillator synchron	ization signal output		
Output impedance	50Ω±10%		
Maximum delay	25 ms (from R wave crest to pulse raise)		
Amplitude	High level: 3.5~5V, providing a maximum of 1 mA output		
	current;		
	Low level: <0.5V, receiving a maximum of 5 mA input current.		
Pulse width	100ms±10%		
Rise and drop time	<1ms		
Alarm output	Alarm output		
Indicates the inherent	≤1s		
delay in determining			
the alarm status.			
Alarm delay time	The alarm delay time from the monitor to remote equipment is ≤		
from the monitor to	2s, measured at the monitor signal output connector.		
remote equipment			
Alarm signal sound	Within a distance of one meter, the peak volume range of the		
pressure level range	audible alarm generated by the equipment is $45 dB(A) \sim 85 dB(A)$.		

A.6 Data Storage

Trend data	Long trend: 1800h, minimum resolution is 10 min
	Medium trend: 180h, minimum resolution is 1 min
	Short trend: 6h, minimum resolution is 5 second.
Parameter alarm	At least 3000 parameter alarm events and associated parameter
event	waveform at the moment.
ARR events	3000 ARR events, and the parameter waveform related to the
	time of event occurrence.

NIBP measurement	At least 2400 groups.
result	
Holographic	At least 72 hours. The specific storage time depends on the
waveform	waveforms stored and the number of stored waveform.

A.7 Wireless network

Conforming	IEEE802.11a/b/g/n
standards	
Operating	$2.4 \mathrm{GHz} \sim 2.495~\mathrm{GHz},~5.15 \mathrm{GHz} \sim 5.35 \mathrm{GHz},~5.47 \mathrm{GHz} \sim$
frequency	5.725GHz, 5.725 GHz~5.82GHz
Data security	WPA-PSK、WPA2-PSK
Encryption	AES、TKIP

A.8 Measurement Specifications

The product shall meet the following measurement specifications. If there is no special indication, the definition of the index shall preferentially refer to the special standard of the parameter.

A.8.1 ECG Specifications

A.8.1.1 Standard

Meet standards of IEC 60601-2-27	
Meet standards of IEC 60601-2-25	

A.8.1.2 Performance indicators

Electrogramacony	Cut mode: 300W
Electrosurgery	Coagulate mode: 100W
protection	Recovery time: ≤10s

	In compliance with the requirements in clause 202.6.2.101 of		
	IEC 60601-2-27.		
Lead-off detection	Measuring electrode: <100nA		
current	Driving electrode (RL): <1 uA		
Tall T-wave	15		
rejection capability	1.5mV		
	Under normal circumstances, the 12 most recent RR intervals		
	are averaged to compute the HR.		
IID	If the last 3 consecutive RR intervals are greater than 1200ms		
HR averaging	(i.e., HR is less than 50bpm), the 4 most recent RR intervals are		
method	averaged to compute the HR.		
	The HR value displayed on the monitor screen is updated every		
	second.		
	Meet the requirements of Clause 201.7.9.2.9.101 b) 4) of IEC		
	60601-2-27.		
	The heart rate value displayed after the 20-seconed stabilization		
Response to	period is:		
irregular rhythm	Waveform 3a (Ventricular bigeminy): 80bpm;		
	Waveform 3b (Slow alternating ventricular bigeminy): 60bpm		
	Waveform 3c (Rapid alternating ventricular bigeminy): 120bpm		
	Waveform 3d (Bidirectional systoles): 90bpm		
Response time to	HR change from 80bpm to 120bpm: < 10s.		
heart rate change	HR change from 80bpm to 40bpm: <10s.		
Time to alarm for	<11s		
Tachycardia	~115		
Dago mulgo es estera	Amplitude: ±2mV~±700mV		
Pace pulse markers	Pulse width: 0.1~2.0ms		
Pacemaker pulse	Amplitude: ±2mV~±700mV, Pulse width: 0.1~2.0ms		
rejection capability	Heart rate calculation should not be affected.		
without overshoot.	Treat rate calculation should not be affected.		
Minimum slew rate			
for pacing pulse	$12.5 \text{V/s} \pm 20\%$		

detection.	
Pacing pulse display method in auxiliary output	Suppression
Pacing function switch	Pace markers function can be switched on and off.
Defibrillation output delay	25ms
ECG Analog Output Delay	25ms

A.8.1.3 ECG Measurement

	3-lead: I, II, III		
	5-lead: I, II, III, aVR, aVL, aVF, V-		
Lead type	6-lead: I, II, III, aVR, aVL, aVF,Va, Vb		
	12-lead: I, II, III, aVR, aVL, aVF,V1~V6		
	Auto: identify leads automatically		
Indication of lead-off shall	Every electrode		
be provided	Every electrode		
ECG abnormal work	Every amplification channel shall have an indication of		
indications	abnormal ECG operation (polarization).		
	Diagnostic mode: 0.05~150Hz		
D 1 (141. (24D)	Monitor mode: 0.5~40Hz		
Bandwidth (-3dB)	Operation mode: 1~25Hz		
	ST mode: 0.05~40Hz		
Signal quality display	Expression way: numerical display and waveform color.		
	Breakdown Voltage: 4000V 50Hz/60Hz		
Defibrillation Protection	Anti-defibrillation effect protection: baseline recovery		
	time: 5s (after defibrillation).		
Input signal range	Input signal range -10.0mV~+10.0mV		
Input signal Lange	Electrode offset ±500 mV d.c.		

		potential		
Input impedance		≥5.0MΩ		
System noise		≤30µVpp RTI		
Waveform Display		6.25mm/s, 12.5mm/s, 25mm/s, 50mm/s, error ≤±5%		
sweep speed	Recorder	12.5mm/s, 25mm/s, 50mm/s, error ≤±5%		
Waveform	Display	×0.25, ×0.5, ×1 (10mm/mV), ×2, ×4, error ≤±5%.		
Gain	Бізрійу	Auto		
Guin	Recorder	$\times 0.25, \times 0.5, \times 1 \text{ (10mm/mV)}, \times 2, \times 4, \text{ error } \leq \pm 5\%.$		
CMRR		Diagnostic mode	≥100 dB	
CMIKI		Monitor, Operation mod	le ≥110 dB	
Calibration	voltage	<±5% (×1)		
accuracy				
Input offset current		<0.1uA		
Time constant		Monitoring mode: ≥0.3s		
		Diagnostic mode: ≥3.2s		

A.8.1.4 ECG analysis calculation

Measurement parameter	HR, PVCs, ST, QT, and Arrhythmia analysis		
Multi-lead	Supports synchro	nous analysis of at least 2 leads, one of which	
synchronous	is the key monito	oring lead and the other is the auxiliary lead. It	
analysis function	is on except 3-lead mode.		
Smart Lead Switch	Automatic, Manual; Default manual (3-lead mode is fixed as manual) Automatic mode: the algorithm automatically identifies the current smart leads, and the host automatically switches the key monitoring leads according to the identification of the algorithm.		
HR measurement range and accuracy	Measurement Adult: 10~300 bpm range Pediatric/Neonatal: 10~350bpm Resolution 1 bpm Accuracy ±1% or ±1 bpm, whichever is greater		

	Detection sensitivity	0.20mVp-p	
ST Display	Display 12-lead ST segment values at the same time and support ST graphic display.		
	Measurement range	-2.0mV~+2.0mV	
ST measurement	Resolution	±0.01mV	
range and accuracy		$-0.8 \text{mV} \sim +0.8 \text{mV}$: $\pm 0.02 \text{mV}$ or $\pm 10\%$,	
	Accuracy	whichever is greater	
		Other: Unspecified	
ST Update period	10s		
PVCs measurement	Indicates the number of PVC in the past minute, ranging from 0/min ~150/min.		
Arrhythmia analysis type	27 (see Table 2)		
QT analysis function	Pediatric / neonatal: 15bpm~180bpm		
	Resolution QT, QTc, Δ QTc: 1ms; QT-HR: 1bpm		
	Accuracy	QT: ±30ms	
Sampling rate	1000Hz (The time bias among every channels ≤100us)		
Amplitude quantisation	≤1 uV/LSB		

Table 2 Arrhythmia Events list

No.	Full Name	Prompt information
1	Asystole	Asystole
2	Ventricular Fibrillation/	Vent Fib/Tach
	Ventricular Tachycardia	vent F10/ Facil
3	Ventricular Tachycardia	V-Tach
4	Ventricular Bradycardia	Vent Brady
5	Extreme Tachycardia	Extreme Tachy
6	Extreme Bradycardia	Extreme Brady
7	R on T	R on T
8	Tachycardia	Tachy
9	Bradycardia	Brady
10	Nonsustained Ventricular Tachycardia	Nonsustained V-Tach
11	Ventricular Rhythm	Vent Rhythm
12	Pacer Not Captured	PNC
13	Pacer Not Pacing	PNP
14	Heartbeat Pause	Pause
15	Pauses/min High	Pauses/min High
16	Run PVCs	Run PVCs
17	Couplet	Couplet
18	VentricularBigeminy	Bigeminy
19	VentricularTrigeminy	Trigeminy
20	Frequent PVCs	Frequent PVCs
21	Premature ventricular contraction	PVC
22	Missed Beat	Missed Beat
23	Atrial Fibrillation	A-Fib
24	Atrial Fibrillation End	A-Fib End
25	ECGNoise	ECG Noise
26	Irregular Rhythm	Irregular Rhythm
27	Irregular Rhythm End	Irregular RhythmEnd

A.8.2 RESP Specifications

A.8.2.1 Measurement specification

Measurement parameter	Respiration Rate and respiration waveform		
Source	RA-LA, RA-LL (default)		
Excitation waveform	Excitattion frequency: 6	64 kHz; Error: ≤±10%	
Excitation current	≤0.3mA RMS		
Respiration Apnea	Fixed high priority alarm Adjustable delay time: 10~60s, error ±3s or ±10%, whichever is greater.		
Cardiac interference alert	Fixed high priority alarm		
	Measurement range	0~150 rpm	
RR measurement	Resolution	1 rpm	
range and accuracy	Accuracy	±2 rpm or ±2% of reading, whichever is greater.	
Bandwidth	0.2 Hz ~2.5Hz (-3dB~+	0.4dB)	
Sweep speed	6.25mm/s, 12.5mm/s, 25mm/s, 50mm/s, error is within ±10%.		
Baseline impedance	$200\sim2500\Omega$ (using defibrillator proof cable with resistance of		
range	1kΩ)		
Measuring impedance range	0.3Ω~3Ω		
Gain	×0.25, ×0.5, ×1, ×2, ×4		

A.8.3 NIBP Specifications

A.8.3.1 Standard

Meet standard of IEC80601-2-30

A.8.3.2 Measurement Specification

> BLT NIBP

Measurement parameters	SYS, DIA, MAP,PR		
Mode of operation	Manual, Auto, STAT, Sequence		
Periodic measurement	The monitor automatically starts NIBP measurement at set intervals.		
Sequence mode	At least 5 groups are supported, and each group individually sets the interval and number of periodic measurement.		
STAT mode cycle time	5 min		
Measurement range of cuff pressure	0~300 mmHg		
Initial inflation	Adult: 100-	~280mmHg, default 160mmHg	
pressure	Pediatric: 100~240 mmHg, default 130mmHg		
pressure	Neonatal: 60~140mmHg, default 100mmHg		
Sensor calibration time	One year (recommend)		
Unit	mmHg, kPa		
	Systolic	Adult: 30~270 mmHg (4.0~36.0 kPa)	
		Pediatric: 30~235 mmHg (4.0~31.3 kPa)	
		Neonatal: 30~135 mmHg (4.0~18.0 kPa)	
		Adult: 10~220 mmHg (1.3~29.3 kPa)	
Dynamic pressure	Diastolic	Pediatric: 10~220 mmHg (1.3~29.3 kPa)	
measurement range		Neonatal: 10~110 mmHg (1.3~14.6 kPa)	
		Adult: 20~235 mmHg (2.7~31.3 kPa)	
	Mean	Pediatric: 20~225 mmHg (2.7~30.0 kPa)	
		Neonatal: 20~125 mmHg (2.7~16.6 kPa)	

Dynamic Pressure Measurement Error of Simulator	±8 mmHg (±1.1kPa)	
Static pressure accuracy	±3 mmHg (±0.4kPa)	
Pressure resolution	1 mmHg or 0.1k	rPa
PR measurement range and accuracy	Measurement range	40 ~ 240 bpm
	Accuracy	±3bpm or ±3%, whichever is greater
Maximum	Neonatal: <90s	
measurement time	Adult, Pediatric: <120s	
First overvoltage protection point	Adult: 297±3 mmHg Pediatric: 252±3 mmHg Neonatal: 147±3 mmHg	
Second overvoltage protection point	Adult: 315±10 mmHg Pediatric: 265±10 mmHg Neonatal: 155±10 mmHg	

Note: The accuracy of NIBP cannot be determined by using a simulator, but under many conditions, it is still necessary to use a simulator to test its performance (for example, a simulator is required for quality control in the production process), and the simulator of the model specified by the manufacturer shall be used for this performance test.

Clinical index	
Evaluation method for	Follow ISO 81060-2 standard, in which
clinical accuracy of blood	Adult (including pediatric): auscultation method
pressure	Neonatal: invasive method
	Systolic and diastolic pressures: mean error: ±5mmHg,
Accuracy	standard deviation: ≤8 mmHg
	Mean pressure: not participating in evaluation
Overall measurement time	20~45s (typical value)

> Suntech NIBP (optional)

Way of measurement	Oscillometric.		
		Adult: 40~260 mmHg (5.3~34.7 kPa)	
	Systolic	Pediatric: 40~160 mmHg (5.3~21.3 kPa)	
		Neonatal: 40~130 mmHg (5.3~17.3 kPa)	
D 6		Adult: 20~200 mmHg (2.7~26.7 kPa)	
Range of measurement	Diastolic	Pediatric: 20~120 mmHg (2.7~16.0kPa)	
measurement		Neonatal: 20~100 mmHg (2.7~13.3kPa)	
		Adult: 26~220 mmHg (3.5~29.3 kPa)	
	Mean	Pediatric: 26~133 mmHg (3.5~17.7 kPa)	
		Neonatal: 26~110 mmHg (3.5~14.7kPa)	
Static pressure accuracy	±3 mmHg (±0.4kPa)		
Unit	mmHg, kPa	-	
Pulse rate range	30 ~ 220 bpn	1	
Pulse Rate Accuracy	±2bpm or =	±3%, whichever is greater	
Inflation time for cuff	<75s		
Measurement	Adult: <180s		
protection time	Pediatric: <180s		
processor same	Neonate: <90s		
Initial inflation	Adult: 120~280mmHg, default 160mmHg		
pressure	Pediatric: 80~170mmHg, default 120mmHg		
prossure	Neonate: 60~140mmHg, default 90mmHg		
Overpressure			
Protection	Hardware and software double protections		
Adult	<300 mmHg		
Pediatric	<300 mmHg		

Neonate	<150 mmHg
Clinical accuracy	Systolic and diastolic pressures: mean error: ±5mmHg, standard
	deviation: ≤8 mmHg

A.8.4 SpO₂ Specifications

A.8.4.1 Standard

Meet the standard of ISO 80601-2-61

A.8.4.2 Specification

► BLT SpO₂

Measurement	Measure two channels of SpO $_2$, PR, PI, Δ SpO $_2$ and RR, and			
parameter	show SpO ₂ waveform and respiration waveform.			
Sensitivity	High, Mediun	High, Medium, Low		
	Measurement range		0~100%	
			70%~100%: ≤3% (SpO ₂ probe	
Measurement range	Clinical accur	acy	included in Appendix);	
Measurement range and accuracy			0~69%: unspecified	
and accuracy	Simulator measurement		70%~100%: ±2% (non-motion	
			conditions)	
	CITOI		0~69% unspecified	
SpO ₂ Update period	Normal		≤2s	
Spo ₂ epalite period	Maximum		≤25s	
		High	≤8s	
SpO ₂ Response time	Sensitivity	medium	≤11s	
		Low	≤15s	
	Measurement range		25 bpm ~300 bpm	
PR	Resolution		1 bpm	
	Accuracy		± 3bpm (non-motion conditions)	

	Measurement range	At least 0.05~20.00%
PI	Resolution	0.01%
	Accuracy	$\pm 0.1\%$ or $\pm 10\%$ of reading, whichever is greater
	Measurement range	0 rpm ~90 rpm
RESP (from pleth)	Resolution	1 rpm
	Accuracy	±2 rpm

Sensor model	Number of subjects	Number of Data	Arms
SRA-A11	11 (male 5, female 6)	236 pts	1.67
SRA-A12	11 (male 5, female 6)	236 pts	1.63
SRA-P11	10 (female)	208 pts	2.04
SRA-P12	10 (female)	208 pts	1.84
SRA-N13	10 (female)	208 pts	1.69
SRA-N15	10 (female)	208 pts	1.8
SDA-N14	10 (female)	208 pts	2.04
	11 (male 5, female 6)	236 pts	2.15

Remarks:

- a) *The accuracy of SpO₂ cannot be determined by simulator, but in many conditions, it is still necessary to use a simulator to test its performance (for example, a simulator is required for quality control in the production process), and the simulator of the model specified by the manufacturer shall be used for testing.
- b) Confirmation of measurement accuracy: The accuracy of SpO_2 has been confirmed in human experiments by comparing with the arterial blood oxygen reference value measured by CO blood gas analyzer. The measurement results of arterial oxygen saturation confirm to statistical distribution. Compared with the measurement results of CO blood gas analyzer, only two thirds of the measurement results are expected to be within the specified accuracy.
- c) The accuracy of blood oxygen under low perfusion conditions cannot be determined using a simulator. The definition here only refers to the PI index, and does not mean that SpO_2 can reach the claimed accuracy under the low perfusion condition during clinical use.

➤ Nellcor SpO₂

Measurement range	0% to 100%		
Resolution	1%		
Accuracy	70% to 100%: ±2% (adult/pediatric, non-motion) 70% to 100%: ±3% (neonate, non-motion) 0% to 69%, unspecified		
Alarm range and	Alarm range	0% to 100%, high/low limit can be adjusted continuously.	
error	Alarm error	±1%	
Average time	8s, 16s		
	Measurement range	20 bpm to 300 bpm	
	Accuracy	20 bpm to 250 bpm: ±3 bpm (non-motion conditions) 251 bpm to 300 bpm: unspecified	
PR	Resolution	1 bpm	
	Alarm range	0bpm∼300bpm, high/low limit can be adjusted continuously.	
	Alarm error	±1bpm	
	Measurement range	At least 0.05~20.00%	
PI	Resolution	0.01%	
	Accuracy	$\pm 0.1\%$ or $\pm 10\%$ of reading, whichever is greater	
Update period	Normal	≤2s	
opuate periou	maximum	≤25s	

➤ Masimo SpO₂

Measurement parameter	SpO ₂ , SpMet, PVI, SpHb, SpOC, PI and SpCO		
Measurement range	0% to 100%		
Resolution	1%		
Accuracy	70% to 100%:±2% (adult/pediatric, non-motion conditions) 70% to 100%:±3% (neonate, non-motion conditions) 0% to 69%,unspecified		
Average time	2-4s, 4-6s, 8s, 10s, 12	2s, 14s, 16s	
Alarm range and	Alarm range	0% to 100%, adjusted conti	high/low limit can be nuously.
error	Alarm error	±1%	
	Measurement range	25 bpm to 240 bpm	
	Accuracy	±3 bpm (non-motion conditions)	
PR	Alarm range and error	Alarm range	0bpm to 300bpm, high/low limit can be adjusted continuously.
		Alarm error	±1bpm
	Measurement range	0% to 100%	1
SpCO	Accuracy	0% to 40%: ±3%(non-motion conditions) >40%, unspecified	
	Resolution	1%	
	Measurement range	range 0.0% to 100.0%	
SpMet	Accuracy	0% to 15%:±1%(non-motion conditions) >15%, unspecified	
	Resolution	0.1%	

	Measurement range	At least 0.05% to 20.00%
PI	Resolution	0.01%
	Accuracy	$\pm 0.1\%$ or $\pm 10\%$ of reading, whichever is greater
Measurement range		0.0 g/dl to 25.0 g/dl
SpHb	Accuracy	8 g/dl to 17 g/dl: ±1 g/dl (non-motion conditions) <8 g/dl or >17 g/dl, unspecified
	Resolution	0.1g/dl

A.8.5 TEMP Specifications

A.8.5.1 Standard

Meet the standard of ISO 80601-2-56.

A.8.5.2 Measurement Specification

Parameter	$T1,T2,T_D$		
Probe	YSI400 series probe (2252Ω@25°C, accuracy ±0.1°C)		
Measurement site	Surface and coelom		
Massuroment rongs and	Measurement range	0.0°C ~50.0°C (32.0°F ~122.0°F)	
Measurement range and accuracy	Resolution	0.1°C or 0.1°F	
·	Accuracy of circuit	± 0.1 °C (± 0.2 °F) (without sensor)	
Alama wan sa	0.0°C to 50.0°C (32.0°F \sim 122.0°F) , high/low limit c be adjusted continuously.		
Alarm range			
Alarm error	±0.1℃ or ±0.1°F		

Updated time	Every about 1~2s
Minimum measurement	Surface: ≤100s
time	Coelom: ≤80s

A.8.6 IBP Specifications

A.8.6.1 Standard

Meet the standard of IEC 60601-2-34.

A.8.6.2 Functional specification

Measurement parameters Scale Unit	Dual channels IBP parameters (including systolic blood pressure, diastolic blood pressure, average pressure, PR) and waveforms Manual, interval and automatic scale settings mmHg, kPa, cmH ₂ O		
PPV	Measurement range 0%~50% Resolution 1%		
Static pressure measurement range and accuracy	Measurement range Resolution Accuracy	-6.7kPa ~ + 48.0kPa (-50mmHg ~ + 360mmHg) 1 mmHg ±0.3kPa (±2mmHg) or ±2%, whichever is greater (without sensor)	
Dynamic pressure measurement range and accuracy	Measurement range Accuracy	-6.7kPa \sim + 48.0kPa (-50mmHg \sim + 360mmHg) ±0.3kPa (±2mmHg) or ±2%, whichever is greater (without sensor)	
Frequency response IBP zero range	$ \begin{array}{ccc} \text{Including sensor} & 0\text{Hz}{\sim}10\text{Hz} \\ \\ \text{Only host} & 0\text{Hz}{\sim}12\text{Hz} \\ \\ \text{-200mmHg}{\sim}+200\text{mmHg} \\ \end{array} $		

PR	Measurement range	30bpm ~300bpm	
	Resolution	1bpm	
	Accuracy	±1% or ±1bpm whichever is greater	
	Nominal	5uV/V/ mmHg	
	sensitivity		
	Output	300Ω~3000Ω	
Pressure sensor	impedance	30032 -300032	
	Volumetric	<0.04 mm ³ /100 mmHg	
	displacement	0.04 mm /100 mming	
	Error	±2%	
IBP analog output	<35ms		
delay	2551119		

A.8.7 CO₂ Specifications

A.8.7.1 Standard

Meet the standard of ISO 80601-2-55.

A.8.7.2 Functional Specification

Measurement parameter	EtCO ₂ , FiCO ₂ , a CO ₂ waveform and awRR
Measurement method	Mainstream, Sidestream/Microflow
Unit	mmHg, kPa and %

A.8.7.3 Performance Specification

EtCO ₂ /FiCO ₂ measurement range	0%~19.7% (0mmHg~150mmHg)			
EtCO ₂ /FiCO ₂ measurement accuracy	\pm (0.43% + 8% of reading	$\pm (0.43\% + 8\% \text{ of reading})$		
Accuracy drift	Meets the requirement hours.	is for measurement accuracy within 6		
EtCO ₂ /FiCO ₂ display resolution	0.1% or 1mmHg			
awRR measurement range	0∼150 bpm			
awRR measurement accuracy	±1 bpm			
awRR display resolution	1 bpm			
awRR alarm range	0bpm to 150bpm, high/low limit can be adjusted continuously.			
Sampling frequency and accuracy of gas	C1 50 mL/min~200mL/min, ±10%, can be adjusted.			
(only sidestream)	Other sidestream CO ₂ 50 mL/min±10mL/min			
Response time	Sidestream CO ₂	<3s		
•	Mainstream CO ₂	<1s		

A.8.7.4 The effects on CO₂ measuring values caused by the interfering gases

> C11, C1 & C6

The accuracy of CO_2 is affected by interfering gases and water vapor. For example, N_2O , a halide-containing anesthetic gas can raise the CO_2 reading (about 2%-10%), and helium and oxygen can reduce the CO_2 reading (1%-10%), so in the presence of

interfering gas, the user should send relevant command to the module (the instrument's compensation menu to adjust the interference gas data), so that the module (instrument) can meet the nominal accuracy requirements.

> C5 & C2

Gas	Gas level	CO ₂		Agents	N ₂ O
		ISA CO ₂	ISA AX+	1	
			ISA OR+		
N ₂ O ⁴⁾	60 vol%	_ 2)	_ 1)	- ¹⁾	_ 1)
Hal ⁴⁾	4 vol%	_ 1)	- ¹⁾	- ¹⁾	_ 1)
Enf, Iso, Sev ⁴⁾	5 vol%	+8% of	- ¹⁾	- ¹⁾	_ 1)
		reading 3)			
Des 4)	15 vol%	+12% of	- ¹⁾	- ¹⁾	_ 1)
		reading 3)			
Xe (Xenon) ⁴⁾	80 vol%	−10% of read	ing 3)	- ¹⁾	_ 1)
He (Helium) 4)	50 vol%	−6% of readir	ng ³⁾	- ¹⁾	_ 1)
Metered dose	Not for use	with metered do	ose inhaler pro	pellants	•
inhaler					
propellants 4)					
C ₂ H ₅ OH	0.3 vol%	- ¹⁾	- ¹⁾	- 1)	_ 1)
(Ethanol) 4)					
C ₃ H ₇ OH	0.5 vol%	- ¹⁾	- ¹⁾	- ¹⁾	_ 1)
(Isopropanol) 4)					
CH ₃ COCH ₃	1 vol%	- ¹⁾	- ¹⁾	- ¹⁾	_ 1)
(Acetone) 4)					
CH ₄ (Methane) ⁴⁾	3 vol%	_ 1)	- ¹⁾	_ 1)	- ¹⁾
CO (Carbon	1 vol%	- ¹⁾	_ 1)	- ¹⁾	_ 1)
monoxide) 5)					
NO (Nitrogen	0.02 vol%	- ¹⁾	- ¹⁾	- ¹⁾	- ¹⁾
monoxide) 5)					
O ₂ 5)	100 vol%	_ 2)	_ 2)	_ 1)	

Note 1: Negligible interference, effect included in the specification "Accuracy, all conditions" above.

Note 2: Negligible interference with N_2O / O_2 concentrations correctly set, effect included in the specification "Accuracy, all conditions" above.

Note 3: Interference at indicated gas level. For example, 50 vol% Helium typically decreases the CO_2 readings by 6%. This means that if measuring on a mixture containing 5.0 vol% CO_2 and 50 vol% Helium, the actual measured CO_2 concentration will typically be (1-0.06) * 5.0 vol% = 4.7 vol% CO_2 .

Note 4: According to the EN ISO 80601-2-55:2011 standard.

Note 5: In addition to the EN ISO 80601-2-55:2011 standard.

A.8.8 AG Specifications

A.8.8.1 Standard

Meet the standard of ISO 80601-2-55.

A.8.8.2 Specification

Measurement way	Infrared spectrum
Measurement mode	Mainstream, Sidestream
Measurement parameters	Respiratory end gas fraction (Et), inhaled gas fraction (Fi), airway respiration rate (awRR) of Halothane (Hal), Enflurane (Enf), Isoflurane (Iso), Sevoflurane (Sev), Desflurane (Des), CO ₂ , N ₂ O, O ₂ (only be applicable for ISA OR+ module)
Resolution	Hal, Enf, Iso, Sev, Des, CO ₂ : 0.1% N ₂ O, O ₂ : 1%
Warm up time	< 20s (meets the requirement of accuracy)
Total system Response Time	Mainstream: <1s Sidestream: <4s
Sampling rate of sidestream	50mL/min±10mL/min

anesthesia gas		
Measurement range and accuracy of gas		
Gas	Range	Accuracy ¹⁾
CO ₂	0.0%~15.0%	± (0.43%+8% of reading)
N ₂ O	0%~100%	± (2%+8% of reading)
O ₂	0%~100%	± (2.5%+2.5% of reading)
Hal, Iso, Enf	$0.0\% \sim 8.0\%$ $\pm (0.2\% + 15\% \text{ of reading})$	
Sev	0.0%~10.0%	± (0.2%+15% of reading)
Des	0.0%~22.0%	± (0.2%+15% of reading)
awRR measurement range	0 bpm to 150 bpm	
awRR measurement accuracy	±1bpm	

A.8.9 ICG Specifications

Measurement	Measurement of thoracic electrical bioimpedance
way	
Measurement	C.O., C.I., SV, SI, SVR, SVRI, TFI, TFC, HR (the HR is obtained
parameters	directly from the ICG module)
Measurement	HR: 44 bpm to 185 bpm
range	SV: 5.0 mL to 250.0 mL
	C.O.: 1.4 L/min to 15.0 L/min
Resolution	HR: 1bpm
	SV: 0.1mL
	C.O.: 0.1 L/min
Alarm range	C.I.:1.4 L/min/m ² to 15.0 L/min/m ² , alarm error is ± 0.1 L/min/m ²
	TFC: 19 /k Ω to 125 /k Ω , alarm error is \pm 1/k Ω .

A.8.10 RM Specifications

Measure method	Flow sensor		
Paw	Measurement range -20.0 to 100.0 cmH ₂ O		
	Accuracy	±1 cmH ₂ O or ±3% of reading, whichever is	
		greater	
	Resolution	0.1 cmH ₂ O	
TVe/TVi	Measurement range	Adult/ Pediatric: 150 ml to 1600 ml	
		Neonate: 15 ml to 300 ml	
	Accuracy	Adult/Pediatric: ±10% or ±30 ml, whichever	
		is greater	
		Neonate: ±10% or ±6 ml, whichever is	
		greater	
	Resolution	1 ml	
MVe/MVi	Measurement range	Adult/Pediatric:2.00 L/min to 20.00 L/min	
		Neonate: 0.50 L/min to 5.00 L/min	
	Resolution	0.01 L/min	
RR	Measurement range	Adult/Pediatric: 4 bpm to 35 bpm	
		Neonate: 4 bpm to 50 bpm	
	Resolution	1 bpm	

A.8.11 EEG Specifications

A.8.11.1 Standard

Meet the standard of IEC 60601-2-26.

A.8.11.2 Specification

Sampling frequency	250 dots/s
Signal input Range	≥-2mV~+2mV
Polarization resistance	Add ±320mV d.c. bias voltage, the EEG waveform

voltage	amplitude deviation is within $\pm 5\%$.
Input impedance	≥15MΩ@10Hz
Amplitude frequency	Default 0.5 Hz ~30Hz (-3dB~+0.4dB); can expand to
characteristics	$0.25 \text{ Hz} \sim 110 \text{Hz} (-3 \text{dB} \sim +0.4 \text{dB})$ when low filter is
	turned off.
Noise level	≤3.0uV (0.5~30Hz)
CMRR	≥89dB (turn off the power frequency filter)
	≥100dB (turn on the power frequency filter)
Measurement range	SEF: 0.5 Hz -30.0 Hz
	MDF: 0.5 Hz -30.0 Hz
	PPF: 0.5 Hz -30.0 Hz
	TP: 0.0 Hz -100.0 dB
	Delta: 0.0%-100.0%
	Theta: 0.0%-100.0%
	Alpha: 0.0%-100.0%
	Bata: 0.0%-100.0%
Resolution	SEF: 0.1Hz
	MDF: 0.1Hz
	PPF: 0.1Hz
	TP: 0.1dB
	Delta: 0.1%
	Theta: 0.1%
	Alpha: 0.1%
	Beta: 0.1%

A.8.12 BIS Specifications

	Bispectral index (BIS), myoelectric activity (EMG), signal
Measurement Index	quality index (SQI), suppression ratio (SR), break count
	(BC), total power (TP), and spectral edge frequency (SEF)
Wave shape	Electroencephalo-graph waveform (EEG)
Parameter measurement	a) BIS measurement range is 0~100;

range	b) SQI measurement range is 0%~100%;
	c) SEF measurement range is 0.5 Hz ~30Hz;
	d) TP measurement range is 40.0dB~100.0dB.
	a) Duration: error ≤±5%;
	b) Noise(RTI) < 2uV;
	c) Amplitude frequency characteristics: 6Hz~30Hz
EEG measurement	(-3dB \sim +0.4dB) when turn on the filter; at least
specifications	0.5 Hz \sim 70Hz (-3dB \sim +0.4dB) when turn off the filter.
	d) CMRR: ≥100dB;
	e) Polarization resistance voltage: plus $\pm 300 \text{mV}$ d.c. bias
	voltage, EEG waveform amplitude deviation is within $\pm 5\%$.
	The monitor screen shows sensor graphs with numbers
	indicating the impedance value of electrodes.
Sensor detection display	Detection results are shown in two different colors: Green
	means passing, Yellow means high impedance, Grey
	means noise, Red means lead off.
Alarm range and error	BIS: 0~100, with the upper and lower limits continuously
setting	adjustable; alarm error: ±1.

A.8.13 NMT Specifications

Stimulation Modes	TOF、DB、PTC、SMC, Auto, TWI and TET
Current Range	0 mA \sim 80mA, error: \pm 5% or \pm 2mA, whichever is greater.
Maximum Stimulation	<400V
Voltage	
Stimulus	Monophasic square wave, pulse width was 0.2ms and
	error is $\pm 5\%$.
Stimulating Frequency	1Hz, 2Hz, 5Hz, 50Hz, 100Hz, error ±5%
	When the load impedance is 5 k Ω , the maximum peak
Maximum Load	output of the stimulus current is 70mA
Impedance	When the load impedance is 4 k Ω , the maximum peak
	output of the stimulus current is 80mA

A.8.14 rSO₂ Specifications

> Masimo rSO₂

Measurement Range		0%-99%
Calibration Range		45%-85%
		Invasive Co-oximeter
		The measured regional hemoglobin oxygen
		saturation value (rSO ₂) of the sensors is
		compared to arterial/venous hemoglobin
Calibration Stan	dard	oxygen (SavO ₂) value, determined from
		venous and arterial blood samples. The
		model used for blood in the brain is 70%
		venous and 30% arterial, which is applicable
		under normocapnic conditions
Accuracy (RMS,	trend)	≤3%
Accuracy (RMS,	absolute value)	Adult: ≤4%; Pediatric: ≤5%
Resolution		1%
$\Delta \mathrm{SpO}_2$	Measurement range	0%~99%
Δορο ₂	Resolution	1%
Percentage of	Measurement range	-100%~890%
rSO ₂ deviation Resolution		1%
baseline (ΔBL)		
Area under the Measurement range		0min·%∼9999min·%
curve (AUC)	Resolution	1min·%
Signal reliability		Has the signal credibility measurement
		function

➤ Nonin rSO₂

Measurement range	0%~99%
Resolution	1%

∆SpO ₂	Measurem range	ent	0%~99%			
	Resolution		1%			
Percentage of	Measurem	ent	-100%~890	%		
rSO ₂ deviation	range					
baseline (ΔBL)	Resolution		1%			
Area under the curve (AUC)	Measurem range	ent	0min⋅%~9999min⋅%			
curve (rice)	Resolution		1min·%			
Signal reliability	Has the signal credibility measurement function					
	1) 8004CA.	1) 8004CA, 8204CA: 50%-100%				
	Accuracy	Right	Left	Both	Hyperca pnia	Hypocap nia
Accuracy	absolute	4.1%	3.8%	3.9%	5.1%	3.3%
	trend	1.9%	3.0%	2.5%	3.4%	3.8%
	2) 8004CB, 8004CB-NA: 45-95%: ±5.9%					

Note: Accuracy specifications are based on the measured regional hemoglobin saturation value (rSO₂) of cerebral sensors calibrated to arterial / venous hemoglobin oxygen (SavO₂) value.

A.8.15 DM Specifications

A.8.15.1 Functional Specification

	Alarm and stop liquid when drip rate is abnormal.
Liquid stop function	Alarm and stop liquid when infusion is completed.
Enquiu stop runoutus	When the module is powered off, the liquid stop clip is opened
	without affecting the infusion.
	Drops/min, mL/h, can be automatically converted (for
Unit	conversion, 1mL of conventional tube =20 drops is mainly
	used.)

A.8.15.2 Performance Specification

Drip rate	5 Drops/min~200 Drops/min (1mL of conventional tube =20
measurement range	drops)
Drip accuracy	± 2 Drops/min and $\pm 2\%$, whichever is greater
Resolution	1 Drops/min

A.9 Alarm Specification

If no special instructions are given in the following specifications, the adjustable range of the alarm limit is the same as the measuring range of the signal.

A.9.1 ECG Alarm Specification

Alarm limit	Range	Step
ST High	(low limit +0.01 mV) ~2.00 mV	0.01 mV
ST Low	-2.00 mV ~ (high limit -0.01 mV)	0,01 111 1
HR High	(HR low limit +1bpm) ~350bpm	1bpm
HR Low	0bpm~ (HR high limit -1bpm)	Тори
QTc High	200ms~700ms	1ms
ΔQTc Low	-500ms~500ms	11110

A.9.2 RESP Alarm Specification

Alarm limit	Range	Step
RR High	(low limit +1 rpm) ∼150rpm	1rpm
RR Low	0rpm~ (high limit -1rpm)	r

A.9.3 NIBP Alarm Specification

BLT NIBP

Alarm limit	Range	Step
NIBP-S-High	Adult: (low limit+1 mmHg) ~270 mmHg	1mmHg
THE STINGS	Pediatric: (low limit+1 mmHg)~235 mmHg	111111115

Alarm limit	Range	Step
	Neonatal: (low limit+1 mmHg) ~135mmHg	
NIBP-S-Low	30 mmHg ~ (high limit-1 mmHg)	-
NIBP-M-High	Adult: (low limit+1mmHg) ~235 mmHg	
	Pediatric: (low limit+1 mmHg) ~225 mmHg	1mmHg
	Neonatal: (low limit+1 mmHg) ~125 mmHg	Tillining
NIBP-M-Low	20 mmHg ~ (high limit-1 mmHg)	
NIBP-D-High	Adult: (low limit+1mmHg) ~220 mmHg	
	Pediatric: (low limit+1 mmHg) ~220 mmHg	1mmHg
	Neonatal: (low limit+1 mmHg) ~110 mmHg	Timming
NIBP-D-Low	10 mmHg ~ (high limit-1 mmHg)	1

SunTech NIBP

Alarm limit	Range	Step
	Adult: (low limit+1 mmHg) ~260 mmHg	
NIBP-S-High	Pediatric: (low limit+1 mmHg)~160 mmHg	1mmHg
	Neonatal: (low limit+1 mmHg) ~130mmHg	Tillilirig
NIBP-S-Low	40 mmHg ~ (high limit-1 mmHg)	
NIBP-M-High	Adult: (low limit+1mmHg) ~220 mmHg	
	Pediatric: (low limit+1 mmHg) ~133 mmHg	1mmHg
	Neonatal: (low limit+1 mmHg) ~110 mmHg	Tillilling
NIBP-M-Low	26 mmHg ~ (high limit-1 mmHg)	
NIBP-D-High	Adult: (low limit+1mmHg) ~200 mmHg	
	Pediatric: (low limit+1 mmHg) ~120 mmHg	1mmHg
	Neonatal: (low limit+1 mmHg) ~100 mmHg	immig
NIBP-D-Low	20 mmHg ~ (high limit-1 mmHg)	

A.9.4 SpO₂ Alarm Specification

Alarm limit	Range	Step
SpO ₂ High	(low limit+1%)~100%	1%
SpO ₂ Low	$(SpO_2 Desat +1\%) \sim (high limit-1\%)$	1,0

SpO ₂ Desat	0%~ (low limit-1%)	
PR High	(PR low limit+1bpm)~350bpm	1bpm
PR Low	0bpm~ (PR high limit-1bpm)	Торш
\triangle SpO ₂ High	(low limit+1%)~100%	1%
△SpO ₂ Low	0bpm~ (high limit-1%)	
RR High	(low limit+1%) ~150bpm	1bpm
RR Low	0bpm~ (high limit-1bpm)	Topin

A9.5 TEMP Alarm Specification

Alarm limit	Range	Step
T1/T2 High	(low limit+0.1°C) ~50.0°C	0.1 °C
T1/T2 Low	0 °C~ (high limit-0.1°C)	0.1 °C
TD High	0°C~5.0°C	0.1 °C

A.9.6 IBP Alarm Specification

Alarm limit	Range	Step
IBP-M-High	(low limit +1mmHg) ~360mmHg	1mmHg
IBP-M-Low	-50mmHg~ (high limit -1mmHg)	111111111111111111111111111111111111111
IBP-D-High	(low limit +1mmHg) ~360mmHg	1mmHg
IBP-D-Low	-50mmHg~ (high limit -1mmHg)	111111111111111111111111111111111111111
NIBP-S-High	(low limit +1mmHg) ~360mmHg	1mmHg
NIBP-S-Low	-50mmHg~ (high limit -1mmHg)	111111111111111111111111111111111111111

A.9.7 CO₂ Alarm Specification

Alarm limit	Range	Step
Apnea delay time	20 s~60 s	5s
EtCO ₂ High	(low limit+1mmHg) ~152mmHg	1 mmHg
Et CO ₂ Low	0mmHg~ (high limit-1mmHg)	115
Fi CO ₂ High	0~152mmHg	1 mmHg
awRR High	(low limit+1bpm) ~150 bpm	1 bpm

Alarm limit	Range	Step
awRR Low	0bpm~ (high limit-1bpm)	

A.9.8 AG Alarm Specification

Alarm limit	Range	Step
Apnea delay time	20 s~60 s	5s
EtCO ₂ High	(low limit+1mmHg) ~114mmHg	1 mmHg
Et CO ₂ Low	0mmHg~ (high limit-1mmHg)	1 mminig
Fi CO ₂ High	0~152mmHg	1 mmHg
awRR High	(low limit+1bpm) ~150 bpm	1 bpm
awRR Low	0bpm~ (high limit-1bpm)	1 Opin
EtO ₂ / FiO ₂ High	(low limit+1%) ~100%	1%
EtO ₂ / FiO ₂ Low	18% ~ (high limit-1%)	1%
EtN ₂ O/ FiN ₂ O High	(low limit+1%) ~100%	1%
EtN ₂ O/ FiN ₂ O Low	0% ~ (high limit-1%)	1/0
EtHal/ EtEnf/ EtIso High	(low limit+0.1%) ~8.0%	
FiHal/ FiEnf/ FiIso High	1	0.1%
EtHal/ EtEnf/ EtIso Low	0% ~ (high limit-0.1%)	0.1%
FiHal/ FiEnf/ FiIso Low	1	
EtSev/ FiSev High	(low limit+0.1%) ~10.0%	0.10/
EtSev/ FiSev Low	0% ~ (high limit-0.1%)	0.1%
EtDes/ FiDes High	(low limit+0.1%) ~22.0%	0.1%
EtDes/ FiDes Low	0% ~ (high limit-0.1%)	0.170

A.9.9 ICG Alarm Specification

Alarm limit	Range	Step
C.I. High	$(low limit +0.1 L/min/m2) \sim 15.0 L/min/m2$	0.1 L/min/m ²
C.I. Low	$1.4 \text{ L/min/m}^2 \sim \text{(high limit-0.1 L/min/m}^2\text{)}$	0,11 2,11111,111
TFC High	(low limit $+1/k\Omega$) $\sim 125/k\Omega$	1 /kΩ
TFC Low	19 / kΩ~ (high limit-1 / kΩ)	1 / 1100

A.9.10 BIS Alarm Specification

Alarm limit	Range	Step
BIS High	(low limit +1) ~100	1
BIS Low	0 ~ (high limit-1)	

A.9.11 RM Alarm Specification

Alarm limit	Range	Step
PEEP High	(low limit +1 cmH ₂ O) ~100 cmH ₂ O	1 cmH ₂ O
PEEP Low	$0 \text{ cmH}_2\text{O} \sim \text{(high limit-1 cmH}_2\text{O)}$	1 6
PIP High	(low limit +1 cmH ₂ O) ~100 cmH ₂ O	1 cmH ₂ O
PIP Low	$1 \text{ cmH}_2\text{O} \sim \text{(high limit-1 cmH}_2\text{O)}$	
MVe High	Adult/Pediatric: (low limit +0.1 L/min) ~20.0 L/min	
III V V TIIGII	Neonate: (low limit +0.1 L/min) ~5.00 L/min	0.1 L/min
MVe Low	Adult/Pediatric: 2.00 L/min ~ (high limit-0.1 L/min)	011 2/11111
	Neonate: 0.50 L/min ~ (high limit-0.1 L/min)	
awRR High	(low limit+1bpm) ~150 bpm	1 bpm
awRR Low	0bpm~ (high limit-1bpm)	F

A.9.12 rSO₂ Alarm Specification

Alarm limit	Range	Step
rSO ₂ High	(low limit +1%) ~99%	1%
rSO ₂ Low	1% ~ (high limit-1%)	170
\triangle SpO ₂ High	(low limit +1%) ~99%	1%
\triangle SpO ₂ Low	0% ~ (high limit-1%)	170

Appendix B EMC and Radio Regulatory Compliance

B.1 EMC

The monitor complies with IEC 60601-1-2. All accessories listed in the accessories listed in the accessories of this manual meet the requirements of IEC 60601-1-2 when used with this equipment.



CAUTION:

- The monitor conforms to the electromagnetic compatibility requirements in IEC 60601-1-2, ISO 80601-2-55, IEC 80601-2-30, IEC 80601-2-49, ISO 80601-2-61, IEC 60601-2-34 standards.
- The user shall install and use according to the electromagnetic compatibility information provided by the accompanying documents.
- Portable and mobile RF communication equipment may affect the performance of this monitor, and strong electromagnetic interference should be avoided during use, such as close to mobile phones, microwave ovens, etc.
- The guidelines and the manufacturer's statement are detailed in the appendix.
- The device needs special precautions regarding EMC and needs to be installed and put into service according to the EMC information provided below.
- Portable and mobile communication equipment may affect the performance of this monitor.
- Other devices that have RF transmitter or source may affect this device (e.g. cell phones, PADs, PCs with wireless function).



WARNING:

■ The monitor should not be used close to or stacked on top of other equipment. If it must be used close to or stacked on top of other equipment, it should be observed and verified that it can operate normally under its used configuration.

- Class A equipment is intended to be used in industrial environment. Due to conduction disturbance and radiation disturbance of this monitor, there may be potential difficulties in ensuring electromagnetic compatibility in other environments.
- In addition to cables sold by the manufacturer of this monitor as spare parts for internal components, the use of accessories and cables other than those specified may result in increased emission or reduced immunity of this monitor.
- Even if other equipment meets the emission requirements of corresponding national standards, this monitor may still be interfered by other equipment.
- A warning that operation of the EQUIPMENT or SYSTEM below the minimum amplitude or value may cause inaccurate results. The minimum amplitude or value of patient physiological signal: the minimum amplitude of ECG signal is 0.5mV, the minimum value of PR is 30bpm and the minimum value of SpO₂ is 70%.

Table 1

Guidance and manufacture's declaration – electromagnetic emission

The monitor is intended for use in the environment specified below. The customer or the user of the monitor should assure that it is used in such environment.			
Emission test	Compliance	Electromagnetic environment - guidance	
RF emissions	Group1	The monitor uses RF energy only for its internal	
CISPR11		function. Therefore, its RF emissions are very low	
CISTRIT		and are not likely to cause any interference in	
		nearby electronic equipment.	
RF emission	Class A		
CISPR 11			
Harmonic emissions		The monitor is suitable for use in all establishments	
IEC 61000-3-2	Class A	other than domestic and those directly connected to the public low-voltage power supply network that	
Voltage fluctuations /		supplies building used for domestic purposes.	
flicker emissions	Complies		
IEC 61000-3-3			

If the system is operated within the electromagnetic environment listed in Table Guidance and Declaration – Electromagnetic Immunity, the system will remain safe and provide the following essential performance:

- **♦** Operating mode
- **♦** Accuracy
- **♦** Function
- **♦** Data stored
- **♦** Alarm
- **♦** Parameter
- **♦** Detect for connection

Table 2

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 monitor should assure that it is used in such an environment.

Immunity test	IEC60601 test level	Compliance level	Electromagnetic environment - guidance
Electrostatic	±8 kV contact	±8 kV contact	Floors should be wood,
discharge	±15 kV air	±15 kV air	concrete or ceramic tile. If
(ESD)			floors are covered with
IEC 61000-4-2			synthetic material, the
			relative humidity should be
			at least 30%.
Electrical fast	±2 kV for power	±2kV for power	Mains power quality should
transient/burst	supply lines	supply lines	be that of a typical
IEC 61000-4-4	±1 kV for input/output	±1 kV for	commercial or hospital
	lines	input/output lines	environment.
Surge	±1 kV line(s) to	±1 kV line(s) to	
	line(s)	line(s)	
IEC 61000-4-5	±2 kV line(s) to earth	±2 kV line(s) to	
		earth	
Voltage dips,	0 % UT; 0.5 cycle At	0 % UT; 0.5 cycle	Mains power quality should
short	0°, 45°, 90°, 135°,	At 0°, 45°, 90°,	be that of a typical
interruptions	180°, 225°, 270° and	135°, 180°, 225°,	commercial or hospital

and voltage	315°	270°and 315°	environment. If the user of
variations on			the monitor requires
power supply input lines IEC 61000-4-11	0 % UT; 1 cycle and 70 % UT; 25/30 cycles Single phase: at 0°	0 % UT; 1 cycle and 70 % UT; 25/30 cycles Single phase: at 0°	continued operation during power mains interruptions, it is recommended that the monitor be powered from an uninterruptible power supply or a battery.
	0 % UT; 250/300	0 % UT; 250/300	
	cycles	cycles	
Power frequency (50/60 Hz) magnetic field IEC 61000-4-8	30A/m	30A/m	Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.

NOTE: UT is the a.c. mains voltage prior to application of the test level.

Table 3

Guidance and	d manufact	ture's d	leclarat	ion – ele	ectromagneti	ic immunity
--------------	------------	----------	----------	-----------	--------------	-------------

The monitor is intended for use in the electromagnetic environment specified below. The customer or the user of monitor should assure that it is used in such an environment.

Immunity test	IEC 60601 test level	Complianc e level	Electromagnetic environment - guidance
Conducted RF IEC 61000-4-6	3Vrms 150kHz~ 80MHz	3Vrms	Portable and mobile RF communications equipment should be used no closer to any part of the <i>monitor</i> including cables, than the recommended separation
Radiated RF IEC 61000-4-3	3V/m 80MHz~ 2.7GHz	3V/m	distance calculated from the equation applicable to the frequency of the transmitter. Recommended separation distance: $d = \left[\frac{3.5}{V_1}\right]\sqrt{P} 150 \text{ KHz to } 80 \text{ MHz}$

	$d = \left[\frac{3.5}{E_1}\right] \sqrt{P} \qquad 80 \text{ MHz} \sim 800 \text{ MHz}$
	$d = \left[\frac{7}{E_1}\right] \sqrt{P} \qquad 80 \text{MHz} \sim 2.7 \text{GHz}$
	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:
	$(({\bf r}_{\bf A}))$

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.

Note 3: The device that intentionally receives RF electromagnetic energy at the exclusion band (2400-2483.5MHz) is exempt from the ESSENTIAL PERFORMANCE requirements, but remains safe.

- Field strengths from fixed RF 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 monitor is used exceeds the applicable RF compliance level above, the monitor should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as re-orienting or relocating the monitor.
- b Over the frequency range 0.15 MHz to 80 MHz, field strengths should be less than 3 V/m.

Table 4

Recommended separation distances between portable and mobile RF communications equipment and the monitor

The monitor is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the monitor can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the monitor.

Rated maximum	Separation dis	Separation distance according to frequency of transmitter				
output power of transmitter (w)	$150 \text{kHz} \sim 80 \text{MHz}$ $d = 1.2 \sqrt{P}$	$80MHz \sim 800MHz$ $d = 1.2\sqrt{P}$	80 MHz \sim 2.7 GHz $d = 2.3\sqrt{P}$			
0.01	0.12	0.12	0.23			
0.1	0.38	0.38	0.73			
1	1.20	1.20	2.30			
10	3.80	3.80	7.30			
100	12.00	12.00	23.00			

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.

B.2 The management compliance of Radio

RF Parameter

Radio frequency	Operating frequency	Modulation	Transmission power
transmitter			
WiFi	2.4GHz~2.495GHz	DSSS and	<20dBm (average value)
IEEE802.11a/b/g/n	5.15 GHz~5.35GHz	OFDM	<30dBm (Peak)
	5.47 GHz~5.725GHz		
	5.725 GHz~5.82GHz		

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The radio module used in the device is complied with the essential requirements and other relevant provisions of The Radio Equipment Directive.



WARNING:

■ Keep a distance of at least 20 cm away from the monitor when WIFI function is in use.

Appendix C Default Settings

This chapter lists some important factory default settings for monitors. The user cannot change the factory default settings, but the monitor can be restored to the factory default settings when necessary.

C.1 ECG, Arrhythmia, ST and QT Default Settings

C.1.1 ECG Default Settings

Item		Default Setting
HR	Alarm switch	ON
	Alarm high limit	Adult: 120 bpm
		Pediatric: 160 bpm
		Neonatal: 200 bpm
	Alarm low limit	Adult: 50 bpm
		Pediatric: 75 bpm
		Neonatal: 100 bpm
	Alarm priority	Med
	Alarm print	Off
	Alarm source	HR
ECG 1		II
ECG2 (5-lead	d, 6-lead, 12-lead)	I
Gain		×1
Wave Speed		25 mm/s
Filter Mode		Monitor
Notch Filter		On
Lead type		3-lead
QRS volume		3
Paced		Adult: Unspecified
		Pediatric/Neonatal: No
Pacer Reject		Off
		•

C.1.2 Arrhythmia Default Settings

Item	Alarm switch	Alarm priority	Alarm print
Asystole	ON	HIGH	OFF
Vent Fib/Tach	ON	HIGH	OFF
V-Tach	ON	HIGH	OFF
Vent Brady	ON	HIGH	OFF
Extreme Tachy	ON	HIGH	OFF
Extreme Brady	ON	HIGH	OFF
R on T	OFF	MED	OFF
Tachy	OFF	MED	OFF
Brady	OFF	MED	OFF
Nonsustained V-Tach	OFF	MED	OFF
Vent Rhythm	OFF	MED	OFF
PNC	OFF	MED	OFF
PNP	OFF	MED	OFF
Pause	OFF	MED	OFF
Pauses/min High	OFF	MED	OFF
Run PVCs	OFF	MED	OFF
Couplet	OFF	LOW	OFF
Bigeminy	OFF	LOW	OFF
Trigeminy	OFF	LOW	OFF
Frequent PVCs	OFF	LOW	OFF
PVC	OFF	LOW	OFF
Missed Beat	OFF	LOW	OFF
A-Fib	OFF	LOW	OFF
A-Fib End	OFF	LOW	OFF
ECG Noise	OFF	LOW	OFF
Irregular Rhythm	OFF	LOW	OFF
Irregular Rhythm End	OFF	LOW	OFF

C.1.3 ST Default Settings

Item		Default Setting
ST-I, ST-II,	Alarm switch	ON
ST-III, ST-aVR,	Alarm high limit	0.2 mV
ST-aVL,	Alarm low limit	-0.2 mV
ST-aVF, ST-V1,	Alarm priority	MED
ST-V2, ST-V3,	Alarm print	OFF
ST-V4, ST-V5,		
ST-V6, ST-Va,		
ST-Vb		
ST Analysis		OFF
ST Mark		OFF
Auto Adjust		OFF
ST Point		J + 60 ms
ISO		-80 ms
J		48 ms

C.1.4 QT Default Settings

Item		Default Setting
QTc	Alarm switch	OFF
	Alarm high limit	Adult: 500
		Pediatric: 480
		Neonate: 460
	Alarm priority	MED
	Alarm print	OFF
ΔQΤc	Alarm switch	OFF
	Alarm high limit	60
	Alarm priority	MED
	Alarm print	OFF
QT Analysis		OFF

C.2 RESP Default Settings

Item		Default Setting
RR	Alarm switch	ON
	High limit	Adult / Pediatric: 30
		Neonatal: 100
	Low limit	Adult / Pediatric: 8
		Neonatal: 30
	Alarm priority	MED
	Alarm print	OFF
Apnea	Alarm switch	ON
	Alarm priority	HIGH, unadjustable
	Alarm print	OFF
Apnea De	elay	20 s
RR Source	ce	Auto
RESP Le	ad	RA_LL
Gain		×1
Wave Speed		6.25 mm/s
Auto Threshold Detection		ON
Respirato	ory anti-drift	ON

C.3 SpO₂ Default Settings

Item		Default Setting
SpO_2	Alarm switch	ON
	High limit	Adult / Pediatric:100%
		Neonatal:95%
	Low limit	90%
	Alarm priority	MED
	Alarm print	OFF
Desat	Alarm switch	ON

Item		Default Setting
	Low limit	85%
	Alarm priority	HIGH
	Alarm print	OFF
NIBP Simul		OFF
Sensitivity		MED
Display PI		ON
Waveform S	peed	25 mm/s
PR	Alarm switch	ON
	High limit	Adult: 120 bpm
		Pediatric: 160 bpm
		Neonatal: 200 bpm
	Low limit	Adult: 50 bpm
		Pediatric: 75 bpm
		Neonatal: 100 bpm
	Alarm priority	MED
	Alarm print	OFF
	Alarm source	HR
	PR source	Auto
	QRS volume	3
	Pitch Tone	ON

C.4 TEMP Default Settings

Item		Default Setting
T1, T2	Alarm switch	ON
	High limit	38.0 °C
	Low limit	36.0 °C
	Alarm priority	MED
	Alarm print	OFF
ΔΤ	Alarm switch	ON

Item		Default Setting
	High limit	2.0 °C
	Alarm priority	MED
	Alarm print	OFF
Unit		°C

C.5 NIBP Default Settings

Item		Default Setting
NIBP-S	Alarm switch	ON
	High limit	Adult: 160 mmHg
		Pediatric: 120 mmHg
		Neonatal: 90 mmHg
	Low limit	Adult: 90 mmHg
		Pediatric: 70 mmHg
		Neonatal: 40 mmHg
	Alarm priority	MED
	Alarm print	OFF
NIBP-D	Alarm switch	ON
	High limit	Adult: 100 mmHg
		Pediatric: 70 mmHg
		Neonatal: 60 mmHg
	Low limit	Adult: 60 mmHg
		Pediatric: 40 mmHg
		Neonatal: 20 mmHg
	Alarm priority	MED
	Alarm print	OFF
NIBP-M	Alarm switch	ON
	High limit	Adult: 115 mmHg
		Pediatric: 90 mmHg
		Neonatal: 70 mmHg
	Low limit	Adult: 70 mmHg

Item		Default Setting
		Pediatric: 50 mmHg
		Neonatal: 25 mmHg
	Alarm priority	MED
	Alarm print	OFF
NIBP-sdp	Alarm switch	ON
	High limit	60mmHg
	Low limit	20mmHg
	Alarm priority	MED
	Alarm print	OFF
Initial Pressur	e	Adult: 160 mmHg
		Pediatric: 130 mmHg
		Neonatal: 100 mmHg
Interval		manual
Start Mode		Clock
NIBP End Tone		OFF
Venipuncture pressure		Adult: 80mmHg
		Pediatric: 60mmHg
		Neonatal: 40mmHg
Unit		mmHg

C.6 IBP Default Settings

Item		Default Setting
IBP-S	Alarm switch	ON
	High limit	ART/Ao/UAP/BAP/FAP/LV/P1/P2 arterial pressure
		Adult: 160 mmHg
		Pediatric: 120 mmHg
		Neonatal: 90 mmHg
		PA/PAWP:
		Adult: 35 mmHg
		Pediatric/Neonatal: 60mmHg

Item		Default Setting
	Low limit	ART/Ao/UAP/BAP/FAP/LV/P1/P2 arterial pressure
		Adult: 90 mmHg
		Pediatric: 70 mmHg
		Neonatal: 55 mmHg
		PA/PAWP:
		Adult: 10 mmHg
		Pediatric/Neonatal: 24 mmHg
	Alarm priority	MED
	Alarm print	OFF
IBP-D	Alarm switch	ON
	High limit	ART/Ao/UAP/BAP/FAP/LV/P1/P2 arterial pressure
		Adult: 90mmHg
		Pediatric: 70 mmHg
		Neonatal: 60mmHg
		PA/PAWP:
		Adult: 16 mmHg
		Pediatric/Neonatal: 4 mmHg
	Low limit	ART/Ao/UAP/BAP/FAP/LV/P1/P2 arterial pressure
		Adult: 50 mmHg
		Pediatric: 40 mmHg
		Neonatal: 20 mmHg
		PA/PAWP:
		Adult: 0 mmHg
		Pediatric/Neonatal: -4 mmHg
	Alarm priority	MED
	Alarm print	OFF
IBP-M	Alarm switch	ON
	High limit	ART/Ao/UAP/BAP/FAP/LV/P1/P2 arterial pressure
		Adult: 110 mmHg
		Pediatric: 90 mmHg
		Neonatal: 70 mmHg
	1	ļ

		PA/PAWP
		ra/rawr
		Adult: 20 mmHg
		Pediatric/Neonatal: 26 mmHg
		CVP/ICP/RAP/LAP/UVP Venous pressure
		Adult: 10 mmHg
		Pediatric/Neonatal: 4 mmHg
	Low limit	ART/Ao/UAP/BAP/FAP/LV/P1/P2 arterial pressure
		Adult: 70 mmHg
		Pediatric: 50 mmHg
		Neonatal: 35 mmHg
		PA/PAWP
		Adult: 0 mmHg
		Pediatric/Neonatal: 12 mmHg
		CVP/ICP/RAP/LAP/UVP venous pressure
		Adult: 0 mmHg
		Pediatric/Neonatal: 0 mmHg
	Alarm priority	MED
	Alarm print	OFF
CPP	Alarm switch	ON
	High limit	Adult: 130 mmHg
		Pediatric: 100 mmHg
		Neonatal: 90 mmHg
	Low limit	Adult: 50 mmHg
		Pediatric: 40 mmHg
		Neonatal: 30 mmHg
	Alarm priority	MED
	Alarm print	OFF
Unit		ART/Ao/UAP/BAP/FAP/LV/RAP/LAP/UVP/PA/PAW
		P/P1/P2: mmHg
		CVP /ICP/CPP: cmH ₂ O
Sensitivity		MED

Item		Default Setting
Wave Speed		25 mm/s
Scale Type		Manual
Scale	Upper scale	ART /Ao/UAP/BAP/FAP/LV/P1/P2 arterial pressure:
		160mmHg
		CVP /ICP/RAP/LAP/UVP venous pressure : 20mmHg;
		PA/PAWP: 30mmHg
	Lower scale	0mmHg
High-precisio	n cursor	OFF
switch		
High-precisio	n cursor	ART/ Ao/ UAP/BAP/FAP/LV/P1/P2 arterial pressure:
		80mmHg
		CVP /ICP/RAP/LAP/UVP venous pressure: 10mmHg
		PA/PAWP: 15mmHg
PPV Measure	ment	OFF
PPV Source		Auto
Overlapping	Left Scale	0~160mmHg
IBP	Right Scale	P1/P2: 0~160mmHg
Waveforms		CVP/RAP/LAP/ICP/UVP: 0 ~20 mmHg
	Wave speed	25 mm/s
	Grid	OFF

C.7 CO₂ Default Settings

Item		Default Setting
EtCO ₂	Alarm switch	ON
	High limit	Adult/Pediatric: 50 mmHg
		Neonatal: 45 mmHg
	Low limit	Adult/Pediatric: 25mmHg
		Neonatal: 30mmHg
	Alarm	MED
	priority	

Item		Default Setting
	Alarm print	OFF
FiCO ₂	Alarm switch	ON
	High limit	4 mmHg
	Alarm	MED
	priority	
	Alarm print	OFF
Apnea delay	y	Adult/Pediatric: 20s; Neonate:15s
Wave Speed	d	6.25 mm/s
Scale		50 mmHg
Wave Mode	e	Draw
Operation Mode		Measurement mode
Unit		mmHg
Gas temperature		35 °C
Barometric	Pressure	760mmHg
O ₂ Concentration		16%
N ₂ O Concentration		0%
Zero Gas Type		Air
Anesthetics Gas		0%
Balance Gas		Air

C.8 DM Default Settings

Item	Default Setting
Unit	Drops/min
Drop Per Milliliter	20

C.9 AG Default Settings

Item		Default Setting
EtCO ₂	Alarm switch	ON
	High limit	Adult/Pediatric: 50 mmHg
		Neonatal: 45 mmHg

Item		Default Setting
	Low limit	Adult/Pediatric: 25mmHg
		Neonatal: 30mmHg
	Alarm	MED
	priority	
	Alarm print	OFF
FiCO ₂	Alarm switch	ON
	High limit	4 mmHg
	Alarm	MED
	priority	
	Alarm print	OFF
EtO ₂	Alarm switch	ON
	High limit	88%
	Low limit	18%
	Alarm	MED
	priority	
	Alarm print	OFF
FiO ₂	Alarm switch	ON
	High limit	Adult/Pediatric: 100%
		Neonatal: 90%
	Low limit	18%
	Alarm	MED
	priority	
	Alarm print	OFF
EtN ₂ O	Alarm switch	ON
	High limit	55%
	Low limit	0%
	Alarm	MED
	priority	
	Alarm print	OFF
FiN ₂ O	Alarm switch	ON

Item		Default Setting
	High limit	53%
	Low limit	0%
	Alarm	MED
	priority	
	Alarm print	OFF
EtHal/	Alarm switch	ON
EtEnf/	High limit	3.0%
EtIso	Low limit	0.0%
	Alarm	MED
	priority	
	Alarm print	OFF
FiHal/	Alarm switch	ON
FiEnf/	High limit	2.0%
FiIso	Low limit	0.0%
	Alarm	MED
	priority	
	Alarm print	OFF
EtSev	Alarm switch	ON
	High limit	6.0%
	Low limit	0.0%
	Alarm	MED
	priority	
	Alarm print	OFF
FiSev	Alarm switch	ON
	High limit	5.0%
	Low limit	0.0%
	Alarm	MED
	priority	
	Alarm print	OFF
EtDes	Alarm switch	ON

Item		Default Setting
	High limit	8.0%
	Low limit	0.0%
	Alarm	MED
	priority	
	Alarm print	OFF
FiDes	Alarm switch	ON
	High limit	6.0%
	Low limit	0.0%
	Alarm	MED
	priority	
	Alarm print	OFF
Apnea Dela	ıy	20s
Wave Speed	d	6.25 mm/s
Scale		CO ₂ : 50 mmHg
		O ₂ : 400 mmHg
		N ₂ O: 50%
		Hal/Enf/Iso: 2.5%
		Sev: 4.0%
		Des: 9.0%
Wave Mode		Draw
Operation Mode		Measurement mode
Unit		mmHg
Gas temperature		35 °C
Atmospheric		760mmHg
O ₂ Compensation		OFF

C.10 RM Default Settings

Item		Default Setting
PEEP	Alarm switch	ON
	High limit	10 cmH ₂ O

Item		Default Setting
	Low limit	0 cmH ₂ O
	Alarm	MED
	priority	
	Alarm print	OFF
PIP	Alarm switch	ON
	High limit	40 cmH ₂ O
	Low limit	1 cmH ₂ O
	Alarm	MED
	priority	
	Alarm print	OFF
MVe	Alarm switch	ON
	High limit	8.0 L/min
	Low limit	2.0 L/min
	Alarm	MED
	priority	
	Alarm print	OFF
Wave Speed	d	6.25 mm/s
Paw Scale		40 cmH ₂ O
Flow Scale		60 L/min
Vol Scale		1200ml
Atmosphere Temp		25 °C
Relative Humidity		55%
Paw		PIP, PEEP, Pmean, Pplat, RR
Flow		IE,Resi,Compl
Vol		MVe, MVi, TVe, TVi

C.11 BIS Default Settings

Item		Default Setting
BIS	Alarm switch	ON
	High limit	70

Item		Default Setting
	Low limit	20
	Alarm	MED
	priority	
	Alarm print	OFF
Average tin	ne	30s
Waveform Speed		25 mm/s
Scale		100μV
Dispaly		EEG
Filter		ON

C.12 EEG Default Settings

Item		Default Setting
Scale		100μV
Wave Spe	eed	25 mm/s
Lowpass	Filter	0.5Hz
Highpass	Filter	30Hz
Notch Fil	ter	ON
SEF Thre	shold	95%
EEG exp	oand screen	
EEG	EEG Channel	EEG1, EEG2, EEG3, EEG4
	Scale	100μV
	Speed	25 mm/s
Trend	EEG Channels	EEG2, EEG3, EEG4
	Parameters	SEF
	Trend Length	60min
DSA	EEG Channels	All
	Parameters	SEF
	Trend Length	20min
	Power Scale	1 × 64 dB
CSA	EEG Channels	All

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Item		Default Setting
	Parameters	SEF
	Trend Length	20min
	Power Scale	1 × 64 dB
	CSA Clipping	ON

A.13 ICG Default Settings

Item		Default Setting
C.I.	Alarm switch	ON
	High limit	5.0 L/min/m ²
	Low limit	1.5 L/min/m ²
	Alarm	MED
	priority	
	Alarm print	OFF
TFC	Alarm switch	ON
	High limit	60/kΩ
	Low limit	20/kΩ
	Alarm	MED
	priority	
	Alarm print	OFF
Wave Speed	d	25 mm/s
Setup Parameters		C.I., SQI, SVI, SVRI, TFC

C.14 rSO₂ Default Settings

Item		Default Setting
rSO ₂ -1,	Alarm switch	ON
rSO ₂ -2	High limit	90%
	Low limit	40%
	Alarm priority	MED
	Alarm print	OFF
rSO ₂ -1 %BL,	Low limit	-20

Item		Default Setting
rSO ₂ -2 %BL		
Auto Low Limit		OFF
rSO ₂ -1 Sensor S	ite	The left forehead
rSO ₂ -2 Sensor S	ite	The right forehead
AUC Mode		Below Baseline Percentage
Fixed		50
Below Base Per	centage	25
Select parameter	r	Baseline

C.15 NMT Default Settings

Item	Default Setting
Stimulation current	15mA
Stimulation mode	TOF
Stimulation frequency	50Hz

C.16 Alarm Default Settings

Item	Default Setting
Alarm Volume	2
High Alarm Volume	Alarm volume +2
Reminder Volume	5
Apnea Delay	20s
Alarm Record Duration	8s

C.17 Screen Setup Default Settings

Item	Default Setting
Screen Select	Standard
Screen Lock Duration	2min
Brightness	5
Brightness On Battery	1

C.18 Color of parameters Default Settings

Item	Default Setting
ECG	Green
NIBP	White
SpO_2	Yellow
TEMP	Purple
RESP	Cyan
CO ₂	White
DM	Yellow-green
IBP	Red
O2	Blue
N ₂ O	Yellow
AA	Pink
ICG	White
BIS	White
EEG1	Red
EEG2	Blue
EEG3	Yellow
EEG4	Green
RM	White
rSO ₂ -1	Blue
rSO ₂ -2	Yellow
NMT	White

C.19 Recorder Default Settings

Item	Default Setting
Waveform 1	П
Waveform 2	Pleth
Waveform 3	RR
Record Speed	25mm/s
Record Duration	8s

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Cycle Record Interval	OFF
Cycle Record Duration	8s
Alarm Record Duration	8s
NIBP Trigger	OFF

C.20 Other Default Settings

Item		Default Setting
Keypad tone		ON
Night	Brightness	1
Mode	Alarm volume	2
	QRS volume	1
	Touch Tone	OFF
	NIBP End	OFF
	Tone	

C.21 Maintenance Item Default Settings

Item	Default Setting
Network Type	LAN
LAN IP	Use the Following Address
Frequency	2.4G
Device No.	8
Alarm Pause Duration	2min
Minimum alarm volume	2
Alarm Sound	ISO
High Alarm Interval (s)	10
Medium Alarm Interval (s)	20
Low Alarm Interval (s)	20
Reset Remote Bed's Alarms	OFF
Alarm Reset By Other Bed	OFF
Alarm Off	ON
Reminder Interval	5min
ECG Lead Off Alarm Level	MED

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Alarm delay		OFF
Notch filter		50 Hz
Nurse	Signal type	Continuous
call	Trigger method	N.O.
	Alarm level	All
	Alarm type	Technical Alarm & Physiological Alarm

Appendix D Alarm Message

This chapter lists some of the most important physiological and technical alarm information, and some alarm information may not be listed.

D.1 Physiological alarm

D.1.1 General physiological alarm

Alarm messages	Default priority	Cause and solution
XX High	MED	The measured value of the corresponding parameter is higher than the alarm high limit. Please check the patient's physiological condition and confirm whether the patient type and alarm limit settings are applicable to the patient.
XX Low	MED	The measured value of the corresponding parameter is lower than the alarm high limit. Please check the patient's physiological condition and confirm whether the patient type and alarm limit settings are applicable to the patient.

Note: XX represents the nominal name of physiological parameter, such as HR, ST, RR, SpO_2 or PR, etc.

D.1.2 Arrhythmia alarm information

Alarm messages	Default priority	Alarm messages	Default priority
Asystole	HIGH	Pauses/min High	MED
Vent Fib/Tach	HIGH	Run PVCs	MED
V-Tach	HIGH	Couplet	LOW
Vent Brady	HIGH	Bigeminy	LOW
Extreme Tachy	HIGH	Trigeminy	LOW
Extreme Brady	HIGH	Frequent PVCs	LOW
R on T	MED	PVC	LOW

Alarm messages	Default priority	Alarm messages	Default priority
Tachy	MED	Missed Beat	LOW
Brady	MED	A-Fib	LOW
Nonsustained V-Tach	MED	A-Fib End	LOW
Vent Rhythm	MED	ECG Noise	LOW
PNC	MED	Irregular Rhythm	LOW
PNP	MED	Irregular Rhythm End	LOW
Pause	MED		

D.1.3 RESP Physiological Alarm

Alarm messages	Default priority	Cause and solution
		The patient is not breathing or the respiratory
		signal is too weak to measure the respiratory
DECD Anno	High	rate. Please check the patient's condition,
RESP Apnea	High	check whether the electrode plate is placed
		correctly and whether the connection of
		electrode plate, cable and lead wire is firm.
	High	The patient's heartbeat interferes with
		breathing, thus making it impossible to
RESP Artifact		measure the breathing rate correctly. Please
REST AITHACT		check the patient's condition and check the
		connection of electrode plates, cables and lead
		wires.

D.1.4 SpO₂ Physiological Alarm

Alarm messages	Default priority	Cause and solution
SpO ₂ Search Pulse Timeout	High	Can't find a pulse for a long time. Please immediately check the patient's condition. If the patient is normal condition, please replace placement position of blood oxygen probe.

Alarm messages	Default priority	Cause and solution
SpO ₂ Desat	High	SpO ₂ measurement is below the desaturation
		limit. Please check the patient's status and
		confirm whether the alarm limit setting is
		applicable to the patient.

D.1.5 CO₂ Physiological Alarm

Alarm messages	Default priority	Cause and solution
		The patient is not breathing or the
		respiratory signal is too weak to measure
CO ₂ Apnea	High	the respiratory rate. Check the patient's
		condition and whether the air circuit
		connection is correct.

D.1.6 AG Physiological Alarm

Alarm messages	Default priority	Cause and solution
		The patient is not breathing or the
	High	respiratory signal is too weak to measure
CO ₂ Apnea		the respiratory rate. Check the patient's
		condition and whether the air circuit
		connection is correct.
O ₂ Fi Extremely low	High	Check the patient's condition, O ₂
(<18%)	High	concentration and aire connection.

D.1.7 RM Physiological Alarm

Alarm messages	Default priority	Cause and solution
		The patient is not breathing or the
		respiratory signal is too weak to measure
CO ₂ Apnea	High	the respiratory rate. Check the patient's
		condition and whether the air circuit
		connection is correct.

D.2 Technical Alarm

This chapter lists the main technical alarms, the level of technical alarms, the cleared status of alarm reset alarm prompts, and the measures to be taken after the alarm occurs. Some alarm messages may not be listed.

After different technical alarm is reset, the alarm prompt will be cleared to different degrees. The following three types of technical alarms are given in this section according to the status of alarm being cleared.

- Completely clear: the technical alarm is completely clear. The monitor has no alarm indication.
- Sound and light can be cleared: the technical alarm displays as prompt information.
- Not clearable: the sound of technical alarm is shielded.

D.2.1 General Technical alarm

Alarm messages	Default	Alarm clear	Cause and solution
Atai iii iiiessages	priority	method	
XX			XX measurement module failure or
Communication error	Med	Not clearable	communication failure.

Note: "XX" represents the module name, such as ECG, SpO₂, IBP, TEMP, etc.

D.2.2 ECG Technical Alarm

A1.	Default	Alarm clear	Cause and solution
Alarm messages	priority	method	
ECG Self-test	M- J	NI-4 -1	Board failure. Please contact the
Error	Med	Not clearable	manufacturer for repair.
			All ECG leads fall off or ECG cables
ECG Leads Off	Med	Sound and light	are not connected. Please check the
ECG Leads Off	Med	can be cleared	connection of ECG electrode plates,
			lead wires and cables.
ECG XX Off	Med	Sound and light	The electrode is not firmly connected

A1	Default	Alarm clear	Cause and solution	
Alarm messages	priority			
		can be cleared	with the patient or falls off, causing the	
			corresponding ECG lead to fall off.	
			Please check the connection of ECG	
			electrode plates, lead wires and cables.	
ECG YY		Sound and light	ECG electrode polarization or poor	
Polarized	Low	can be cleared	contact. Please check the connection of	
Folarized		can be cleared	ECG electrodes plates.	
ECC Iin-	D	/	Relearn is triggered manually or	
ECG Learning	Prompt	/	automatically	
ECG Cable	Med	Not clearable	Use non-factory cables. Please replace	
Incompatible	Mcd	Not clearable	the original cable.	
ECG Cable Has	Med	Not clearable	ECG cable has expired. Please replace	
Expired	Med	Not clearable	the cable.	
ECG Cable is	Prompt	/	ECG cable is about to expire. Please	
About to Expire	Trompt		replace the cable in time.	
			Pacing signal has been detected by	
ECG Suspected			non-pacing patients. Please check	
Pacing Signal	Prompt	/	whether the patients have pacemakers	
1 acing Signal			and check the connection of ECG	
			electrode sheets.	
Note: XX represents RA, LA, LL, RL, V1, V2, V3, V4, V5, V6,				

YY represents I, II, III, aVR, aVL, aVF, V1, V2, V3, V4, V5 or V6.

D.2.3 RESP Technical Alarm

Alarm messages	Default priority	Alarm clear method	Cause and solution
RESP Leads Off	Med	Completely clear	ECG lead-off or the ECG cable is not connected. Check the communication of ECG electrode and lead wires.

D.2.4 SpO₂ Technical Alarm

	Default	Alarm clear	Cause and solution
Alarm messages	priority	method	
SpO ₂ / SpO ₂ L Self-test Error	Med	Not clearable	Board failure. Please contact the manufacturer for repair.
SpO ₂ / SpO ₂ L Sensor Off	Med	Sound and light can be cleared	The SpO ₂ sensor is falled off from the patient end. Check the connection of the sensor. If the alarm still exists, replace the sensor.
SpO ₂ / SpO ₂ L Sensor Disconnected	Low	Sound and light can be cleared	The SpO ₂ main cable falls off from the module end or the connection between the SpO ₂ sensor and the SpO ₂ main cable falls off. Confirm that SpO ₂ main cable and sensor are connected normally. If the alarm still cannot be eliminated, replace the sensor.
SpO ₂ / SpO ₂ L Low Confidence	Low	Not clearable	PI<0.3% or signal quality <60.
SpO ₂ / SpO ₂ L Update Timeout	Low	Not clearable	25s SpO ₂ measurement data not updated.
SpO ₂ / SpO ₂ L Motion Interference	Low	Not clearable	Patients move too much, affecting measurement.
SpO ₂ / SpO ₂ L Searching Pulse	Prompt	/	SpO ₂ module is searching for pulse.
SpO ₂ / SpO ₂ L Sensor Incompatible	Med	Not clearable	Non-factory SpO ₂ sensor are used. Please replace the original sensor.
SpO ₂ / SpO ₂ L Sensor Has Expired	Med	Not clearable	SpO ₂ sensor has expired. Please replace the sensor.

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SpO ₂ / SpO ₂ L Sensor	Prompt	/	SpO ₂ sensor is about to expire.
is About to Expire	Frompt	/	Please replace the sensor in time.

D.2.5 TEMP Technical Alarm

Alarm messages	Default priority	Alarm clear method	Cause and solution
TEMP Self-test	Med	Not clearable	Board failure. Please contact the
Error			manufacturer for repair.
⟨TEMP label⟩	Med	Completely	Check the connection of the sensor and
Sensor Off		clear	reconnect the sensor.

D.2.6 NIBP Technical Alarm

Alarm messages	Default	Alarm clear	Cause and solution
Alarm messages	priority	method	
NIBP Self-test	Med	Not clearable	Board failure. Please contact the
Error	Med	Not clearable	manufacturer for repair.
NIBP System	Low	Not clearable	System operation failure.
Failure	Low	Not clearable	System operation failure.
NIBP Air Pressure	Low	Completely	Pressure error, unable to maintain stable
Error	Low	clear	cuff pressure, such as tracheal knot.
NIBP		Completely	NIBP air leakage was found in the
Air Leakage	Low	clear	inspection. Please check the sleeve and
All Leakage			airpipe for air leakage.
NIBP Air System	Low	Completely	Damaged cuff, hose or joint.
Leak	Low	clear	Damaged curr, nose of joint.
			The cuff used does not match the patient
			type set. Please confirm that the patient
NIBP Cuff Type Error	Low	Completely clear	type is set correctly and select correct
			cuff according to the patient type. If the
			patient type and cuff selection are
			correct, please check whether the airway

Alarm messages	Default priority	Alarm clear method	Cause and solution
			and airpipe are bent or blocked.
NIBP Overpressure Detected	Med	Completely clear	The pressure exceeds the specified safety limit.
NIBP Loose Cuff	Low	Completely clear	The cuff is not tight; Or the cuff is not connected. Select the correct cuff according to the patient type, place the cuff according to the manual, and connect the airpipe.
NIBP Excessive Motion	Low	Completely clear	The patient moved frequently during the measurement. Or violent movement during measurement; Or irregular pulse rate, such as arrhythmia.
NIBP Signal Saturated	Low	Completely clear	Great movement.
NIBP Weak Signal	Low	Completely clear	The cuff is too loose or the patient's pulse is too weak. Please check the patient's condition or whether the cuff is placed correctly.
NIBP Out of Range	Low	Completely clear	The measurement range exceeds the specified upper limit.
NIBP Time Out	Low	Completely clear	Measurement time exceeds 120s (adult/pediatric) or 90s (neonatal). Please check the patient's condition and the connection of accessories or replace the cuff, and conduct the measurement again.
NIBP Cycle Abort	Low	Completely clear	Three consecutive measurement failures occurred during periodic measurement.

Alaum magagag	Default	Alarm clear	Cause and solution
Alarm messages	priority	method	
			Please check whether the patient's
			condition or cuff placement is correct.
NIBP Zero Failed	Duament	/	At zero, the pressure is beyond the zero
	Prompt		range or the pressure is unstable.

D.2.7 IBP Technical Alarm

A1	Default	Alarm clear	Cause and solution
Alarm messages	priority	method	
IBP Self-test Error	Med	Not clearable	The board is failure. Please contact the factory for repair.
XX Sensor Off	Med	Completely clear	The XX cable is off the monitor.
XX Pressure Calibrate Failed	Med	Not clearable	When the XX sensor is zeroed, the sensor is not connected or the pressure is out of range or the pressure is unstable. The catheter is pulled out from the
XX Catheter Off	HIGH	Not clearable	patient. Please check the connection.
Zero Required	Prompt	/	/
XX Pressure Calibration Succeed	Prompt	/	The IBP module is zeroing successful
Note: XX represents IBP labels, such as PA, CVP, FAP, P1, etc.			

D.2.8 ICG Technical Alarm

Alarm messages	Default	Alarm clear	Cause and solution
Alai iii iiiessages	priority	method	
ICG Hardware	Med	Not clearable	The ICG module is damaged,
Error	Med	Not clearable	please replace it with a new one.
ICG Sensor Off	Med	Completely clear	Please check the connection status

Alaum magagag	Default	Alarm clear	Cause and solution
Alarm messages	priority	method	
			of the sensor and reconnect the
			sensor.
			The electrode is not firmly
			attached to the patient or falls off
ICG Electrodes Off	Med	Sound and light	causing the corresponding ICG
ico Electrodes Off	Med	can be cleared	lead off. Please check the
			connection of ICG electrode, lead
			wire and cable.
			ICG monitoring only be applicable
ICG Invalid Patient	Low	Sound and light	for patients in height of 122 to
Setting		can be cleared	229cm, weight of 30 to 155 kg,
			and in age no less than 13.
ICG Invalid Blood		Sound and light	The blood pressure parameter
Pressure Setting	Low	can be cleared	cannot be obtained from IBP or
Tressure Setting		can be cleared	NIBP. Manual input is required.
ICG External			The two sensors must be placed
Measurement	Low	Sound and light	right on the opposite position
Setting	Low	can be cleared	(180°). The current status of cable
Setting			isn't ready.
ICG Invalid	Med	Not clearable	ICG hardware is invalid.
Hardware	ivieu	TAUL CICALAUIC	

D.2.9 CO₂ Technical Alarm

Alarm messages	Default priority	Alarm clear method	Cause and solution
CO ₂ Sensor Off	Med	Completely clear	The CO ₂ sensor is detached from the patient or monitor.
CO ₂ Out of Range	Low	Not clearable	The measured data of CO_2 module is out of range and needs zero.

Alarm messages	Default	Alarm clear	Cause and solution
Alai iii iiicssages	priority	method	
CO ₂ Zero Required	Low	Not clearable	The sensor needs zero.
CO ₂ Sensor Over Temp	Low	Not clearable	Check sensor.
CO ₂ Compensation Not Set	Low	Not clearable	The CO ₂ sensor was not initialized. Set compensation and initialize.
CO ₂ Sleep Mode	Prompt	/	The CO_2 sensor is in sleep mode. Please select the measurement mode, CO_2 can enter the working state.
CO ₂ Check Sampling Line	Low	Not clearable	The CO ₂ sampling tube is blocked or damaged; The sampling tube is kinked or compacted; The exhaust pipe is blocked. Check the sampling tube.
CO ₂ Check Adapter	Low	Not clearable	Reinstall the airway adapter.
CO ₂ Zero In Progress	Prompt	/	The CO ₂ module is being zeroed.
CO ₂ Sensor Warm up	Prompt	/	The CO ₂ module is warming up.
CO ₂ Self-Test	Prompt	/	Module initialization
CO ₂ Sensor Faulty, E*	Low	/	Hardware or software errors, contact after-sales personnel to check maintenance.
CO ₂ Self-test Error	Low	Not clearable	Hardware errors, Replace sensor, if the problem cannot be solved, please contact the after-sales personnel to check the maintenance.
CO ₂ Motor Speed	Low	Not clearable	Check whether the sampling tube

Alarm messages	Default priority	Alarm clear method	Cause and solution
Error			is blocked.
CO ₂ Factory Calibration lost	Low	Not clearable	Contact the after-sales personnel to check the maintenance.
CO ₂ Sampling Line Clogged	Low	Not clearable	Check sampling line.
CO ₂ No Sample Line	Low	Completely clear	Check sampling line.
CO ₂ Internal Temp.Out	Low	Not clearable	Hardware error, and contact the after-sales personnel to check the maintenance.
CO ₂ Ambient Pressure Out of Range	Low	Not clearable	Recalibrate atmospheric pressure.
CO ₂ Span Calibration Command Failed	Low	Not clearable	Contact the after-sales personnel to check the maintenance.
CO ₂ Span Calibration in Progress	Prompt	/	Disappear after success.
CO ₂ Replace Adapter	Low	Not clearable	Check adapter.
CO ₂ No Adapter	Low	Completely clear	Check adapter.

D.2.10 AG Technical Alarm

Alarm messages	Default priority	Alarm clear method	Cause and solution
O ₂ Sensor Error	Low	Not clearable	Hardware or software error, contact the after-sales personnel to check the maintenance.
O ₂ Port Failure	Low	Not clearable	Hardware or software error, contact the after-sales personnel to check the maintenance.
O ₂ Out of Range	Low	Not clearable	The measured data of O_2 module is out of range.

Alarm messages	Default priority	Alarm clear method	Cause and solution
N ₂ O Out of Range	Low	Not clearable	The measured data of N_2O module is out of range.
AA Agent ID Are Unreliable	Low	Not clearable	Hardware or software error, contact the after-sales personnel.
AA At Least One Agent Outside Range	Low	Not clearable	Hardware or software error, contact the after-sales personnel.

D.2.11 DM Technical Alarm

Alarm messages	Default priority	Alarm clear method	Cause and solution
DM Finished	Low	Completely clear	The infusion container is empty and the infusion is complete.
DM Drip Speed	Low	Sound and light	During infusion, the drip rate
Abnormal		can be cleared	changes by more than 20%.

D.2.12 BIS Technical Alarm

Alanm massagas	Default	Alarm clear	Cause and solution
Alai iii iiiessages	Alarm messages priority	method	
BIS Sensor Off	Med	Completely clear	Check the connection status of sensors, re-connect the sensor if accessory. If the problem still exist, replace the sensor.
BIS Bad SQI	Med	Not clearable	SQI<15. 1. Check the patient's status. 2. Check whether the application sites of sensors are correct, and if the contact with the patient's skin is good.

Alarm massages	Default	Alarm clear	Cause and solution
Alarm messages	priority	method	
			3. Check if the BIS is away from
			RF device.
			SQI<50.
			1. Check the patient's status.
	Med		2. Check whether the application
DIG B GOI		Sound and light	sites of sensors are correct, and if
BIS Poor SQI		can be cleared	the contact with the patient's skin
			is good.
			3. Check if the BIS is away from
			RF device.
			The BIS simulator is connected
BIS Simulator	Low	Sound and light	and is for testing and
Connected		can be cleared	demonstration purposes only. Not
			for clinical use.

D.2.13 EEG Technical Alarm

Alarm mossages	Default	Alarm clear	Cause and solution
Alarm messages	priority	method	
EEG XXElectrode Off	Med	Completely clear	The electrode is not firmly attached to the patient or falls off causing the corresponding EEG lead off. Please check the connection of EEG electrode, lead wire and cable.
EEG XX Polarized	Med	Completely clear	Corresponding EEG electrodes is polarized. Please check the connection of the electrodes, re-connect the electrodes if necessary.

Alarm messages	Default priority	Alarm clear method	Cause and solution
EEG XX Impedance High	Med	Sound and light can be cleared	The connection of corresponding electrodes is poor. Check the connection of electrodes and re-connect the electrodes if necessary.
Note: XX represents EEG channel CH1, CH2, CH3, CH4.			

D.2.14 RM Technical Alarm

Alarm messages	Default priority	Alarm clear method	Cause and solution
RM Sensor Off	Med	Completely clear	Check the connection of sensor, re-connect the sensor.
RM Check Zero Failure	Low	Completely clear	Replug the module, if the problem still exists, contact the after-sales personnel.
RM Power Err	Low	Completely clear	Replug the module, if the problem still exists, contact the after-sales personnel.

D.2.15 rSO₂ Technical Alarm

Alaum massagas	Default	Alarm clear	Cause and solution
Alarm messages	priority	method	
rSO ₂ -1/rSO ₂ -2 Light Absorption Out of Range	Prompt	/	Sensor Off or ambient light is too strong. Replace the sensor to the patient's measurement site or adjust the ambient light intensity.
rSO ₂ -1/rSO ₂ -2 Noise	Prompt	/	Noise interference (e.g. electrosurgical unit). Check for any possible sources of signal

Alarm messages	Default priority	Alarm clear method	Cause and solution
			noise.
rSO ₂ -1/rSO ₂ -2 Unauthorized Sensor	Med	Not clearable	The sensor used is not authorized. Replace the sensor.
rSO ₂ -1/ rSO ₂ -2 No Cable Connected	Med	Sound and light can be cleared	The main cable is not connected, the connection is not firm or falls off. Please check the connection between the main cable and the rSO ₂ -1/rSO ₂ -2 module.
rSO ₂ -1/rSO ₂ -2 Incompatible Cable	Med	Not clearable	Use non-factory rSO ₂ -1/ rSO ₂ -2 cables. Please replace the original cable.
rSO ₂ -1/rSO ₂ -2 No Sensor	Med	Sound and light can be cleared	rSO ₂ sensor is detached, or the sensor cable is disconnected from the pre-amplifier. Re-connect the sensor to the pre-amplifier.
rSO ₂ -1/rSO ₂ -2 Sensor Fault	Med	Not clearable	Replug the module, if the problem still exists, contact the after-sales personnel.
rSO ₂ -1/rSO ₂ -2 Sensor Life Near Expiration	Prompt	/	SpO ₂ sensor is about to expire. Please replace the sensor in time.
rSO ₂ -1/rSO ₂ -2 Sensor Life Expired Sensor	Med	Not clearable	SpO_2 sensor has expired. Please replace the sensor in time.

D.2.16 NMT Technical Alarm

Alarm massages	Default	Alarm clear	Cause and solution
Alarm messages	priority	method	
NMT Cable Off	Low	Completely clear	The main cable is not connected, the connection is not firm or falls off. Please check the connection between the main cable and the module.
NMT Stimulus Electrode Off	Low	Completely clear	Check and confirm that the connection of the stimulation cable and main cable is normal. If the alarm still exists, detect the connection status of stimulation cable and patient.
NMT Low Battery	High	Not clearable	Check and replace the battery of NMT device.

D.2.17 System Alarm

Alarm messages	Default priority	Alarm clear method	Cause and solution
Battery Low	High	Not clearable	Please connect AC power supply for power supply and charge the battery.
Battery Fault	Prompt	/	Please replace the battery.
Recorder No Paper	Low	Sound and light can be cleared	The recorder is not loaded with paper or the recorder door is not closed. Please check the recorder to make sure the paper is loaded or the recorder door is closed.
Recorder Not Exist	Prompt	/	The recorder module is not plugged in. please insert the recorder module.
Recorder Too Hot	Low	Not clearable	The recorder works too long. Please

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Alarm massagas	Default	Alarm clear	Cause and solution
Alarm messages	priority	method	
			restart the recording task after the recorder head cools down.
CMS Disconnected	Med	Completely	The monitor is disconnected from the CMS. Please check the network connection.
Disk Full	Med	Not clearable	The storage space of the monitor is full. Please clear the patient related data in time.
Disk Will Be Full	Low	Sound and light can be cleared	The storage space of the monitor is almost full. Please clear the patient related data.

Appendix E Cybersecurity

This chapter mainly describes the information related to cybersecurity of the monitor.

E.1 Operating environment

- Hardware environment
 - Monitor software is only applicable to P series patient monitor hardware platform.
 - Screen:

P12: 12.1" LCD screen with 1280*800 pixels

P15: 15.6" LCD screen with 1920*1080 pixels

P18: 18.5" LCD screen with 1920*1080 pixels

P22: 22" LCD screen with 1920*1080 pixels

- Peripherals: nurse call module, recorder.
- ◆ Software environment
 - Main board: P12MB
 - > Operating system: LinuxLinux-3.2.0 kernel + Busybox filesystem.
 - Database: sqlite-3.16.2
- Network environment.
 - Apply to LAN

E.2 Network data interface

The communication interface between the monitor and the CMS is wired or wireless Ethernet, using the standard TCP/IP protocol family, and the application layer data format follows *the Central Monitoring System Network Communication Protocol* during transmission.

E.3 User access control mechanism

- a) User identification method: after entering the authorization password, you have the corresponding user type setting authority.
- b) User types: medical personnel, hospital equipment maintenance personnel, factory maintenance personnel.

c) User authority:

- 1) Authority of medical staff: No password. Automatically enter the monitoring interface after starting up, and can be routinely set as required.
- 2) Authority of hospital equipment maintenance personnel: Enter the maintenance menu by entering the hospital maintenance password, and at least have settings for language configuration, automatic clearing of NIBP results, automatic release of waveform freezing time and alarm related contents.
- 3) Manufacturer's authority: Enter the maintenance menu by entering the manufacturer's maintenance password. In addition to the contents that can be set by the authority of hospital equipment maintenance personnel, the manufacturer can at least set the power frequency and module configuration.

E.4 Software Environment

The list of system software is as follows:

Software name	Version
Linux	V3.2.0

◆ The supporting software is as follows:

Software name	Version
Sqlit3	V3.16.2

◆ The list of application software is as follows:

Software name	Supplier
P series monitor	Guangdong Biolight Meditech Co., Ltd.
software	

Appendix F Terminology and Definitions

F.1 List of units

Abbreviation	Full name
μΑ	microampere
μV	microvolt
μs	microsecond
A	ampere
Ah	ampere hour
bpm	beat per minute
bps	bit per second
°C	centigrade
cc	cubic centimeter
cm	centimeter
dB	decibel
DS	dyne second
°F	fahrenheit
g	gram
GHz	gigahertz
GTT	gutta
h	hour
Hz	hertz
in	inch
k	kilo
kg	kilogram
kPa	kilopascal
L	litre
lb	pound
m	meter
mAh	milliampere hour
Mb	mega byte

Abbreviation	Full name
mcg	microgram
mEq	milli-equivalents
mg	milligram
min	minute
mL	milliliter
mm	millimeter
mmHg	millimetes of mercury
cmH ₂ O	centimeters of water
ms	millisecond
mV	millivolt
mW	milliwatt
MΩ	megaohm
nm	nanometer
rpm	breaths per minute
S	second
V	volt
VA	volt ampere
Ω	ohm
W	watt

F.2 Symbol list

Symbol	Explanation
_	Minus
-	Negative
%	Percent
/	Per; Divide;Or
~	То
+	Plus
=	Equal to
<	Less than

Symbol	Explanation
>	Greater than
<u>≤</u>	Less than or equal to
<u> </u>	Greater than or equal to
±	Plus or minus
×	Multiply
©	Copyright

F.3 Terminology list

Abbreviation	Full name			
AAMI	Association for Advancement of Medical Instrumentation			
AC	Alternating current			
Adu	Adult			
AHA	American Heart Association			
ANSI	American National Standard Institute			
Ao	Aortic pressure			
Art	arterial			
ATMP	borometric pressure			
AUC	Area under the curve			
aVF	Left foot augmented lead			
aVL	Left arm augmented lead			
aVR	Right armaugmented lead			
awRR	Airway respiratory rate			
BAP	Brachial arterial pressure			
BC	Burst count			
BL	Baseline			
BIS	Bispectral index			
BP	Blood pressure			
BSA	Body surface area			
BT	Blood temperature			

Abbreviation	Full name			
CaO ₂	Arterial oxygen content			
CE	Conformité Européenne			
C.I.	Cardiac index			
CIS	Clinical information system			
CISPR	International Special Committee on Radio Interference			
CMOS	Complementary metal oxide semiconductor			
CMS	Central monitoring system			
C.O.	Cardiac output			
CO ₂	Carbon dioxide			
СОНЬ	Carboxyhemoglobin			
Compl	Compliance			
CVP	Central venous pressure			
DC	Direct current			
Des	Desflurane			
Dia	Diastolic			
DVI	Digital video interface			
DO_2	Oxygen delivery			
ECG	Electrocardiograph			
EDV	End-diastolic volume			
EEC	European Economic Community			
EMC	Electromagnetic compatibility			
EMG	Electromyograph			
EMI	Electromagnetic interference			
Enf	enflurane			
ESU	Electrosurgical unit			
Et	End-tidal			
EtCO ₂	End-tidal carbon dioxide			
FAP	Femoral arterial pressure			
FCC	Federal Communication Commission			

Abbreviation	Full name			
FDA	Food and Drug Administration			
FiCO ₂	Fraction of inspired carbon oxygen			
Flow	Flow			
FV	Flow-volume			
Hal	halothane			
Нь	Hemoglobin			
Hb-CO	Carbonmono-xidehemoglobin			
HbO ₂	Oxyhemoglobin			
HIS	Hospital information system			
HR	Heart rate			
IBP	Invasive blood pressure			
ICG	Impedance cardiography			
ICP	Intracranial pressure			
ICU	Intensive care unit			
ID	Identification			
I:E	Inspiatory time: expiratory time ratio			
IEC	International Electrotechnical Commission			
IEEE	Institute of Electrical and Electronic Engineers			
IP	Internet protocol			
Iso	isoflurane			
LA	Left arm			
LAP	Left atrial pressure			
Lat	Lateral			
LCD	Liquid crystal display			
LCW	Left cardiac work			
LCWI	Left cardiac work index			
LED	Light emitting diode			
LL	Left leg			
LVDS	Low voltage differential signal			

Abbreviation	Full name			
LVSW	Left ventricular stroke work			
LVSWI	Left ventricular stroke work index			
MAC	Minimum alveolar concentration			
MAP	Mean arterial pressure			
MDD	MedicalDeviceDirective			
MetHb	Methemoglobin			
MRI	Magnetic resonance imaging			
N/A	Not applied			
N_2	Nitrogen			
N ₂ O	Nitrous oxide			
Neo	Neonate			
NIBP	Noninvasive blood pressure			
O ₂	Oxygen			
OR	Operating room			
oxyCRG	Oxygencardio-respirogram			
PA	Pulmonary artery			
PAWP	Pulmonary artery wedge pressure			
Paw	Airway pressure			
PD	Photodetector			
Ped	Pediatric			
PEEP	Positive end expiratory pressure			
PEF	Peak expiratory flow			
PIF	Peak inspiratory flow			
PIP	Peak inspiratory pressure			
Pleth	Plethysmogram			
PO ₂	Oxygen supply pressure			
PPV	Pulse pressure variation			
PR	Pulse rate			
PVC	Premature ventricular contraction			

Abbreviation	Full name			
R	Right			
RA	Right arm			
RAM	Random access memory			
RAP	Right atrial pressure			
Raw	Airway resistance			
Rec	Record,recording			
RESP	Respiration			
RHb	Reduced hemoglobin			
RL	Right leg			
RM	Respiratory mechanics			
RR	Respiration rate			
RSBI	Rapid shallow breathing index			
SaO ₂	Arterial oxygen saturation			
SEF	Spectral edge frequency			
Sev	sevoflurane			
SFM	Self-maintenance			
SI	Stroke index			
SpO_2	Arterial oxygen saturation from pulse oximetry			
SQI	Signal quality index			
STR	Systolic time ratio			
SV	Stroke volume			
SVI	Stroke volume index			
SvO ₂	Venous oxygen saturation			
Sync	Synchronization			
Sys	Systolic pressure			
Taxil	Axillary temperature			
ТВ	Blood Temperature			
TD	Temperature difference			
TEMP	Temperature			

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Abbreviation	Full name				
TFC	Thoracic fluid content				
TFI	Thoracic fluid index				
TFT	Thin-film technology				
TP	Total power				
Trect	Rectal temperature				
TVe	Expiratory tidal volume				
TVi	Inspiratory tidal volume				
UAP	Uninterruptible power pressure				
UPS	Uninterruptible power supply				
USB	Universal serial bus				
UVP	Umbilical venous pressure				
VAC	Volts alternating current				
VEPT	Volume of electrically participating tissue				
VI	Velocity index				

Appendix G Toxic and Harmful Substances or elements

Components		Lead	Mercury	Cadmiu	Hexavalent	Polybrominat	Polybrominat
		Pb	Hg	m	chromium	ed biphenyls	ed diphenyl
				Cd	Cr(VI)	PBB	ethers
							PBDE
Host	Shell	0	0	0	0	0	0
	(plastic						
	parts)						
	Label	0	0	0	0	0	0
	Internal	0	0	0	0	0	0
	sheet						
	metal						
	EMI	0	0	0	0	0	0
	Gasket						
	Silicone	0	0	0	0	0	0
	Piece						
Package	Package	0	0	0	0	0	0
	materials						
General	Adapting	0	0	0	0	0	0
	piece						
	Power	0	0	0	0	0	0
	cord						
Battery	Lithium	0	0	0	0	0	0
	battery						
Accessor	ECG	0	0	0	0	0	0
у	accessory						
	SpO_2	0	0	0	0	0	0
	accessory						
	TEMP	0	0	0	0	0	0
	accessory						
	NIBP	0	0	0	0	0	0
	accessory						
	IBP	0	0	0	0	0	0
	accessory						
	ICG	0	0	0	0	0	0

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Compon	ents	Lead Pb	Mercury Hg	Cadmiu m Cd	Hexavalent chromium Cr(VI)	Polybrominat ed biphenyls PBB	Polybrominat ed diphenyl ethers PBDE
	accessory						
	CO ₂	0	0	0	0	0	0
	accessory						
	AG	0	0	0	0	0	0
	accessory						
	RM	0	0	0	0	0	0
	accessory						
	EEG	0	0	0	0	0	0
	accessory						
	NMT	0	0	0	0	0	0
	accessory						
	BIS	0	0	0	0	0	0
	accessory						
	rSO ₂	0	0	0	0	0	0
	accessory						
Stand	Carts stand	0	0	0	0	0	0
	Wall stand	0	0	0	0	0	0

o: It means that the content of the toxic and harmful substances in all homogeneous materials of the component is below the limit specified in SJ/T11363-2006.

^{×:} Indicates that the content of the toxic and harmful substances in at least one homogeneous material of the component exceeds the limit requirements specified in SJ/T11363-2006.

Product name: Patient Monitor

Product model: P12/P15/P18/P22

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