KC7 Series Multi parameter module Patient Monitor Instruction for Use

MDKMed Medical Technology Co., Ltd.

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Product Information

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The registered trademark used on the product is:佳士比医疗

Product name: Multi parameter Patient Monitor

Specifications and models:

This IFU is the KC7 series product manual, applicable to the following models:

KC7: KC7A, KC7AC.

Registered address: 502A, Building 7, No. 22, Xinyan Road, Donghu Street, Linping District, Hangzhou City, Zhejiang Province, P. R China 311323

Production address: No.22, Cangling Rd, Huzhen Town, Jinyun County, Lishui City.Zhejiang Province, P. R. China 321404

Production date/expiration date: as per label

Structural composition: The product consists of a main unit, a power adapter, and accessories. Accessories include ECG lead wires, temperature sensor, blood oxygen probe, blood pressure cuff and external CO2 module (optional).

Our company has passed ISO13485 product quality system certification.

Statement

This manual applies to our company's monitors, and the content may not be changed without consent. Our company reserves the right to improve products in terms of technology, parts, software and hardware. If you need further information about the product, you can contact our company or dealer.

Copyright

Our company owns the copyright of this non-publicly published manual and treats it as confidential information. This manual is only used as a reference for the use, maintenance and repair of this product. Others have no right to disclose the contents of this manual to others.

This manual contains proprietary material protected by copyright laws. All rights reserved; No part of this manual may be photographed, copied, or translated into other languages without the written consent of our company.

Our Company does not make any form of warranty for this information, including (but not limited to) the implied warranties of merchantability and fitness for a particular purpose. Our Company is not responsible for errors contained in this information or for incidental or consequential damages resulting from the provision, actual performance, and use of this manual.

The contents contained in this manual may be changed without notice.

Manufacturer's Responsibilities

Our Company considers itself responsible for the safety, reliability and performance of our products only under the following circumstances, namely:

(1) Assembly operations, expansion, readjustment, improvements and repairs are all performed by personnel approved by the company;

(2) Relevant electrical equipment complies with national standards;

(3) The product should be used according to the operating instructions.

Our company does not make any form of guarantee for errors in this manual, installation errors, or operation errors, and does not assume any legal responsibility for incidental or inevitable damage.

Motice

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- •This product is not for home use.
- •This product is restricted to one patient at a time.

•According to the requirements of YY9706.261-2023 (YY 0784-2010), claims for blood oxygen saturation accuracy should be supported by clinical research measurements covering the entire range. The functional tester cannot be used to evaluate the accuracy of pulse oximeter probes and pulse oximeter monitors.

•Expired products or batteries should be disposed of in accordance with local electronic product scrapping regulations or returned to the manufacturer for recycling.

WARNING

•For safe and continuous use of this equipment, please follow the instructions below. The instructions listed in this leaflet do not replace medical procedures already being performed.

•You cannot rely solely on audible alarm systems to monitor patients. When monitoring a patient, if the volume is set too low or is completely turned off, the safety of the patient may be endangered. The most reliable method of patient monitoring combines the proper use of monitoring equipment with close personal monitoring of the patient.

•This device is intended for use in healthcare facilities by trained caregivers only.

•To reduce the risk of electric shock, do not disassemble the equipment. If necessary, ask professionals to perform repairs.

•This device may interfere with the ultrasound imaging system, such as interference signals appearing on the ultrasound display. Place the two devices as far apart as possible.

•Exposing electrical contacts or connecting devices to saline or other liquids and conductive glue is extremely dangerous. Electrical contacts and other connecting devices such as cable connectors, power supplies, parameter module plug-in connectors, rack connectors, etc. must be kept clean and dry. If they are contaminated with liquid, they must be dried thoroughly. If further decontamination is required, please contact your medical equipment department or our company.

•KC7 monitor can not be used in ambulances.

▲ WARNING

•This product is not a therapeutic device.

•If hospitals or medical institutions responsible for maintaining this product fail to maintain this product as planned, it may cause the product to fail and ultimately endanger the patient's health.

Quality Assurance

Free service scope:

All equipment that meets the scope of our company's warranty service regulations can

enjoy free service.

Paid service scope:

(1) For equipment that exceeds the scope of the company's warranty service regulations,

the company will provide paid services;

(2) Even within the warranty period, the product needs repair due to the following reasons: man-made damage; the network power supply voltage exceeds the rated range

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of the equipment; irresistible natural disasters.

Our company is not responsible for direct, indirect or final damages and delays caused by the following situations (including but not limited to): Components are disassembled, stretched, and re-adjusted; Replace accessories not approved by our company or have the machine repaired by personnel not authorized by the company.

Goods Return

If you need to return a product, please follow these steps:

Obtain the right of return. Contact our company's customer service department and inform them of the product series number. If the series number is not clearly legible, returns will not be accepted. Please indicate the product model and serial number, and briefly describe the reason for return.

Freight: The product is shipped to our company for repair, and the user must bear the freight (including customs fees).

After-sales service

When you need product technical service support, please contact agents and dealers, or contact the nearest office of our company. Also, you could contact our company's service department, please see the last page of this manual for detailed information.

Please make sure to record the following information before you make a call:

- 1. Product model and factory number;
- 2. Production date;
- 3. Product validity period;
- 4. Malfunction details

Introduction

This manual introduces the performance, operation methods and other safety information of the monitor in detail. Reading this manual is the first step for users to familiarize themselves with and make full use of the product, ensuring proper use and ensuring patient and operator safety.

The following symbols indicate certain important reminders that the user should pay attention to:

 \triangle **Danger:** Indicates an imminent hazard which, if not avoided, may result in death, serious personal injury or property damage.

WARNING: Indicates a potentially hazardous or unsafe operation that, if not avoided, may result in death or serious personal injury or property damage.

Careful: Indicates potentially dangerous or unsafe operations which, if not avoided, may result in minor personal injury or product failure, damage or property loss.

 \triangle Notice: Highlight important precautions, provide instructions or explanations for better use of this product.

This manual is intended for people who are familiar with various measurements and need to use monitoring equipment.

This monitor is a multi-parameter monitor that can be used to monitor the vital signs of adults and children during surgery, surgery/anesthesia recovery, emergency rooms, etc. This monitor can be powered by the built-in battery or AC power. Easy to carry and transport.

Scope of application:

This monitor is used to measure and monitor the patient's ECG, respiration, non-invasive blood pressure, pulse oximeter, pulse rate and body temperature, and end-tidal CO_2 parameters.

Intended users:

Patients who require monitoring of electrocardiogram, respiration, non-invasive blood pressure, pulse oxygen saturation, pulse rate, body temperature, and CO₂ parameters. This device is not suitable for newborns.

Contraindication

Patients with sickle cell disease and anyone with skin lesions or expected lesions; For patients with severe coagulation disorders or severe thrombosis;

Patients with intravenous fluids or catheters in their limbs;

Patients with perfusion disorders or skin sensitivities.

∕∕∕Danger

•Do not use this device in an environment rich in oxygen or containing flammable or explosive items such as anesthetics to prevent fire or explosion. At the same time, the product and surrounding areas should be kept clean and dry.

•Do not touch the patient during defibrillation, or keep a sufficient distance from metal objects connected to the patient. Failure to do so may result in serious injury or death.

AWARNING

•Product only plays a supporting role in diagnosis, doctors made diagnosis in conjuctions with patient's clinical situations.

•If this equipment is not secured properly, it may fall, resulting in personal injury or equipment damage. To prevent personal injury or equipment damage, install the equipment in a fixed location.

•Before use, you must check the equipment, connecting cables and supporting accessories to ensure that they can work normally and safely. The equipment should self-check normally when it is turned on.

• This equipment cannot be used in the presence of magnetic resonance (MR) equipment, otherwise the induced current may cause burns to the patient.

•This equipment cannot be used in places with excessive electromagnetic radiation or EMC equipment, otherwise the measurement accuracy will be affected.

•To avoid personal injury, the equipment cannot be repaired by anyone except professional technicians.

•Do not replace the power cord of this device. Do not connect the three-core power cord of this device to a two-core or multi-hole socket without protective grounding. If the reliability of the grounding cannot be confirmed, please use the internal battery for power supply.

Motice

•Prior to use, calibration should be properly verified and the equipment operating properly.

•Please ensure that the battery has been charged so that the device can still work normally when the network power is cut off for more than 30 seconds.

•Pay attention to the placement of power cords, conduits, and all electrical cables so that

they will not strangle the patient or trip others.

•The back of this device must not be blocked to facilitate heat dissipation.

•If liquid is spilled into the casing of this equipment, please disconnect the power supply immediately and contact service personnel.

•The product should be stored in an environment with temperature ($5^{\circ}C$ ~40°C), humidity (15° ~80%, no condensation), and atmospheric pressure (86.0kPa~106.0kPa). If the product has been stored for more than one year, it must be re-tested according to the factory inspection procedures before it can be used again. For related accessories with storage life requirements (such as ECG electrodes, batteries, etc.), follow their respective instructions.

•To avoid contaminating or infecting people, the environment, or other equipment, equipment and accessories that have reached the end of their useful life must be disposed of in accordance with relevant local regulations or hospital policies.

•Equipment should be installed in a location that is easy to observe, operate and maintain.

WARNING

•The multi-parameter monitor is used for clinical patient monitoring. Only doctors and nurses are allowed to use this monitor.

•Do not open the product's casing to avoid possible electric shock hazard. Any repairs and upgrades to the monitor must be performed by service personnel trained and authorized by our company.

•Do not use this product where flammable items such as anesthetics are placed to prevent an explosion.

•Before use, users should check whether this product and its accessories can work properly and safely.

•To prevent delays in treatment, set adequate alarm settings for each patient. At the same time, it should be ensured that the alarm sound can be heard when alarming.

•Do not use mobile phones near the monitor. Mobile phones can produce excessively strong radiation fields that can interfere with monitor functionality.

•During defibrillation, keep away from the patient, table, and product.

•The equipment interconnected with the monitor should form an equipotential body (effective connection to the protective ground).

•When the monitor is shared with electrosurgical equipment, the user (doctor or nurse)

should pay attention to ensuring the safety of the patient being monitored.

•Packaging must be disposed of in accordance with current waste control practices and kept out of the reach of children.

•For patients with a pacemaker, the heart rate monitor may include pacemaker pulses in the heart rate value during cardiac arrest or arrhythmia. Don't rely entirely on heart rate monitor alarms. Patients with pacemakers should be closely monitored and the device's ability to suppress pacing pulses should be discussed elsewhere in this manual.

ACareful

•When the products described in this manual are about to expire, they must be disposed of in accordance with the relevant product handling regulations. If you would like further information, please contact our company or its agents.

•When you have doubts about the completeness and arrangement of the external grounding of the monitor, you must use device's internal battery for operation.

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Description of equipment symbol

Â	Warning	\sim	Alternating current	-1h-	ECG
-+	Battery		Protective ground (earth)	\checkmark	Equipotential
€-	NIBP		computer network	I, I	Lead selection
1	Unlock	\sim	graph recorder		Input /Output
\square	gas output	€	USB interface		gas input
X	Temperature limit	$[m]{}$	Manufacturing date	SN	Serial number
$(((\cdot, \cdot)))$	Non-ionizing radiation	4	Dangerous electric shock	ᠿ	Video output
	Manufacturing measuring instrument license mark	<u>ک</u>	Humidity limits	Ģ	Air pressure limit
00	Power on and off	┦₩₽	The application part of defibrillation prevention CF type	1 1	The application part of defibrillation prevention BF type
\bowtie	Alarm off	X	Alarm pause	.X.	Alarm reset
×	Alarm SILENCE		Follow the operating instructions		Universal warning symbols (safety symbols)
?	Signal incompletenes s indicator				

Chapter 1 Product Description

- For a complete understanding of the monitor, read this overview
- For an introduction to the various information displayed on the screen, read Screen Display Introduction
- To master the operation method, please read the monitor button functions and basic operations.
- To learn the location of the various interfaces, please read Monitor External Interfaces
- To learn about the precautions for using the monitor on battery power, read Built-in Rechargeable Battery

1.1 Basic Information Introduction

Structural features and working principles

The Multi Parameter Patient Monitor controls multiple parameter function modules through a CPU, and forms a monitoring system with physiological parameter measurement modules such as ECG, RESP, NIBP, SPO₂, TEMP, PR, and CO₂. It can monitor physiological parameters such as ECG, RESP, TEMP, NIBP, SPO₂, PR, and CO₂ for adult and pediatric patients. Through serial port communication with the host computer, 7-lead ECG waveform, 1 SPO₂ waveform, 1 RESP waveform, and 1 CO₂ waveform data can be provided in real time. Through analysis and calculation, physiological parameters such as HR, RR, SPO₂, PR, CO₂, non-invasive blood pressure (systolic blood pressure, diastolic blood pressure, mean blood pressure) can be provided. The product consists of a main unit, power adapter, external CO₂ module (optional) and accessories. Accessories include ECG lead wires, temperature sensors, blood oxygen probes, and blood pressure cuffs.

Working Environment:

Temperature: Working temperature 5 \sim 40 (°C), transportation and storage temperature -20 \sim 55(°C);

Humidity: Working humidity $15\% \sim 85$, no condensation, transportation and storage humidity $\leq 93\%$;

Atmospheric pressure: 86.0kPa~106.0kPa;

Power supply voltage: a.c.100V~a.c.240V 50Hz/60Hz, d.c.12V input;

Internal battery specifications: 7.4V/4800mAh.

 \triangle Notice: Do not use this monitor outside the temperature and humidity range specified by the manufacturer, otherwise it will not meet the performance

specifications claimed in Appendix II.

Abbreviation definition:

Name	Definition, Abbreviation	Name	Definition, Abbreviation
ECG	Electrocardiogram	HR	Heart rate
RESP	Breathe	RR	Respiratory rate
TEMP	Body temperature	TD	Temperature difference
SPO2	Pulse oximetry	SaO2	Blood oxygen saturation
Pleth	Plethysmography	PI	Perfusion index
NIBP	Non-invasive blood pressure	SYS	Systolic blood pressure
MAP	Mean arterial pressure	DIA	Diastolic blood pressure
CO2	Carbon dioxide	EtCO2	End-tidal carbon dioxide content
FiCO2	Inhaled carbon dioxide content	awRR	Airway respiratory rate (breaths per minute)
INS	Inspiratory air minimum CO2	InsCO2	Minimum amount of carbon dioxide inhaled
IBP	Invasive blood pressure	CVP	Central venous pressure
RAP	Right atrial pressure	LAP	Left atrial pressure
ICP	Intracranial pressure	ART	Arterial pressure
РА	Pulmonary artery pressure	PR	Pulse rate
BIS	EEG bispectral index		

1.2 Introduction to display interface

The display screen of this monitor is a color LCD screen, which can simultaneously display the collected patient parameters, waveform parameters, alarm information, bed position, monitor status, clock and other prompt information provided by the monitor. The main screen is divided into four areas (as shown in Figure 1-1):



Information		Parameter Area	Shortcut Are
Area	22:46 Bed 1 Adult ECG Leads Off	*** CO2 Apres	<u>چ</u>
•	xt Monitor 25.0em/s	ECG bps	 More
Water Area		50 NIBP 00:00 Mercual mailing	ロ) Silence
Wave Alea	Lav		QC NIBP
	7 Iao	TEMP *C	E Switch
		Sp02	Pa Patient
		C02 Ficoz O miłło	Layout
	602	50 V .	(2) Setup

Figure 1-1 Main Display Interface

Shortcut area button icon:

More	Patient information
Mute	Interface layout
Blood pressure measurement	Setting menu
Interface selection	

Information area introduction:

The information area is at the top of the screen and displays the current status of the monitor and patient.

Examples of the content of the information area are as follows: "Bed": refers to the bed number of the patient being monitored. "Adult": refers to the type of patient being monitored. "11:01": refers to the current time.

Other prompt messages in the information area appear and disappear at the same time as the reported status, and are divided into:

(1) The monitor prompts information to report the status of the monitor or sensor, and the area behind "Adult" appears fixedly;

(2) Monitor alarm information (see the "Alarm" chapter for specific setting methods).

•When the waveform on the screen is frozen, the corresponding prompt "Freeze"

window appears at the bottom of the monitor screen.

•Patient parameter alarm information always appears in the rightmost area.

(3) "----" indicates that the monitored parameters are in an invalid state.

Introduction to waveform/menu area:

The waveform area displays 5 waves, and the order of waveform display can be adjusted.

Under the maximum configuration, the system can display 7 ECG waves, SPO2

plethysmography waves, respiratory waves, etc. in the waveform area.

The name of the waveform is displayed on the upper right side of each waveform. ECG leads can be selected upon request. Each ECG wave also displays the gain of the channel

and the ECG filtering method. There is a 1mV scale on the left side of the ECG waveform.

When a menu pops up during screen operation, the menu always occupies a fixed position on the left side of the waveform area, making part of the waveform temporarily invisible. After exiting the menu, the original screen display is restored.

The waveform is refreshed at the set rate. To adjust the refresh rate of each waveform, see the settings of each parameter.

Introduction to parameter area:

The parameter area is on the right side of the waveform area and is basically placed corresponding to the waveform. The parameters displayed in the parameter area are:

(1) ECG - heart rate or pulse rate (unit: beats/minute)

(2) Blood oxygen saturation SPO₂ - blood oxygen saturation SPO₂ (unit: %), pulse rate (unit: beats/minute) (when the heart rate source selects the "simultaneous" option)

(3) Non-invasive blood pressure NIBP - from left to right are systolic blood pressure, mean blood pressure, and diastolic blood pressure (unit: mmHg or kPa)

(4) Body temperature TEMP - temperature (unit: Celsius ° C or Fahrenheit °F)

(5) Respiratory RESP - respiratory rate (unit: times/minute)

(6) End-tidal carbon dioxide CO₂ - carbon dioxide concentration (unit: mmHg or kPa)

Alarm lights and alarm status:

In normal state, the alarm light does not light up. When an alarm occurs, the alarm light flashes or stays on. The color of the light represents a certain alarm level. For details, please refer to the "Alarm" chapter.

For the specific content of alarm information and prompt information, please refer to the relevant descriptions of each parameter in the relevant chapters.

1.3 Front view

Monitor power switch" is on the front panel of the monitor. The AC indicator light "AC" is on the lower left side of the power switch with the identifier """. When the product is powered by AC power, this light turns on green, and the battery indicator light is on the right side of the AC indicator light "AC" with the symbol """, when the monitor uses the internal battery to work, this light does not light up; when the product is equipped with a battery and is in the charging state, this light stays green; when the battery is fully charged, this light goes out. The alarm light is on the whole machine on

the upper left and when an alarm occurs, this light flashes. The sensor jack is on the left side of the front panel of the product. The recorder is on the right side of the machine. Other jacks and power sockets are on the rear panel.

This monitor has a user-friendly operation interface, and all operations can be completed through the buttons and knobs on the front panel. For details, please refer to the system menu section.



Function description:

1. Alarm indication area 2. Parameter and waveform display area 3. Function key area

4. Measurement cable connection area 5. Shortcut function area

Chapter 2 Monitor Installation

2.1 Key functions and basic operations

Operations on the monitor can be completed through buttons and knobs.



Press and hold for 2 seconds to power on or off



Start or stop blood pressure measurement.

2.2 Monitor external interface



Figure 2-1 Monitor parameter interface

Rear panel jack description:

The upper part is the data transmission interface, and the lower part is the power interface.



AWARNING

•All analog and digital devices connected to this monitor must be products certified by specified IEC standards (such as IEC 60950 data processing equipment standard and IEC 60601-1 medical equipment standard). And all configurations should comply with the contents of the valid version of the IEC 60601-1-1 system standard. The person responsible for connecting additional equipment to the input/output signal ports configures the medical system and is responsible for the system's compliance with IEC 60601-1-1 standards. If in doubt, please contact your supplier.

2.3 Built-in rechargeable battery

KC7 built-in 7.4V/4800mAh Lithium-ion battery, not removable.

When working on battery power, the monitor will alarm when the battery is low. When the battery is exhausted, the monitor will trigger a high-level alarm, make a continuous "beep..." sound, and prompt "Battery voltage is too low" in the information area. At this time, the DC power supply should be plugged in to charge the battery immediately. If it is still powered by batteries, the monitor will automatically power off before running out of power (approximately 10 minutes after the alarm).

Battery and AC status light indication: When the network power supply is powered, the AC indicator light is on, the battery light is on during the charging process, and when the battery is fully charged, the battery indicator light is off; when the battery is powering the device, the AC indicator light is off and the battery indicator light is flashing. The warranty period of the battery is 1 year, and it needs to be charged and discharged

every 3 months, otherwise the battery life will be seriously shortened. Often due to manufacturer inventory - logistics inventory - dealer inventory - customer inventory, the battery is often damaged or life performance is reduced due to this reason. If the monitor will not be used for a long time, then the battery need to be removed. If not used for a long time > 3 months, the battery should be removed from the monitor.

2.4 Unpack and inspect

Carefully remove the product from the packaging box. If any damage is found, please contact the shipping company immediately and save the packaging materials for future transportation or storage. Please check the accessories according to the packing list.

(1) Check for any mechanical damage.

(2) Check all exposed wires.

When installing, leave at least 2 inches (5 cm) of space around the monitor to allow air circulation. The environment in which the monitor is used must be reasonably free from vibration, dust, corrosive or explosive gases, extreme temperatures and moisture, etc.

If you have any questions, please contact our sales department or agent immediately.

ANotice

•When equipment is moved from one environment to another, condensation may occur on the equipment due to differences in temperature or humidity. At this time, you must wait for the condensation to disappear before use.

WARNING

•Packaging materials may cause pollution to the environment. When disposing of packaging materials, you must comply with relevant local regulations or the hospital's waste disposal system. Please keep packaging materials out of the reach of children.

•The equipment may be contaminated by microorganisms during storage, transportation and use. Please confirm whether the packaging is intact before use, especially single-use accessories. If damage is found, please do not use it.

2.5 Power connection

Steps to connect the DC power cord:

(1) Make sure the AC power supply meets the following specifications: a.c.100-240V, 50/60Hz.

(2) Use the adapter provided with the monitor. Plug the adapter into a power outlet. The DC output end of the adapter is plugged into the DC socket of the host computer. \triangle Notice

•The battery must be charged after the product has been transported or stored. If you turn on the product without connecting it to AC power, the product may not work properly due to insufficient battery power. Plug in AC power and charge the battery whether the monitor is turned on or off.

•Please ensure that this equipment works under the specified environmental requirements, otherwise it will not be able to meet the technical specifications claimed in this manual, and may cause equipment damage and other unpredictable consequences.

•When the power supply of this monitor network is cut off for more than 30 seconds, the product switches to the internal battery power supply. All alarm limit settings and other related parameter settings are the values before the power outage.

• This monitor has no power-off time record. Power outage does not affect real-time operation, and logs are recorded according to time without change.

2.6 Power on and start up

Press and hold the power button for 3 seconds. At this time, the alarm light flashes red once. After 2 seconds of powering on, the company logo or model number will be displayed for about 9 seconds. Enter the monitoring screen. Touch anywhere on the display to pop up the touch screen unlock switch. Click "Yes" The touch screen is unlocked and the user can perform operations.

WARNING

•If you find signs of damage to the monitor function, or if an error self-test prompt appears, do not use this equipment to monitor patients, and please contact the hospital's medical equipment engineer or our company's maintenance engineer.

Motice

•Check all available monitoring functions to make sure the monitor is functioning properly.

•If equipped with a battery, the battery must be charged after each use to ensure sufficient power reserve.

•Wait 1 minute after shutting down before turning it on again.

2.7 Sensor connection

Connect the required sensors to the monitor and patient monitoring site.

For the correct connection methods and related requirements of various sensors, please refer to the relevant chapters.

2.8 Start working

1. Confirm which monitoring mode is required.

2. Check whether the patient cable and sensor are correct.

3. Connect the patient cable and sensor.

4. Enter the required working mode and check whether the various settings and alarm parameters meet the current needs.

For other parameters or system settings, please refer to the relevant chapters.

2.9 End of work

Press and hold the power button for 2 seconds to shut down. If using power cable, the power cord should also be unplugged. Before turning off your monitor, perform the following checks:

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- 1. Ensure patient monitoring has been completed.
- 2. Disconnect the cables and sensors from the patient.
- 3. Make sure to save or clear patient monitoring data as needed.

2.10 Disconnecting the network power method

To unplug the adapter from the network power supply.

Chapter 3 System Menu

The configuration of this monitor is flexible, and the monitoring content, waveform scanning speed, etc. can be configured by the user according to needs. Press the "Menu" key on the front panel, the menu shown in Figure 3-1 will pop up, and you can perform the following operations:

Main X
Patient Information >>
Night Mode Setup >>
Monitor Setup >>
Alarm Setup >>
Maintenance >>
Review >>
Drug Calc >>

Image 3-1 Main menu

3.1 Patient information management

Select "Patient Management" in the main menu, and the menu shown in Figure 3-2 will pop up.



Figure 3-2 Patient Management Figure 3-3 Confirm to update patient data

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Project	Illustrate
Bed no.	1-200 beds optional
Gender	Patient gender
Detiont	Adults, children (with pacemaker (this monitor does not provide this
Fatient	function))
	In this menu, the user can select the "Update Patient" item to pop up the
	"Confirm Update Patient" dialog box to determine whether to clear the
Undata nationt	data. As shown in Figure 3-3.
Opdate patient	Select: "Yes" to delete all information of the currently monitored patient
	and exit the menu.
	Select: "No" to continue saving patient information and exit the menu.

ANotice: If you select "Yes", all information of the currently monitored patient will be deleted.

3.2 Default configuration

In the maintenance menu, users can set the current system configuration to the manufacturer's default configuration. At this time, the system will automatically save all parameter menu settings, ECG leads, gain, and filtering according to the patient type as the corresponding manufacturer's default configuration content, and a dialog box will pop up as shown in Figure 3-4:

Select "Yes" to save all configurations of the current patient type as user default configurations.

Select "No" to abandon the current operation, and the system will keep the original configuration content unchanged.



Figure 3-4 Default configuration menu

ANotice

•After selecting any item in the "Default Configuration" menu and exiting, the "Confirm Default Configuration" dialog box will pop up. The user can select "Yes" to confirm the selection, or "No" to abandon the selection.



WARNING

•At this time, all configurations in the system will be replaced by "default

configuration".

3.3 Review function

After selecting the "Review Function" option in the "Main Menu", the menu as shown in Figure 3-5 will appear:

Review X	NIBP Revie	W			X
NIBP Review >>	1	SYS 120	MAP 90	DIA 80	Interval Time 04-24-2018 23:14:11
Alarm Event Review >>	2	120	90	80	04-24-2018 23:07:18
Trend Graph >>					
Trend Table >>					
₩ave Recall >>	Number 2				*

Figure 3-5 Information review menu image Figure 3-6 NIBP measurement review When the "NIBP Measurement Review" option in "Information Review" is selected, the menu shown in Figure 3-6 will appear:

3.3.1 NIBP review:

The monitor can display the last 1000 NIBP measurement data in the NIBP review. Select the "Review" function in the "Menu" and then select the "NIBP Measurement Review" item. The last 10 NIBP measurement results and measurement times will be displayed in the window. As shown in Figure 3-6 above.

Data are organized chronologically from near to far. Each screen can display 10 times of measurement data. Select "page forward and backward" to view later or earlier data. Up to 1000 measurement results can be displayed. When the number of measurements exceeds 1,000, the latest 1,000 data will be displayed.

3.3.2 Event alarm review

You can review the alarm parameters and waveforms corresponding to the 200 groups of alarms that occurred recently.

3.3.3 Trend chart review

(1) The trend chart of the last 2 hours can be displayed with a resolution of one data per second or one data every 5 seconds;

(2) The trend graph of the last 72 hours can be displayed with a resolution of one data every 1 minute, every 5 minutes or every 10 minutes.

Select the "Trend Chart Display" item in the "Menu" and the following window will pop up:



Figure 3-7 Trend chart display

The ordinate represents the measured value, and the abscissa represents the measurement time. "****" is the trend graph cursor. The measurement value of the position it indicates is displayed below the trend graph, and the corresponding time is displayed above the trend graph. Except for the NIBP value, other trends are displayed as continuous curves.

Select the trend chart display of different parameters:

Use the cursor to select the "Parameter Selection" option and modify its display content. When the desired parameter appears, press the spin button and the trend chart of the parameter will appear in the window.

Choose 1-hour or 72-hour trend chart:

Use the cursor to select the "resolution" option. If you want to observe the trend for one hour, you can select 1 second or 5 seconds.

If you want to observe the 72-hour trend, you can choose 1 minute, 5 minutes or 10 minutes.

Observe trend curves that are farther or closer in time:

Use the cursor to select " \blacktriangleleft "instruction, press this key to page forward, and you can observe the earlier trend curve; use the cursor to select " \blacktriangleright "instruction, press this key to turn the page backward and observe the later trend curve.

Get trend data at a certain moment on the current trend chart

Use the cursor to select " ◀ "instruction, press this key to move to the left, the cursor will move to the left, and the moment it points to will also move to the left; use the cursor to select " ▶ " instruction, press this key to move to the right, the cursor will move to the right, and the moment it points to will also move to the right.

Operation example

Observe the SPO₂ trend chart in the last hour:

(1) Press the "MENU" key in the function key area to pop up the "Main Menu";

(2) Select the "Review" item in the menu, and then select "Trend Chart Display";

(3) Select parameters: In the "Parameter Selection" item, turn the knob until "SPO₂" appears in the box:

(4) Select "1 second" or "5 seconds" in the "Resolution" item;

(5) Use the cursor to select " \blacktriangleleft " instruction, press this key and observe the time changes of the trend chart and the changes of the trend curve;

(6) Stop in the period that requires careful observation. If the vertical coordinate ratio is inappropriate, for example, some trend values exceed the current maximum value of the vertical coordinate, select "Adjust Amplitude" to adjust;

(7) If you want to know the measurement value at a certain moment, use the cursor to select " \blacktriangleleft " indication, press this key to move the cursor there, the time is displayed at the top, and the measured value is displayed at the bottom of the curve;

(8) Press the "Exit" key to exit trend chart observation.

3.3.4 Trend table review

The trend table data of the last 72 hours can be displayed in the following resolutions: 1 minute, 5 minutes, 10 minutes, 30 minutes, 60 minutes. Select "Trend Table Review" in the "System Menu" and the following trend table will pop up:

Trend Table X									
Time	Event	HR bpm	PVCs <i>I</i> min	ST1 mV	ST2 mV	SpO2 %			
(24)22:46				-,	-,				
(24)22:45				-,					
(24)22:44					-,				
(24)22:43				-,	-,				
(24)22:42				-,	-,				
(24)22:41					-,				
(24)22:40			<u></u> -	-,	-,				
(24)22:39					-,				
(24)22:38				-,	-,				
(24)22:37									
(24)22:36			<u> </u>	-,	-,				
(24)22:35				-,	-,				
RES. Imin 🗘 🛣 🚺 🕨									

Figure 3-8 Trend table setting menu

The time corresponding to each set of trend data is displayed in the leftmost column, and the date is in brackets. Listed under events are events that have been marked, corresponding to the time of the marked event. Each measured parameter value is recorded in the trend table, as shown in Figure 3-8. Select left or right to view more historical parameters.

The display of NIBP trend data has its particularities. In addition to the measurement value, the time when the NIBP measurement was performed is also displayed under "Measurement Point". If there are multiple measurement values during this period, only one group can be displayed, and one is displayed under "MORE". "*" means there are two or more measurement results.

Choose trend tables with different resolutions:

Use the cursor to select a resolution, use the spin button to change its options, and change the trend data time interval.

Observe trend data for earlier or later dates:

Use the cursor to select " \bigstar "instruction, press this key to page forward and observe earlier trend data; use the cursor to select " \clubsuit " instruction, press this key to page backward and observe trend data at a later date.

Observe trend data for different parameters

Use the cursor to select " \P " instruction, press this key to move left; use the cursor to select " \blacktriangleright " instruction, press this key to move right. You can observe the trend data of 14 groups with different parameters.

3.4 Interface layout

Select the "Interface Layout" option in the "Ribbon Area", and the menu as shown in Figure 3-9 will appear:

Face Layout			X
₩ave 1	ECG	Param 1	ECG
₩ave 2	ECG	Param 2	NIBP
Wave 3	ECG	Param 3	TEMP
Wave 4	Sp02	Param 4	Sp02
₩ave 5	RESP	Param 5	RESP

Figure 3-9 Interface layout menu

In the "Interface Layout" menu, you can adjust the parameters and layout of each waveform display.

3.5 Monitor information

In "Maintenance", you can select "Monitor Information" to view the monitor information, as shown in Figure 3-10.



Figure 3-10 Monitor information

3.6 Monitor settings

Select the "Monitor Settings" option in the "Main Menu", and the menu as shown in

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Figure 3-11 will appear:



Figure 3-11 Monitor settings

In the "Monitor Settings" menu, users can set the following items:

3.5.1 Work interface selection

Select the "Work Interface Selection" item in the "Monitor Settings" menu, and you can see that the current option is the standard interface.

3.5.2 Alarm limit display

By default, the alarm level is displayed next to each parameter.

3.5.3 Alarm pause time

Select "Alarm Pause Time" in the "Monitor Settings" menu and turn the knob to set the time for the alarm to briefly terminate. The system will not perform any alarm processing during this period. The alarm hang-up time can be selected from "1 minute" or "2 minutes".

3.5.4 Parameter alarm form

Select "Customer Maintenance" in the "Monitor Maintenance" menu and enter the password. After entering the settings, go to "Parameter Alarm Mode" and turn the knob

to set the alarm latch or non-latch.

3.5.5 Alarm volume

Select "Alarm Volume" in the "Monitor Settings" menu and turn the knob to set the alarm volume. There are four levels available: "low", "medium", "high" and "off". "Off" means all volumes are turned off.

WARNING

•When the alarm volume of the system is turned off ("Off " is selected), if an alarm occurs, the monitor cannot sound the alarm. Therefore, operators should cautiously use this feature.

•If you select "Off" for the alarm volume in the mute or alarm pause state, the system will automatically end the mute state or alarm pause state.

•When the alarm volume is "off", if the operator selects "mute" or "alarm pause", the system will restore the alarm volume to the alarm volume before turning off the sound, and the system will enter the mute state or alarm pause state.

ANotice:

•The "alarm volume" is only valid for the current setting. It will return to the medium volume after shutting down and then turning it on again. Operators should carefully check this function before use to avoid delaying patient treatment due to silent or too small alarm sounds.

3.5.7 System time setting

Select the "System Time Settings" item in "Monitor Settings", and the menu shown in Figure 3-12 will pop up:



Figure 3-12 System time setting Figure 3-13 Enter maintenance password

Motice

•The system time should be set at boot (if the user needs to set it), otherwise incorrect time information may be provided when reviewing content with time prompt information, etc.!

3.5.8 Event settings

Select the "Item Settings" item in "Monitor Settings" to mark items.

3.6 Monitor maintenance

Select the "Monitor Maintenance" item in the system menu, and the "Enter Maintenance Password" dialog box will pop up, as shown in Figure 3-13 above.

Users can perform user maintenance in the user maintenance menu by entering the user password. Except for the company's designated maintenance personnel, others can cannot perform manufacturer maintenance functions.

In the "Enter Maintenance Password" menu, enter the correct user password (2016), and press the "Confirm" button. The "User Maintenance" menu pops up, and you can set the information as shown in Figure 3-14.



Figure 3-14 User maintenance Figure 3-15 Color customization Language selection: Users can set the text displayed on the screen to be "CHINESE" or "ENGLISH". The specific options are determined by the user's configuration.

Lead naming style: Select "AHA" or "EURO". For details, please refer to the relevant content in "ECG/Respiratory Monitoring" for the difference.

Color customization: used to define the display color of waveforms and parameters on the screen, as shown in Figure 3-15 above.

3.7 Demonstration function

Select the "Demo Function" item in the "System Menu" to pop up the "Enter Demo Password" dialog box. After entering the correct password "5188", the system enters the demonstration waveform state. The demonstration waveform is a simulation demonstration waveform set by the manufacturer only to demonstrate the machine performance and help users with training. In actual clinical use, the function of demonstrating waveforms is disabled because it may cause medical staff to mistake the waveforms and parameters of the patient for monitoring, affecting patient monitoring and delaying diagnosis and treatment. Therefore, this menu has a password, as shown in Figure 3-16 shown.



Figure 3-16 Demonstration function

3.8 Alarm settings

Alarm	Setup						X
	Alarm High Limit	Alarm Low Limit	Alarm Level		Alarm High Limit	Alarm Low Limit	Alarm Level
Hr	120	50 🗘	Medium 🗘	RESP	30 🗘	8 🗘	Medium ᅌ
SP02	100 🗘	90 🗘	Medium ᅌ	т1	39.0 🗢	36.0 🗢	Medium ᅌ
PR	120 🗘	50 🗢	Medium 🗘	T2	39.0 🗢	36.0 🗢	Medium 🗢
SYS	160 🗘	90 🗘	Medium 🗘	TD	2.0 🗘		Medium 🗢
MAP	110 🗢	60 🗘	Medium 🗢	P¥Cs	10 🗘		Medium ᅌ
DIA	90 🗘	50 🗘	Medium ᅌ	ST	0.20 🗘	-0.20 🗘	Medium ᅌ
	Alarm Man	age	C02 A	larm Mer	ıu	IBP Alar	m Menu

Figure 3-17 Alarm settings

Used to set alarm limits and alarm levels for each parameter. To enter the alarm management settings, you need advanced permissions and the password "2016" to enter, as shown in Figure 3-17 shown.

Alarm management

Click "Alarm Management" in the "Alarm Settings" menu to pop up the menu. Enter the permission management password, as shown in Figure 3-17 below. Alarm management is used to set the switch status, alarm delay time, alarm pause time, etc. of various parameter alarms.





Figure 3-18 Alarm management

3.9 Network settings

IP SETUP								Х
LAN IP:	200	¢	200	¢	200	÷	1	¢
LAN Mask:	255	¢	255	¢	255	÷	0	÷
LAN Gateway:	200	÷	200	÷	200	÷	1	÷
₩IFI StaticIP:	200	¢	200	\$	200	÷	1	÷
WIFI Mask:	255	¢	255	¢	255	¢	0	÷
₩IFI Gateway:	200	÷	200	¢	200	÷	1	÷
Server IP:	200	¢	200	÷	200	÷	100	÷

When the monitor needs to perform wired or wireless networking and program upgrade related operations, it is necessary to set up the local network so that other devices can discover or connect to the device.

Operation method: Settings - Maintenance - Enter user password 2016 - Confirm - Set IP. The following dialog box will pop up. You can use the touch screen to operate or move the cursor to the setting item, and then change the setting value, as shown in Figure 3-19.

Chapter 4 Safety

This system has anti-defibrillation and surgical knife protection with floating input. If the correct electrodes are used (see ECG and Respiratory chapter) and placed according to the manufacturer's instructions, the screen display can be restored within 5 seconds after defibrillation.

WARNING: Do not touch the patient, bed, or product during defibrillation.

4.1 Environment:

 Please follow the following guidelines to ensure absolute safety in electrical installation. The environment in which the monitoring system is used must be reasonably protected from vibration, dust, corrosive or explosive gases, extreme temperatures, moisture, etc. When installed in a product cabinet, there must be sufficient space in front for easy operation. With the cabinet door open, there should be enough space behind it to facilitate maintenance. The air circulation inside the cabinet should be ensured.
The monitoring system can meet the technical specifications when working in environments with temperatures below 5 °C to 40 °C. If the ambient temperature exceeds this range, it may affect the accuracy of the product and cause damage to components and circuits. At least 2 inches (5 centimeters) of space should be left around the product to ensure air circulation.

3. 1m (operator's position) priority of visual alarm signal:

"*" indicates the low priority, "**" indicates the middle priority, and "***" indicates the high priority

Please refer to the product specifications section for power requirements.

4.2 Condensation:

During operation, ensure that the product is free of condensation, which may form when the product is moved from one room to another. This is because the product is exposed to moist air and varying temperatures.

 \triangle WARNING: If used in areas with flammable anesthetics, there is a risk of explosion.

For descriptions of other safety matters, please see the introduction.

Chapter 5 Maintenance and Cleaning

5.1 Inspection

Before using this equipment, the following checks must be carried out:

(1) Check the product for any mechanical damage.

(2) Check all exposed wires, plug-in parts and accessories.

(3) Check the functions of all products that may be used to monitor patients, and ensure that the products are in good working order.

(4) If you find signs that the product function is damaged, this monitor must not be used to perform any monitoring on the patient. Please contact the hospital's medical equipment engineer or our company's maintenance engineer.

(5) A comprehensive functional inspection, including safety inspection, must be carried out by professionals every 6-12 months and after each repair.

WARNING

•If hospitals or medical institutions responsible for maintaining this product fail to maintain this product as planned, it may cause the product to fail and ultimately endanger the patient's health.

• Equipment or accessories must be powered off before being disinfected or cleaned.

•All safety inspections or maintenance work that requires dismantling the equipment should be performed by professional maintenance personnel. Operations by non-professionals may cause equipment failure and may endanger personal safety.

•If you find a problem with your device, such as when it accidentally gets wet, please contact the maintenance personnel or manufacturer.

•If the accessories (such as ECG cables, blood oxygen probes, body temperature probes) equipped with the equipment are damaged or show signs of deterioration, they are forbidden to be used again. Please replace the original accessories of our company in time.

•If the blood pressure cuff and extension tube used with the device leak, please replace them in time.

•If all accessories are dirty, please clean and disinfect them in time.

•Various disposable accessories shall not be reused. Please recycle and discard disposable accessories according to specifications after use.
•Please use accessories with corresponding specifications for different patient types.

5.2 General cleaning

This device should be placed in a clean environment.

It is recommended to clean the outer surface of the case and the monitor screen. To clean the case, use a non-abrasive cleaner such as soap and water.

5.3 Use of cleaning agents

In addition to the solutions listed in the "Careful" section below, any solution that can be classified as a product with the following properties may be used as a cleaning agent:

The surface of the monitor and its sensors can be wiped with medical alcohol or isopropyl alcohol, and allowed to air dry or clean with a clean, dry cloth.

Our Company is not responsible for the effectiveness of these chemicals as a means of controlling infectious diseases. Please consult with your hospital's infection control leader or infectious disease specialist.

ACareful

•Do not use strong solvents such as acetone.

•Most detergents must be diluted before use. Follow each manufacturer's instructions for dilution.

•Never use abrasive materials for cleaning (such as steel wool or silver polish).

•When the main unit or its accessories are being disinfected or cleaned, liquid must not enter the joints, and liquid must not enter the main unit. Any liquid remaining on the surface after cleaning or disinfection must be removed.

•Accessories must not be cleaned or disinfected by high temperature, high pressure, or liquid immersion.

5.4 Disinfection

You can disinfect this monitor according to the disinfection procedures of your hospital. This monitor should be cleaned before disinfection.

a. ECG lead wire

1) When cleaning or disinfecting, do not connect the lead wire to the monitor;

2) Clean or disinfect before use on new patients;

3) Use 70% isopropyl alcohol or alcohol solution to clean the surface of the ECG electrode.

4) When cleaning, first dip a clean, dry sponge pad into cleaning fluid. Use this sponge pad to wipe all surfaces of the electrodes and cables; then dampen another clean, dry

sponge pad with sterile or distilled water. Use this sponge pad to wipe all surfaces of the electrodes and cables; finally wipe all surfaces of the electrodes and cables with a clean, dry sponge pad;

5) Careful: The connector pins must not come into contact with cleaning fluid, otherwise it will cause damage to the monitoring equipment and human body.

b. Blood pressure cuff

1) Prepare enzymatic cleaner (such as ENZOL produced in the United States or Cidezyme enzyme cleaner produced in the United Kingdom) and 10% bleach in separate spray bottles.

2) Spray the cleaner evenly on the sleeves, ductwork and hoses. If the dirt has dried, leave it on the clean surface for one minute.

3) Use a lint-free cloth to wipe the smooth surface. If there are visible stains or irregular surfaces, use a soft-bristled brush to clean them. Rinse with plenty of distilled water.

4) When disinfecting, spray 10% bleach solution on the cuff until it is saturated, and leave the product for 5 minutes.

5) Wipe off excess solution, rinse with distilled water again, and let the sleeve air dry naturally.

c. Blood oxygen probe

1. Do not connect the probe to the monitor during cleaning or disinfection.

2. The probe must be disinfected before use.

3. To clean the contact surface between the probe and the patient, use a soft medical cloth dipped in medical cleaning fluid to wipe it.

4. Disinfect the probe and wipe the contact surface between the probe and the patient with disinfectant for at least 1 minute. We recommend 75% isopropyl alcohol as a disinfectant.

5. The probe can withstand 200 times of cleaning and disinfection.

6. Patients should not clean and disinfect themselves, but must let medical staff do it.

d. Medical temperature sensor

Disinfection must be carried out by the hospital or user before use. Disinfectants commonly used in hospitals (such as ethanol, isopropyl alcohol, etc.) can be used to wipe the probe and the wire body in contact with the human body: first wipe with medical alcohol cotton balls, and then use 75% ethanol. Soak for more than 5 minutes and wipe clean.

Careful: To prevent damage to the monitor, do not use gas (EtO) or formaldehyde to disinfect the monitor.

5.5 Sterilization

Sterilization of this monitor, related products, or accessories is not permitted unless otherwise stated in the accompanying instructions.

5.6 Maintenance

In order to ensure that the monitor is available at any time, routine maintenance of the monitor is required. Routine maintenance includes the following inspection items: (1) shift inspection; (2) automatic detection; (3) user detection; (4) ECG cable test; (5) Pace test; (6) NIBP test; (7) NIBP overvoltage protection test, etc.

Cables such as ECG cables, blood oxygen probes, and body temperature probes, probes, and sensors are all important components for obtaining biological signals, and they are prone to wear and tear and should be included in the shift inspection list. The unit using the product should make a maintenance plan.

5.6.1 Routine inspection

(1) Power-on self-test. Each time the monitor is turned on, the alarm light will flash red

and yellow alternately, and then it will make a "beep" sound to complete the self-test.

(2) Visual inspection. The operator should visually inspect the accessories before each

use to confirm that they are not damaged.

(3) CO2 zero calibration: The operator should perform zero calibration of CO2 before

each use, enter CO2 Settings - Zero Calibration, and complete the zero calibration.

Motice

•This equipment should be inspected periodically according to hospital requirements,

and the validity period of the equipment and products should be checked regularly.

Equipment and accessories should be inspected on a daily basis (performed by clinical operators) and on an annual basis (as a service activity). The focus should be on how clinicians test visual and audible alarm signals.

Chapter 6 Alarm

- This chapter introduces general information about alarms and the measures to be taken when an alarm occurs.
- Information about alarms and prompts for each parameter can be obtained in the chapters about each parameter setting.

6.1 Overview

The so-called alarm means that when the patient being monitored undergoes abnormal changes in vital signs, or the machine itself malfunctions and the patient monitoring cannot be carried out smoothly, the monitor will notify the user and medical staff through sound, light, text, etc. tips.

WARNING

•Using different alarm presets for the same or similar equipment in any single area, such as an intensive care unit or cardiac operating room, is potentially dangerous.

•Do not set alarm limits on the monitor that exceed the extreme values, as this will cause the alarm system to fail.

6.2 Alarm attributes

6.2.1 Alarm type

(1) Physiological alarm: If this alarm originates from changes in the patient's vital signs, that is, the monitored patient's physiological parameters exceed a specific range or the patient has physiological abnormalities that cannot be measured by a single physiological parameter exceeding the range, it is called a physiological alarm.

(2) Technical alarm: If this alarm originates from the machine itself, that is, an alarm occurs when patient monitoring cannot be performed accurately due to technical obstacles in the use of the monitor or a malfunction of the machine itself.

(3) Prompt information: The monitor will display some prompt information related to the system status. This information generally does not involve the patient's vital signs. Strictly speaking, prompt information is not an alarm. Prompt information is generally displayed in the prompt information area at the bottom of the screen. Treatment prompt information is displayed in the corresponding treatment information area.

Examples of physiological alarms and technical alarms

Patient or machine condition	Category of alarm
	generated

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The patient's heart rate was measured to be 114bpm, which exceeded the heart rate alarm range set by the user.	Physiological alarm
The patient was found to have ventricular fibrillation	Physiological alarm
The ECG measurement module found that the ECG lead was	T 1 · 1 1
detached	Technical alarm
SPO2 measurement module failed	Technical alarm

6.2.1.1 Physiological alarm classification

Physiological alarms are divided into two situations. One is when the physiological parameters of the monitored patient exceed a specific range, and the other is when the patient has physiological abnormalities that cannot be measured by a single physiological parameter exceeding the range.

The latter is an alarm that can temporarily block the former. The specific ones are as follows:

The ECG signal is too weak; asystole; ventricular fibrillation/ventricular tachycardia; pulse not found; RESP cardiac interference; RESP respiratory asphyxia; others belong to the former situation.

6.2.1.2 Alarm level

Each alarm, whether it is a technical alarm or a physiological alarm, has a level characteristic. The higher the level, when the alarm occurs, the system will prompt the alarm in a more alert way. All technical alarm levels cannot be changed by the user. Some physiological alarm levels can be set by the user, while others are specified by the system and are not allowed to be changed.

Alarm level	Physiological alarm	Technical alarm
Advanced alarm	The patient is in critical condition and may be life-threatening and should be rescued immediately. Such as asystole, ventricular fibrillation/ ventricular tachycardia, etc.	Serious machine failure or misoperation may fail to detect the patient's critical state or result in treatment failure, putting the patient's life in danger. If the battery is low.
Intermediate alarm	If the patient's physiological signs appear abnormal, corresponding measures or treatment should be taken	Certain machine failures or misoperations may not threaten patient safety, but they may also affect the normal monitoring of key physiological parameters and patient



	immediately.	treatment.
Low level alarm	If the patient's physiological signs are abnormal, corresponding measures or treatment may be required.	Certain monitoring functions may not function properly due to machine failure or improper operation, but this will not threaten patient safety.

ANotice:

•It is recommended that operators treat alarm conditions that may result in minor injury and delayed generation of potential injury as low priority alarm conditions.

6.3 Alarm prompt form

When an alarm occurs, there will be sound, light and text prompts. Alarm characteristics: sound pressure level, amplitude, frequency, etc.

Motice

•If the sound pressure level of the auditory alarm signal is lower than the ambient noise, it will prevent the operator from identifying the alarm state. The operator should adjust the sound pressure level of the alarm system to prevent the operator from being unable to identify the alarm state.

Sound pressure level range for each alarm priority as follows:

1) The sound pressure range of medium priority alarm: the minimum volume is 45dB, the maximum volume is 85dB;

2) The sound pressure range of high-priority alarm: the minimum volume is 45dB, and the maximum volume is 85dB.

The characteristics of the prompt signal and the time interval between two prompt signals is less than 1 second.

6.3.1 Sound and light characteristics

Different levels of alarm sound characteristics and light characteristics

Alarm laval	A larm cound characteristics	Alarm light	Alarm
Alaliii ievei	Alarm sound characteristics	characteristics	information
Advanced alarm	Themodeis"Beep-Beep-Beep-Beep-Beep",sounding once every 11 seconds(the interval count is from the startof this sound to the next start)	The alarm light flashes in red with a fast flashing frequency	"***" logo appears on the device screen
Intermediate	The mode is "Beep-Beep-Beep",	The alarm light flashes	"**" mark

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alarm	which sounds once every 25	yellow with a slow	appears on
	seconds (the interval count is from	flashing frequency	the device
	the beginning of this sound to the		screen
	beginning of the next sound)		
	The mode is "beep-beep-", which		"*" mark
Low level	sounds once every 25 seconds (the	The alarm light is	appears on
alarm	interval count is from the start of	always on yellow	the device
	this sound to the next start)		screen

6.3.2 Text characteristics

Background color: The background color of high-level alarm is red, the background color of intermediate alarm is yellow, and the background color of low-level alarm is blue.

The color of the string: Except for the NIBP technical alarm prompt area, it is always black regardless of the alarm level. The color of the string displayed in the NIBP technical alarm prompt area has nothing to do with the alarm level. High-level alarms are displayed in red, medium-level alarms are displayed in yellow, and low-level alarms are displayed in blue. When the measured parameter exceeds the set alarm limit to induce a

physiological alarm, the parameter value that triggered the alarm flashes.

6.3.3 Alarm status icon

In addition to the above alarm methods, the following alarm icons will also appear on the screen, indicating different alarm states.

Indicates that all alarms are suspended. All alarm prompts are temporarily turned off until the alarm pause time expires or the operator manually cancels the alarm pause setting. The system then restores the prompt.

Indicates that the alarm has been reset

Indicates that the alarm sound is turned off. The sound prompt function of the alarm has been artificially and permanently turned off until the operator releases the setting of turning off the alarm volume.

KIndicates that the alarm of a single measurement module is turned off.

When multiple alarms of different levels are generated at the same time, the sound and light prompt will be based on the highest level among the current alarms.

ANote: When **X** sign appears, the system will not be able to give an alarm sound prompt, so the operator should use this function with special caution.

6.4 Alarm status

6.4.1 Overview

For each alarm, there are two states: triggered state and cleared state. It can only be in one state at a time.

Trigger state: The state when the alarm exists.

Clear state: The state in which the alarm does not exist.

All possible alarms are in the cleared state when starting work. When the alarm conditions are met at subsequent times, the alarm enters the triggered state.

For the entire alarm system (that is, for all alarms), the following states are available:

1. Normal state: refers to the state where the alarm can carry out all prompts (including sound, light and text) in the triggered state.

2. Alarm pause state: refers to the state where the alarm is triggered, but no audible and visual text prompts are provided temporarily.

3. Alarm mute state: refers to the state where the alarm is triggered, with light and text prompts but no sound prompts.

ANote: At each moment, the entire alarm system can only be in one state.

•After powering on, the status indicator light lights up and the device automatically performs a safety check. If the self-check passes, you will hear a short "beep" sound, indicating that the alarm system is normal. If the status indicator light does not light up or there is no beep after powering on, it means that the alarm system is faulty and the equipment cannot be used normally and needs to be repaired before it can be put into use. When an alarm occurs, the operator can accurately detect it 1m away from the alarm system.

The maximum alarm status delay and maximum alarm signal generation delay time can be set to $0\sim6s$, and the default display time is 6s.

•The difference between the alarm tone and the alert tone:

The pulse tone alert signal is related to the pulse rate; the higher the pulse rate, the higher the pitch of the alert tone. Pulse tone when the pulse rate is 60 then broadcast a pulse tone once a second; heartbeat tone when the heart rate is 60 then broadcast a heartbeat tone once a second.

6.4.2 Alarm silent state

The alarm silent state means that the alarm prompt sound is turned off. **6.4.3 Alarm pause state**

The following processing is performed when the alarm is suspended:

All alarm sounds and light prompts are prohibited.

All physiological alarm text prompts are prohibited.

The physiological alarm description area displays the number of seconds left until the alarm is suspended.

6.4.4 Status switching

Under normal conditions:

1. Press "PAUSE" to enter the alarm pause state, and press "SILENCE" to enter the alarm silent state.

In paused state:

1. Press "PAUSE" to enter the normal state, and press "SILENCE" to enter the alarm silent state.

2. If no button is pressed until the pause end time is reached, it will enter the normal state.

3. During the pause period, if there is a new technical alarm, the alarm pause state will end and enter the normal state.

4. During the pause period, if there is a new physiological alarm, the system will still be in the alarm pause state.

In the alarm silent state:

1. If a new technical or physiological alarm is generated, the current alarm silent state will end and enter the normal state.

2. Press "PAUSE" to enter the normal state, and press "SILENCE" to enter the normal state.

In any state:

1. In the user settings, set the alarm sound switch to off and enter the alarm sound off state.

2. In the user settings, set the alarm sound switch to on and enter the normal state.

6.5 Alarm method

6.5.1 Overview

There are two alarm modes: latching mode and non-latching mode.

Latching: When the alarm condition does not exist, the system still prompts the alarm, which is called latch mode. Only after resetting the alarm system can the alarm no longer be prompted.

Non-latching: The feature of no longer alarming when the alarm condition does not exist is called non-latching mode.

6.5.2 Corresponding latching methods

1. All physiological alarms can work in latching mode.

2. All technical alarms can only work in non-latching mode.

6.5.3 Alarm prompt after latching

When an alarm is latched (meaning that the alarm has occurred but is not in the alarm triggering state at this time), the prompt mode related to the alarm will change as follows:

1. Measurement parameters and related alarm limits no longer flash.

2. After the alarm description prompt entry, there is the system time of the last time it

entered the trigger state.

6.5.4 Clearing the latch method

Clearing the latch mode is also called alarm reset. The user can use the alarm pause function to reset the latch alarm. When the latch alarm is cleared, those alarms that have occurred in the past but are still prompting alarms when the alarm conditions no longer exist due to the latch method will be cleared.

When working in non-latching alarm mode, the alarm pause key on the keyboard module only has the function of pausing the alarm but has no reset function.

6.6 Alarm settings

Public alarm content settings

1. You can set the high alarm limit, low alarm limit and alarm level of each parameter in the "Alarm Settings" menu.

2. The alarm volume can be set in the "Monitor Settings" menu.

a) Alarm limit display: You can see the set upper and lower alarm limits in each display parameter area.

b) Alarm volume: There are four options, namely "low", "medium", "high" and "off"

3. The alarm pause time and parameter alarm form can be set in the "User Maintenance" menu of "Maintenance".

a) Alarm pause time: refers to the alarm pause time. There are four options, namely 1 minute and 2 minutes.

b) Parameter alarm form: latching mode and non-latching mode.

Alarm settings for each measurement parameter

The alarm settings for each parameter are in the corresponding parameter setting menu.

Select the parameters that need to be set in "Parameter Settings", and you can set the

alarm switch and alarm record of each parameter individually.

When the intended use includes monitoring patients who do not receive constant attention from clinical operators, it is recommended that the alarm tone be set to level 2 or above.

Item		Default setting
	Alarm upper limit	Adult: 120;
		Children: 160
Heart Rate	Alarm lawor limit	Adult: 50;
	Alami lower mint	Children: 75
	Alarm level	Middle
Dlaad	Alarm upper limit	Adult: 100; Child: 100
Dioou	Alarm lower limit	Adult: 90; Child: 90
Oxygen	Alarm level	Middle
	Alarm upper limit	Adults: 120; Children: 160
Pulse Rate	Alarm lower limit	Adult: 50; Child: 75
	Alarm level	Middle
Systolic	Alarm upper limit	Adult: 160; Child: 120
Blood	Alarm lower limit	Adult: 90; Child: 70

Default alarm settings

Pressure	Alarm level	Middle	
Maan	Alarm upper limit	Adults: 110; Children: 90	
Program	Alarm lower limit	Adult: 60; Child: 50	
riessuie	Alarm level	Middle	
Diastolic	Alarm upper limit	Adult: 90; Child: 70	
Blood	Alarm lower limit	Adult: 50; Child: 40	
Pressure	Alarm level	Middle	
Descrimentary	Alarm upper limit	Adult: 30; Child: 30	
Respiratory	Alarm lower limit	Adult: 8; Child: 8	
Kale	Alarm level	Middle	
D 1	Alarm upper limit	Adult: 39.0; Child: 39.0	
Tomporatura	Alarm lower limit	Adult: 36.0; Child: 36.0	
Temperature	Alarm level	Middle	
	Alarm upper limit	Adult: 50; Child: 50	
EtCO2	Alarm lower limit	Adult: 15; Child: 20	
	Alarm level	Middle	
	Alarm upper limit	Adults: 4; Children: 4	
FiCO2	Alarm lower limit	/	
	Alarm level	Middle	
AWRR	Alarm upper limit	Adult: 30; Child: 30	
	Alarm lower limit	Adult: 8; Child: 8	
	Alarm level	Middle	

Motice

•Only the alarm settings made 3 seconds before the power interruption can be successfully saved in the alarm settings.

•It is recommended that alarm systems required by the patient population use factory

default settings, preferred alarm settings and alarm presets.

6.6.1 Alarm shutdown

Alarm shutdown refers to the failure of the alarm function of a single measurement module. At this time, even if the measurement module meets the alarm conditions, the system will not issue any alarm prompts, print alarms, or store alarms.

When any measurement module just starts working, all alarms related to the module will be automatically turned off within 30 seconds after the module starts working, and other alarms will not be affected.

6.6.2 The lead falls off when turning on the machine

When powering on, if the open parameter module does not have a lead connected:

1. For ECG or SPO2 modules, the product changes the lead-off alarm prompt to a prompt

message (that is, the sound and light are automatically cleared), and then prompts the

user.

2. There is no lead loss alarm for other modules.

6.7 Parameter alarm

Alarm parameters can be set independently in each parameter menu, and users can set alarm limits and alarm status.

When a parameter alarm is turned off, "⁽²⁾," prompt symbol will display. The alarm switch of each parameter can be set independently.

For parameters that set alarms, when the value of one or several parameters exceeds the alarm limit, the monitor automatically alarms and performs the following processing:

1) A prompt appears on the screen in the form described in Alarm Mode;

2) If the alarm volume is set, the alarm sound will be sounded according to the set alarm level and alarm volume;

3) The alarm light flashes (if the machine has an alarm light).

6.8 Measures to be taken when an alarm occurs

Motice: When an alarm occurs, the patient's condition should be checked first.

The alarm information is displayed in the system information area or system alarm information area. It is necessary to identify this alarm and take corresponding measures according to the cause of the alarm.

(1) Check the patient's condition.

(2) Identify which parameter is alarming or which alarm is occurring.

(3) Identify the cause of the alarm.

(4) If necessary, set the alarm to be muted.

(5) When the alarm condition is relieved, check whether the alarm is eliminated. Alarm information and prompt information about parameters can be found in each parameter monitoring chapter.

6.9 Alarm reset

ANotice: The alarm reset operation can only be performed in the latch state. In

the non-latched state, the alarm cannot be reset.

Alarm reset operation method: Press the alarm reset button to reset the current alarm, and

the alarm reset icon "2" will be displayed in the information bar, and the prompt message "Alarm resetting" will be displayed. The reset alarm information will no longer be displayed if no new alarm occurs.

6.10 Password protected alarm settings

The following alarm settings are password protected:

- 1. Alarm delay time
- 2. Alarm pause time
- 3. Parameter alarm form
- 4. Alarm sound range
- 5. Latching and non-latching settings

Chapter 7 ECG and Respiration (ECG/RESP)

7.1 ECG monitoring instructions 7.1.1 Definition of ECG monitoring

ECG monitoring generates a continuous waveform of the patient's cardiac electrical activity to accurately assess the patient's physiological state at that time. For this reason, the normal connection of the ECG cable should be ensured so that correct measurement values can be obtained. With five-lead or 12-lead device for monitoring, the monitor in the normal working state, the standard interface shows three waveforms, and other interfaces show different waveforms, such as: large font interface shows one ECG waveform; ECG multi-conductor interface shows seven ECG waveforms;

(1) The parameters displayed by the monitoring include heart rate (HR).

(2) All the above parameters can be used as alarm parameters.

AWARNING

•During defibrillation, do not touch the patient or any connected objects such as tables or products that come into contact with the patient.

•When using a monitor for ECG signal monitoring, you must use the ECG cable recommended by our company.

•If the patient needs to be defibrillated, a defibrillation-resistant ECG cable must be used, otherwise the device may be damaged or the patient may be harmed.

•When you connect electrodes or cables, make sure there is absolutely no contact with any other conductive parts or with ground. In particular, make sure that all ECG electrodes, including the neutral electrode, are attached to the patient to prevent them from coming into contact with conductive parts or ground.

•Regularly check the skin and pads where the ECG electrodes are placed. If there are signs of allergy, replace the electrode pads or change the placement of the electrode pads. Do not use expired electrode pads.

•For patients with built-in pacemakers, the detection function switch should be turned on in the patient information and ECG settings to ensure that the pacemaker can calculate normally. Please focus on this item.

Motice

•Interference from ungrounded products near the patient and ESU interference can cause waveform distortion.

•If operated according to the conditions specified in YY 9706.102-2021 (when the anti-radiation capability is 3V/m), the field strength exceeding 1V/m may cause errors in various frequency measurements. Therefore, it is recommended not to use the device in the vicinity of electrocardiographic/respiratory measuring equipment.

•Electrode pads should be of the same model and should not exceed their service life.

7.2 ECG monitoring operation method

7.2.1 Preparation

1) Prepare the patient's skin before placing electrodes.

a. The skin is a poor conductor, so to obtain good contact between the electrode and the skin, the patient's skin pretreatment is very important.

b. If necessary, shave body hair where the electrodes are placed.

c. Wash skin thoroughly with soap and water. (Do not use ether or pure alcohol as this will increase the impedance of the skin).

d. Dry rub the skin to increase capillary blood flow to the tissues and remove skin flakes and oil.

2) Install the spring clip or snap button before placing the electrode.

3) Place the electrode on the patient. If you are using an electrode without conductive paste, apply conductive paste before placement (or operate according to the requirements of the electrode manufacturer).

4) Connect the electrode leads to the patient cable.

5) Confirm that the monitor is powered on.

WARNING

•The electrodes should be placed carefully and make sure the contact is good.

•The ECG electrode patch should be checked daily for irritation of the patient's skin. If there are signs of allergy, replace electrodes or change position every 24 hours.

•It is necessary to check whether the leads are normal before starting monitoring. After unplugging the ECG cable, the screen will display the prompt "Sensor is off" and trigger an audible alarm at the same time.

7.2.2 Install the cardiac conductor

Position of ECG monitoring electrodes

The electrode placement of the five-lead device is shown in Figure 8-1 shown. When there are three leads, only RA, LA, and LL need to be connected.

(1) RA white (right arm) electrode 1 is placed under the clavicle, close to the right shoulder.

(2) LA black (left arm) electrode 1 is placed under the clavicle, close to the left shoulder. Place it on the chest wall as shown below.

(3) RL green (right leg) electrode 1 is placed on the right lower abdomen.

(4) The LL red (left leg) electrode is placed on the left lower abdomen.

(5) The V brown (chest) electrode is placed on the chest wall as shown in Figure 8-2.

The following table lists the lead names in European and American standards

respectively. (In the European standard, each lead is represented by R, L, N, F, and C,

while in the American standard, it is represented by RA, LA, RL, LL, and V.)

U	SA	Euro	pe
Lead name	Color	Lead name	Color
RA	White	R	Red
LA	Black	L	Yellow
LL	Red	F	Green
RL	Green	Ν	Black
V	Brown	С	White



Figure 7-1 5-lead electrode placement location

Motice

•To ensure patient safety, all leads must be connected to the patient.

WARNING

•When using electrosurgical (ES) equipment, place the ECG electrode halfway between the ES ground plate and the electrosurgical blade to avoid burns. Cables from electrosurgical equipment should not be entangled with ECG cables.

•When using electrosurgical (ES) equipment, never place the electrode close to the ground terminal of the surgical electrical equipment, otherwise there will be a lot of interference on the ECG signal.

•The placement of ECG leads depends on the type of surgery being performed. For

example, for open-heart surgery, electrodes can be placed across the side or back of the chest. In the operating room, due to the use of surgical electrocautery equipment, sometimes artifacts may affect the ECG waveform. To help reduce artifacts, electrodes can be placed on the left and right shoulders, near the left and right sides of the abdomen, while chest leads can be placed on the left side of the chest center. Avoid placing the electrode on the upper arm, otherwise the ECG wave will become very small. Characteristics of a good signal:

Tall, narrow and without incision. The R wave is tall and lies completely above or below the baseline. The pacing signal is no greater than the height of the R wave. The T wave is less than 1/3 the height of the R wave. P waves are much smaller than T waves.



Figure 7-2 Standard ECG waveform

You can use the 1mV ECG calibration wave to calibrate the ECG. At this time, the screen prompts "The patient cannot be monitored during calibration."

Using a five-lead ECG device

Users can arrange the leads on channels 1, 2, and 3 according to their needs. The lead labels on the three channels are displayed on the left side of the corresponding waveform and can be changed in the ECG menu. You can select appropriate leads from I, II, III, AVR, AVL, AVF, and V for channels 1, 2, and 3 respectively, as shown in Figure 8-4. When the user selects the same lead, the monitor automatically adjusts to a different lead.



Figure 7-3 ECG lead diagram 53 / 113

Motice

•If the electrodes are attached correctly but the ECG waveform is inaccurate, the lead wires need to be replaced.

•Interference from ungrounded products near the patient and ESU interference can cause waveform problems.

7.3 ECG Menu

ECG setup menu

Turn the knob, move the cursor on the main screen to the ECG hotkey in the parameter area, and then press the knob to pop up the ECG setting menu, as shown in Figure 7-4:



Figure 7-4 ECG setting menu

Heart rate alarm

Select "On" to prompt and store the alarm when the heart rate alarm occurs. Select "Off" to not alarm and prompt "

ANotice

• The upper and lower alarm limits should be set according to the clinical conditions of each patient.

•The heart rate alarm upper limit setting is very important in monitoring. The upper limit should not be set too high. Taking into account changing factors, the upper limit of the heart rate alarm should not be set 20 beats/min higher than the patient's heart rate.

ECG channel

There are three channels in total, namely: ECG 1, ECG 2, and ECG 3. Optional leads are

I, II, III, aVR, aVL, aVF, and V.

ECG gain

There are five options in total, namely $\times 0.25 \times 0.5$, $\times 1$, $\times 2$ and automatic.

Motice

•When the input signal is too large, the wave peak may be truncated. At this time, the user can refer to the actual waveform to manually change the gain level of the ECG waveform to avoid incomplete waveform display.

•The gain of each calculation channel can be selected, and a 1mV scale is given on the left side of each ECG waveform. The height of the 1mV scale is proportional to the amplitude.

Heart rate source

You can choose to detect the heart rate by ECG (electrocardiogram) or SPO2 (blood oxygen volume tracing wave); select "Auto" and the monitor will determine the heart rate source based on signal quality; if "Simultaneous" is selected, the monitor will display the heart rate and heart rate at the same time. Pulse rate. If provided by SPO2, it prompts PULSE (pulse) and has a pulse rate sound.

When the heart rate source is selected as SPO2, the heart rate alarm judgment is not performed, but the pulse rate alarm judgment is performed.

When the "select all" option is selected, the measurement value of PR is displayed on the right side of the main screen SOP2; HR and PR alarm at the same time. The heartbeat sound is based on HR. If HR has data, it will give a sound prompt. If there is no HR data, it will give a sound prompt for PR.

Calculation channel

(1) "Channel 1" "Channel 2" "Channel 3" means calculating the heart rate based on the waveform data of the 1st, 2nd or 3rd ECG wave.

(2) "Auto" means that the monitor automatically selects the channel for calculating heart rate.

Lead type

Choice of 5-lead or 3-lead

Waveform speed

The ECG waveform scanning speed is available in three levels: 12.5, 25.0 and 50.0mm/s.

Other settings

Select this item to enter the "ECG Other Settings" menu, as shown in Figure 7-5:



Figure 7-5 ECG other settings menu

The following functions are available in this submenu:

(1) Heartbeat volume: You can choose the volume levels of "low", "medium", "high" and "off".

(2) Pacing analysis: You can choose "On" or "Off".

(3) Power frequency trap: Suppress grid interference.

(4) ECG calibration: Select this option to automatically calibrate the ECG waveform.

(5) Default settings: Select this item to enter the ECG default configuration dialog box. You can choose the system default configuration.

▲Notice: When the input signal is too large, the wave peak may be truncated. At this time, the user can refer to the actual waveform to manually change the gain level of the ECG waveform to avoid incomplete waveform display.

The gain of each calculation channel can be selected. There are four gain levels: $\times 0.25$, $\times 0.5$, $\times 1$, and $\times 2$. The corresponding scale is given on the left side of each ECG waveform. The height of the corresponding ruler is proportional to the wave amplitude. Filtering can result in cleaner or more precise waves.

Three filtering methods can be selected. The reference mode displays unfiltered ECG waves; the monitoring mode filters out artifacts that may cause false alarms; and the surgical mode reduces artifacts and interference from electrosurgical equipment in the operating room.

7.4 ECG alarm information and prompt information Alarm information

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Alarms that may occur during ECG measurement are divided into physiological alarms and technical alarms. At the same time, the ECG measurement process may also generate various prompt messages. When these alarms or prompts occur, the visual representation and auditory representation of the monitor can refer to the relevant descriptions in the alarm function chapter. On the display screen, physiological alarms and general prompt information (general alarms) are displayed in the alarm area of the monitor, while technical alarms and prompt information that cannot trigger the alarm are displayed in the information area of the monitor. When the alarm recording switch in the relevant menu is turned on, physiological alarms caused by parameters exceeding alarm limits may trigger the recorder to automatically output alarm parameter values and related measurement waves.

The following category list describes the various alarms that may be generated by this measurement part.

Prompt message	Cause	Alarm level
ECG signal is too	Undetectable patient's ECG signal	High
weak		
HR is too high	HR measurement value is higher	User selectable
	than the set alarm high limit	
HR is too low	HR measurement value is lower	User selectable
	than the set alarm low limit	

Physiological alarm:

Technical alarm:

Prompt message	reason	Alarm level	Countermeasures
All ECG leads or one lead reported	The ECG electrodes fall off the patient or the ECG cable falls off the monitor.	Low	Make sure the electrodes, leads and cables are all connected properly.
missing	Intentionally disconnected by clinical operator	Low	The operator can turn off the alarm by pressing the "alarm reset" button.
ECG module communication stopped	ECG measurement module failure or communication failure	High	Stop using the measurement function provided by the ECG module and notify the biomedical engineer or our company's maintenance personnel.
HR alarm limit error	Functional safety failure	High	Stop using the HR alarm function and notify biomedical engineers or our company's maintenance personnel.
ECG interference	ECG measurement	Low	Be sure to keep the patient

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is too great	signals are greatly affected by	quiet, ensure the electrodes are connected reliably, and the AC
	interference	power supply system is well grounded.

Prompt information (including general alarm information):

Prompt message	Cause	Alarm level
HR measurement	HR measurement value exceeds	Uich
out of bounds	measurement range	Tiigii

7.5 Respiration measurement

How is breathing measured?

The monitor measures respiration from the thoracic impedance values of two electrodes, and the change in impedance between the two electrodes (due to thoracic activity) produces a respiratory waveform on the screen.

Respiratory monitoring settings

No additional electrodes are required to monitor respiratory parameters, but the placement of the electrodes is important. In some patients, due to their clinical condition, the thoracic cage expands laterally resulting in negative intrathoracic pressure. In this case, it is best to place the two RESP electrodes on the right midaxillary line and the left side of the thorax when the patient's respiratory activity is maximum to obtain the best respiratory waveform.

ANote: Respiratory monitoring is not suitable for patients with a large range of motion, as this may lead to false alarms.

RESP Guardianship Check:

- 1) Prepare the patient's skin before placing electrodes.
- 2) Install the electrodes on the patient as described below.







II-lead RESP measurement electrode position diagram

Motice

Place the white RA and red LL electrodes diagonally to obtain the best respiratory waveform in the II lead, or place the RA and LA electrodes in order to obtain the I lead respiratory waveform. When selecting different respiratory leads, be careful to select the corresponding calculation lead in RESP Settings - Respiratory Lead Settings, otherwise the correct result cannot be measured. Placing the liver and ventricles in line with the respiratory electrode should be avoided to avoid artifacts from cardiac coverage or pulsatile blood flow.

RESP settings menu

Turn the knob, move the cursor to the RESP hotkey in the parameter area on the main screen, and then press the knob to enter "RESP Settings" menu, as shown in Figure 7-7.



Figure 7-7 RESP setting menu

RESP alarm settings

(1) Alarm switch: Select "On" to prompt and store the alarm when the respiratory rate

alarm occurs. Select "Off" to not alarm and prompt "

The respiratory rate alarm is based on the set high limit and low limit. When the respiratory rate exceeds the set value, an alarm occurs.

The adjustment range of the RESP alarm upper and lower limits is as follows:

Model	Maximum limit	Minimum floor	Single adjustment
			amount
RR adult	120	7	1
RR children	147	7	1

(2) Suffocation alarm: Set the time to judge the patient's suffocation, between 10 seconds

and 40 seconds. Each time the knob is turned, add/subtract 5 seconds.

(3) Waveform speed

The optional respiratory wave speeds are 6.25mm/s, 12.5mm/s, and 25.0mm/s.

(4) Breathing Gain

Users can set the magnification display of RESP waveform, and the magnification options are 0.25, 0.5, 1, 2, and 4.

(5) Default configuration

Select this item to enter the "RESP Default Configuration" dialog box. The user can select "Manufacturer Default Settings" or "User Default Settings". After selecting to exit the dialog box, the system will pop up a dialog box asking the user to confirm their choice.

7.6 RESP alarm information and prompt information

When the alarm recording switch in the relevant menu is turned on, those physiological alarms caused by parameters exceeding the alarm limit will trigger the recorder to automatically output alarm parameter values and related measurement waveforms. The physiological alarms, technical alarms and prompt messages that may occur during RESP measurement are listed in the following table:

Physiological alarm:

Prompt message	Cause	Alarm level
RR is too high	RESP measured value is higher than the	User selectable
	set alarm high limit	
RR is too low	RESP measured value is lower than the set	User selectable
	alarm low limit	
RESP respiratory	Breathing cannot be measured within a	High
apnea	specific time interval	

Technical alarm:

Prompt	Cause	Alarm	Countermeasures
message		level	
RESP alarm limit error	Functional safety failure	High	Stop using the RESP alarm function and notify biomedical engineers or our company's maintenance personnel.

Prompt information (including general alarm information):

Prompt message	Cause	Alarm level
RR measurement	RR measurement value exceeds	High
out of bounds	the measurement range	

Chapter 8 Blood Oxygen Saturation (SPO2)

8.1 Instructions for blood oxygen saturation monitoring

The SPO2 can be used on adults and children, and the pulse oximeter device is calibrated to display functional oxygen saturation. The SPO2 accuracy of the pulse oximeter device is the root mean square of a difference and is less than or equal to 4.0%SpO2 in the range of 70% to 100%SaO2. SpO2 should be expressed as functional oxygen saturation, not as oxyhemoglobin fraction.

Definition of SPO2 monitoring

The SPO2 plethysmography parameter measures arterial oxygen saturation, which is the percentage of total oxyhemoglobin. For example, if 97% of the total hemoglobin molecules in the red blood cells of arterial blood are combined with oxygen, then the blood has an SPO2 oxygen saturation of 97%, and the SPO2 value reading on the monitor should be 97%. The SPO2 value shows the percentage of oxygen-carrying hemoglobin molecules that form oxyhemoglobin. The SPO2 plethysmography parameter also provides pulse rate signals and plethysmography waves.

(1) Blood oxygen saturation is measured using pulse oximetry. This is a continuous, non-invasive method of measuring hemoglobin oxygen saturation. It measures the amount of light emitted from the sensor light source that passes through the patient's tissue (such as a finger or ear) and reaches the receiver on the other side. The wavelength that the sensor can measure is usually 660nm for red LED and 905nm for infrared LED. The maximum optional output power of LED is 4mW.

(2) The amount of light passing through depends on a variety of factors, most of which are constant. However, one of these factors, arterial blood flow, changes over time because it pulsates. By measuring the light absorbed during the pulsation, it is possible to obtain the oxygen saturation of the arterial blood. Detecting the pulsation itself gives a "plethysmographic" waveform and pulse rate signal.

Motice

•Because the measurement values of pulse oximeter equipment are distributed based on statistical probability, only about 2/3 of the measurement values of pulse oximeter equipment fall within \pm Arms of the value measured by the carbon monoxide-blood gas analyzer.

•Functional testers cannot be used to evaluate the accuracy of pulse oximeter probes or

pulse oximeter monitors.

WARNING

•If carboxyhemoglobin, methemoglobin or dye diluting chemicals are present, the SPO2 value will be biased.

•Please use the blood oxygen sensor recommended in the manual and refer to its manual for use.

•Before use, users need to check the compatibility between the monitor, sensors, and cables, otherwise the patient may be injured or the equipment may be damaged.

•When a patient has a tendency to be hypoxic, an oximeter should be used to analyze the blood sample to fully understand the patient's blood oxygen changes.

•Avoid using this device and blood oxygen sensor in the environment of MRI equipment, otherwise nuclear magnetic induction current may cause severe burns to the patient.

•When patients are continuously monitored for a long time, the probe attachment position should be checked every 2 hours, and when the skin changes or the measurement position is changed every 4 hours, some patients (patients with perfusion disorders or skin sensitivity) may require more frequent examine. Continuous long-term monitoring may increase skin risks such as irritation, redness, blistering, or pressure necrosis.

SPO2 plethysmography parameter measurements:

(1) The "SPO2" value and "plethysmography" waveform can be displayed on the main screen.

(2) SPO2 in this manual refers to human functional blood oxygen saturation measured by non-invasive methods.

(3) The measured data update period is 1 second.

(4)In order to stabilize the blood oxygen waveform in the display area, the blood oxygen waveform is normalized..

ANotice:

•The product will detect when a finger falls off by searching for the pulse, and the "SPO2 finger off" alarm will be delayed for 3s-5s.

•If carboxyhemoglobin, methemoglobin or dye diluting chemicals are present, the SPO2 value will be biased.

•Electrosurgical equipment cables must not become entangled with sensor cables.

•Do not place the sensor on a limb that has an arterial catheter or an intravenous line.

•Before starting monitoring, check whether the sensor cable is normal. When the SPO2

sensor cable is unplugged from the socket, the screen will display a "sensor off" error message and trigger an audible alarm.

•If the disposable packaged sensor shows signs of damage, do not use the SPO2 sensor and return it to the manufacturer.

•Do not place the blood oxygen probe on the same limb as the blood pressure cuff for blood pressure measurement, because blood flow occlusion during blood pressure measurement will affect the blood oxygen saturation reading.

- •1) Make sure your nails cover the light.
- •2) The probe wire should be placed on the back of the hand.
- The SPO2 value is always displayed in a fixed place, and the pulse rate is only displayed under the following circumstances:
- ●1) Set "Heart Rate Source" to "SPO2" or "Auto" in the ECG menu.

•2) Set "Heart Rate Source" to "Auto" in the ECG menu, and there is no ECG signal at this time.

•The SPO2 waveform is not proportional to pulse volume.

•Functional test equipment or blood oxygen simulators cannot be used to verify the accuracy of blood oxygen saturation monitors and pulse oximeters. The accuracy of oxygen saturation monitors and pulse oximeter probes needs to be verified by clinical data.

• The blood oxygen probe and extension cable used with the monitor have been confirmed and tested for compliance with YY0784-2010.

8.2 Blood oxygen saturation monitoring operation method SPO2 plethysmography measurement

- 1) Turn on the monitor;
- 2) Place the sensor on the appropriate position of the patient's finger;
- 3) Insert the connector at one end of the sensor cable into the SPO2 hole.



Figure 8-1 Adult blood oxygen probe

ANotice

•If the test site and probe cannot be accurately positioned, it may lead to inaccurate blood oxygen saturation readings, or even failure to search for pulse waves and prevent blood oxygen monitoring. In this case, the probe should be repositioned.

•Excessive movement of the measurement part may cause inaccurate measurement. At this time, the patient should be kept quiet or the measurement part should be changed to reduce the impact of excessive movement on the measurement.

•When the displayed pulse oxygen saturation or pulse rate value may be incorrect, the device will display the signal incompleteness indicator "?" symbol.

AWARNING

•During long-term continuous monitoring, check the peripheral circulation and skin conditions of the measurement site every 2 hours or so. If adverse changes are found, the measurement site should be changed in time.

•During long-term continuous monitoring, the positioning of the probe should be checked periodically to avoid changes in the positioning of the probe due to movement and other factors, which would affect the accuracy of the measurement.

•The way this product confirms the accuracy of blood oxygen measurements is by comparing the reading with the value from the blood gas analysis.

8.3 Blood oxygen saturation monitoring measurement limitations

During operation, the following factors can affect the accuracy of blood oxygen saturation measurement:

1) High-frequency electrical interference, including interference generated by the host system itself or interference from electrosurgical products such as those connected to the system.

2) Do not use photooximeters and blood oxygen sensors during magnetic resonance imaging (MRI) scans. Induced current may cause burns.

3) There is dye injection into the vein.

4) The patient moves too frequently.

- 5) External light radiation.
- 6) The sensor is improperly installed or in improper contact with the object.
- 7) Sensor temperature (the optimal temperature should be in the range of 28°C~42°C).
- 8) Place the sensor on the limb with a blood pressure cuff, arterial catheter, or

intravascular catheter.

9) High concentrations of dysfunctional hemoglobin such as carboxyhemoglobin and methemoglobin.

10) Blood oxygen saturation is too low.

11) Poor circulatory perfusion at the monitoring site.

12) Shock, anemia, hypothermia, and the use of vasoconstrictor drugs may reduce arterial blood flow to unmeasurable levels.

13) The measurement also depends on the absorption of light of specific wavelengths by oxyhemoglobin and reduced hemoglobin. If other substances that absorb the same wavelength are present, they can cause spurious or low SPO2 values in the measurement. Such as: carbonized hemoglobin, methemoglobin, methylene blue, carmine indigo.

14) The recommended SPO2 sensor in the attachment is not used.

8.4 Blood oxygen saturation menu

SPO2 setup menu

Turn the rotary button, move the cursor in the display interface to the SPO2 hot key in the parameter area, and press the rotary button to enter the "SPO2 Settings" menu, as shown in the figure 8-2.



Figure 8-2 SPO2 setting menu

WARNING

•Setting the SPO2 alarm upper limit to 100% is equivalent to turning off the upper limit alarm. High oxygen levels can cause premature infants to develop crystalline posterior fibrous tissue syndrome. Therefore, the upper alarm limit for blood oxygen saturation must be carefully selected based on accepted clinical practice experience.

Alarm switch

Select "On" to prompt and store the alarm when SPO2 alarms. Select "Off" to not alarm

and prompt "X".

SPO2 and PR alarm range:

Parameter	Maximum limit	Minimum floor	Single adjustment amount
SPO2	100	0	1
PR	300	0	1

The default alarm range of SPO2 and PR under default settings:

Parameter		Maximum limit	Minimum floor
SPO2	Adult	100	90
	Child	100	90
PR	Adult	120	50
	Child	160	75

Waveform speed

SPO2 plethysmography waveform scanning speed has two options: 12.5 and 25.0mm/s.

Pulse volume

Volume levels can be selected from "Low", "Medium", "High" and "Off"

Measurement sensitivity

Select the averaging time for calculating SPO2 values. Selecting "High", "Medium" or "Low" means taking the average SPO2 within 4 seconds, 8 seconds or 16 seconds.

Default configuration

Select this item to enter the SPO2 default configuration dialog box. You can choose the system default configuration.

8.5 Blood oxygen saturation alarm information

SPO2 alarm information

When the alarm recording switch in the relevant menu is turned on, those physiological alarms caused by parameters exceeding the alarm limit will trigger the recorder to

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automatically output alarm parameter values and related measurement waveforms. The physiological alarms, technical alarms and prompt messages that may occur during SPO2 module measurement are listed in the table below.

Physiological alarm:

Prompt message	Reason	Alarm level
SPO2 is too high	The SPO2 measured value is higher than	User selectable
	the upper alarm limit.	
SPO2 is too low	The SPO2 measured value is lower than the	User selectable
	lower alarm limit.	
PR is too high	The PR measured value is higher than the	User selectable
	upper alarm limit.	
PR is too low	PR measurement value is lower than the	User selectable
	lower alarm limit	

Technical alarm:

Prompt message	Reason	Alarm level	Remedies	
SPO2 sensor falls off	SpO2 sensor detaches from patient or monitor	Low	Make sure the sensor is placed on the patient's finger or other part, and that the monitor and cable are connected properly.	
SPO2 module communication stops	SPO2 module error or communication error	High	Stop using the SPO2 module measurement function and notify biomedical engineers or our company's distributors.	
PR alarm limit error	Functional safety failure	High	Stop using the SPO2 module measurement function and notify biomedical engineers or our company's distributors.	

Prompt information (including general WARNINGs):

Prompt message	Reason	Alarm level
SPO2 measurement	SPO2 measurement value is out of	Iliah
out of bounds	range	rigi

URAJEDI		
PR measurement	DD magazinement value is out of remov	High
out of bounds	PK measurement value is out of range	
Search pulse	The SPO2 module is searching for a	No alarma
	pulse	no alarm
Pulse not found	The SPO2 module cannot detect the	Iliah
	SPO2 signal for a long time.	підіі

8.6 Blood oxygen saturation accuracy

CDACEDV"

The calculated value of the root mean square of the blood oxygen value (SPO2) measured by the monitor and the blood gas analysis value (SaO2) of the corresponding blood sample is used as an indicator to evaluate the accuracy. The blood oxygen accuracy should be $\leq 2\%$ to be qualified. The calculation formula is:

$$A_{rms} = \sqrt{\frac{\sum_{i=1}^{n} (SpO_{2i} - SaO_{2i})^2}{n}}$$

8.6.1 Main evaluation index-scatter plot (i.e. Bland-Altman plot)

(1) SPO2 and SaO2 scatter plot with Ampcon adult finger clip blood oxygen probe



(2) SPO2 and SaO2 scatter plot of adult finger-clip blood oxygen probe equipped with Zhongcang



(3) SPO2 and SaO2 scatter plot of blood oxygen probe with Zhongcang newborn package



Chapter 9 Body Temperature (TEMP)

9.1 Instructions for temperature monitoring

The monitor can use a temperature probe to measure body temperature data. The operating mode adopted by this equipment is the direct mode, and the output temperature is an unadjusted temperature, which represents the temperature of the measurement part contacted by the probe. The recommended measurement site is the human armpit (rated output range: 36.0°C~37.0°C) or the temperature measurement point required by the user.



Figure 9-1 Illustration of body temperature measurement parts

Temperature measurement settings

(1) If a disposable temperature probe is being used, the temperature cable must be inserted into the socket, and then the probe and cable must be connected. For a reusable temperature probe, you plug it directly into the socket.

(2)Place the temperature probe close to the skin, and the value will be accurate after 10 minutes of heat conduction in the skin or cavity.

WARNING

•Before starting monitoring, check whether the probe cable is normal. Unplug the temperature probe cable from the jack, the screen will display the error message "Body temperature sensor is off" and an alarm sound will sound.

Motice

•Disposable temperature probes can only be used once.

•During the startup process of the monitor, the temperature measurement circuit will automatically self-check and calibrate once, which will not affect the normal operation of temperature monitoring.

9.2 Body temperature menu

The user can use the knob to move the cursor to the TEMP hot key in the parameter area on the main screen and press the knob to enter the TEMP setting menu, as shown in Figure 9-1.



Figure 9-1 TEMP setting menu

Alarm switch

Select "On" to prompt and save the alarm when TEMP (body temperature) alarms. Select "Off" to not alarm and prompt "^(A).

T1 alarm is based on the set high limit and low limit. When the temperature exceeds the

high limit or is lower than T1, it represents the temperature of channel 1.

The adjustment range of the upper and lower alarm limits is as follows:

Parameter	Maximum limit	Minimum floor	Single adjustment
			amount
T1, T2	50	0	0.1

Temperature unit

Select Celsius °C or Fahrenheit °F.

Default setting

Please refer to the "ECG Default Settings" section in "ECG/TEMP Monitoring".

9.3 Body temperature alarm information and prompt information

When the alarm recording switch in the relevant menu is turned on, those physiological alarms caused by parameters exceeding the alarm limit will trigger the recorder to automatically output alarm parameter values and related measurement waveforms. Physiological alarms, technical alarms and prompt messages that may occur during TEMP measurement are listed in the table below.

Physiological alarm:

Prompt message	Cause	Alarm level	
T1/T2, too high	The body temperature measurement value	User selectable	
	is higher than the set alarm high limit		
T1/T2, too low	The body temperature measurement value	User selectable	
	is lower than the set alarm low limit		

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Technical alarm:

Prompt message	Reason	Alarm level	Countermeasures
TEMP sensor falls	Temperature probe detaches	Low	Make sure the probe connection is reliable
TEMP alarm limit error	Functional safety failure	High	Stop using the TEMP alarm function and notify biomedical engineers or our company's maintenance personnel.

Prompt information:

Prompt message	Cause	Alarm level
TEMD magginger out of	Body temperature	
hourda	measurement value is outside	High
bounds	the measurement range	

Chapter 10 Non-Invasive Blood Pressure

(NIBP)

10.1 Instructions for non-invasive blood pressure monitoring

- Non-invasive blood pressure (NIBP) measurement uses the oscillation method and is suitable for adults and children.
- The blood pressure value measured by the monitor is equivalent to the value measured by auscultation or invasive methods.
- The clinical significance of NIBP measurements must be determined by the physician.

Non-invasive blood pressure measurement mode:

Manual, automatic and continuous measurements. Each mode displays systolic, mean and diastolic blood pressure.

(1) Manual mode: only perform one measurement.

(2) Automatic mode: Measurement is repeated. The interval can be set to 1/2/3/4/5/10/15/30/60/90/120/180/240/480 minutes.

(3) Continuous mode: Measure continuously within five minutes.

NIBP measurement will exert a certain pressure on the patient through cuff inflati on. Doctors should determine whether the patient is suitable for NIBP measureme nt based on the patient's actual condition.

Motice

•Non-invasive blood pressure measurements should not be performed on patients with sickle cell disease and anyone with skin lesions or anticipated lesions.

•For patients with severe coagulation disorders or severe thrombosis, the decision whether to perform automatic blood pressure measurement should be based on clinical evaluation, because there is a risk of hematoma at the friction point between the limb and the cuff.

•When taking measurements on pediatric patients, it is important to ensure that the correct mode setting is selected (see Patient Information Menu Settings). Using the wrong patient model has the potential to compromise patient safety, for example higher adult blood pressure levels are inappropriate for children.

•Do not perform NIBP measurements on a limb with intravenous infusion or
cannulation, as infusion slowdown or blockage may occur during measurement inflation, which may result in damage to tissue surrounding the measurement site.

•Continuous cuff pressure caused by bends in the connecting tubing can lead to blood flow disruption and harm to the patient.

•Do not take blood pressure measurements too frequently as this may cause injury to the patient due to interference with blood flow.

•Cuff application and pressurization on any limb undergoing endovascular intervention or treatment, or arteriovenous (AV) shunting, as this temporarily interferes with blood flow and can result in patient injury.

•Cuff application and compression on the arm on the mastectomy or lymphadenectomy side.

•Cuff pressurization can cause temporary loss of function of monitoring ME equipment used simultaneously on the same limb..

10.2 Non-invasive blood pressure monitoring operation methods

10.2.1 Non-invasive blood pressure measurement

The inflatable tube connecting the blood pressure cuff and monitor should be smooth and not tangled.

(1) Insert the inflation tube into the blood pressure cuff interface of the monitor and turn on the product power.

(2) Follow the method below to tie a blood pressure cuff on the patient's upper arm or thigh, as shown in Figure 10-1.



Figure 10-1 Using the cuff

a. Make sure the cuff is completely deflated.

b. Use an appropriately sized cuff on the patient, ensuring that the ϕ mark is exactly on the appropriate artery. Make sure the cuff

Do not wrap the limb too tightly, otherwise it may cause discoloration or even ischemia in the distal limb.

Motice

•The cuff width should be 40% of the limb circumference. (50% for children), or 2/3 of the length of the upper arm. The inflated portion of the cuff should be long enough to surround 50 to 80% of the limb. Choosing an incorrectly sized cuff can produce erroneous readings, such as a cuff that is too small. Use a larger cuff to reduce the error.

Adult/Pediatric Reusable Cuff Sizes

patient type	limb circumference	cuff width	Inflatable tube length
child	18~26cm	10.6cm	
aldult	25~35cm	14cm	1.5m~3m
Adult (obese)	33~47cm	17cm	

Please check to make sure that the edge of the cuff falls within the range of the "<->" symbol, otherwise, please change to a more suitable cuff.

(3) Connect the cuff to the inflation tube. The limb used for manometry should be placed at the same level as the patient's heart. If this is not possible, the following correction methods must be used to correct the measurement results:

a. If the cuff is higher than the level of the heart, 0.75mmHg (0.10kPa) should be added to the displayed value for each centimeter difference.

b. If the cuff is lower than the level of the heart, 0.75mmHg (0.10kPa) should be subtracted from the displayed value for each centimeter difference.

(4) Confirm whether the monitoring mode is correct (the monitoring mode is displayed in the monitor information area, to the right of the bed number). If you need to change the monitoring mode, please enter the "Patient Information Settings" in the "System Menu" and change the "Patient Type".

(5) Select the measurement mode in the NIBP menu. For specific methods, see the "Operation Tips" below.

(6) Press the "blood pressure" (start) button on the front panel to start pressure measurement. **Operation tips**

1) Carry out an automatic measurement

Enter the "NIBP Settings" menu and select the "Interval Time" item. The user can select the time interval value for automatic measurement. After that, press the "NIBP" button on the front panel, and the system will automatically perform inflation measurements according to the set interval.

ANotice

•If the non-invasive pressure measurement time in automatic mode is set too long,

purpura, ischemia and nerve damage may occur in the limbs in contact with the cuff.

While monitoring the patient, frequently check the color, warmth, and sensitivity of the

distal extremities. Once any abnormality is observed, change the cuff placement or stop

blood pressure measurement immediately.

•After the short-term automatic mode measurement is completed, the device should

return to long-term automatic mode or manual mode. The operator can also intentionally activate short-term automatic mode again.

2) Stop automatic measurement

Pressing the "NIBP" key at any time during the automatic measurement process will stop the automatic measurement.

3) Carry out a manual measurement

a. Enter the "NIBP Settings" menu, select the "Interval" item, set the value to "Manual", and then press the "Blood Pressure" key on the front panel to start a manual measurement.

b. During the idle time of automatic measurement, press the "NIBP" key to start a manual measurement. If at this time

Press the "NIBP" key again to stop manual measurement and continue automatic measurement.

4) Perform a manual measurement during the automatic measurement process

Press the "NIBP" button on the control panel.

5) Stop a manual measurement midway

Press the "NIBP" button on the control panel again.

6) Make continuous measurements

Enter the "NIBP Settings" menu and select the "Continuous Measurement" item to start continuous measurement. This process will last for 5 minutes.

7) Stop continuous measurement midway

The continuous measurement can be stopped at any time during the continuous measurement by pressing the "NIBP" button on the control panel.

Motice

•If you have any doubts about the product's measurement parameters, please use other equipment or methods to check the patient's vital signs before checking whether the monitor is functioning properly.

Measurement limitations

Because of the diversity of patient conditions, oscillometric measurement has limitations.

This measurement monitors the regular pulse waveform generated by arterial pressure. Under this method, when the patient's condition makes monitoring difficult, the measured parameters become unreliable and the measurement time increases. Users should be aware that the following conditions may interfere with measurements. In some cases, a patient's condition makes monitoring impossible:

(1) Patient movement

If the patient is moving, shaking or spasming, monitoring will be unreliable or even impossible because these conditions may interfere with the detection of arterial pressure pulses and the pressure measurement time will be extended.

(2) Arrhythmia

If the patient shows arrhythmia resulting in irregular heartbeats, monitoring will be unreliable or even impossible, and the pressure measurement time will be extended.

(3) Heart-lung machine

If the patient is connected to an artificial heart-lung machine, monitoring will not be possible.

(4) Pressure changes

If arterial pressure pulsations are being analyzed to obtain measurements at a time when the patient's blood pressure changes rapidly,

Monitoring will be unreliable or even impossible.

(5) Severe shock

Monitoring will be unreliable if the patient is in severe shock or hypothermic. Because reduced blood flow to the periphery results in reduced arterial pulsation.

(6) Extreme heart rate

Blood pressure monitoring cannot be performed when the heart rate is lower than 40bpm (beats per minute) and higher than 240bpm (beats per minute).

Non-invasive blood pressure parameter setting and adjustment

The measurement results and corresponding information of NIBP are displayed on the screen as follows:



Non-Invasive Blood Pressure Menu

Turn the knob, move the cursor to the NIBP hot key in the parameter area on the screen, and then press the knob to enter "NIBP

"Settings" menu, as shown in Figure 10-2.

NIBP Setup	Х		
Alarm On/Off	ON 🗘		
Unit	mmHg 🗘		
Interval Time	Manua I 🗘		
Inflation	160 \$		
	Reset		
Continue			
Calibration			
Pneumatic			
Default >>			

Figure 10-2 NIBP setting menu

Alarm switch

Select "On" to prompt and store the alarm when the pressure alarm occurs. Select "Off" to

not alarm and prompt next to NIBP in the parameter area of the screen. A. The pressure alarm is based on the set high limit and low limit. When the pressure exceeds the high limit or is lower than the low limit, an alarm is issued. Alarm settings can be set separately for systolic blood pressure, mean blood pressure and diastolic blood pressure. The adjustment range of the upper and lower alarm limits is as follows:

Blood pressure	Adult(mmHg)	Children (mmHg)
Systolic blood pressure alarm high limit	42–260	42–230
Systolic blood pressure alarm low limit	40–258	40–228
Average pressure alarm high limit	22–240	22–170
Average pressure alarm low limit	20–238	20–168
Diastolic blood pressure alarm high limit	12–220	12–160
Diastolic blood pressure alarm low limit	10–218	10–158

Pressure unit

Optional mmHg or kPa.

Intervals

Automatic measurement interval (unit: minutes). You can choose from 1, 2, 3, 4, 5, 10, 15,

30, 60, 90,120, 180, 240, 480 minutes. After selecting the interval, the prompt "Please press the 'blood pressure' (start) key" will appear in the NIBP prompt area. This is the first automatic measurement of inflation by pressing the "NIBP" key. To end automatic measurement, select "Manual" during the measurement interval to return to manual mode.

Pre-inflation value

Default	Default	Manually selectable pre-charge value in NIBP	
Default	pre-inflation value	menu	
configuration	(mmHg/kPa)	(mmHg/kPa)	
Adult default	160	80~ 250	
configuration	100	80,~230	
Child default	140	80~ 200	
configuration	140	80,~200	

Press this key to select the initial pressure value for the next cuff inflation. Under different default configurations, there are different pre-inflation value selection ranges, as shown in the following table:

After the you press the "Menu" button on the front shell, they enter the "Default Configuration" menu in the "System Menu". After confirming the default configuration, please return to the main interface and select the NIBP menu hotkey in the NIBP parameter area to enter "NIBP Settings". It can be seen that the initial value corresponding to the " pre-inflation value " is the initial inflation pressure value corresponding to the selected default configuration, as shown in the table above. Move the cursor to the "pre-inflation value" option and press to see the range of pre-inflation value that can be manually adjusted, as shown in the table above.

ANotice

•The "Pre-inflation value" option is to help the user select the next cuff inflation

pressure, but the pre-inflation value for subsequent measurements will be based on the

last systolic blood pressure measurement of the same patient. The system memorizing

this value can shorten the measurement time of the same patient and increase the

accuracy of the measurement.

Measurement mode

You can choose "automatic measurement" or "manual measurement".

Reset

(1) The measurement status of the blood pressure pump can be reset.

(2) Press this key to restore the inflation value of the blood pressure pump to its initial setting.

(3) It is recommended to use this key when the blood pressure pump is not working properly but the monitor cannot indicate the cause of the problem. This allows the blood pressure pump to perform a self-check, allowing it to automatically recover if the pump is not working abnormally due to unexpected reasons.

WARNING

•Calibration of NIBP measurements should be performed every two years (or as per

your hospital's maintenance schedule). Its performance should be checked in the

following details.

Air leak detection

It is used to detect whether the NIBP measurement pump is leaking. When the NIBP cuff is connected, you can use this key to start the NIBP inflation process to find out whether the sealing condition of the NIBP air path is good. If the air leak test passes, the system will not give any prompt; if it fails, there will be a corresponding error prompt in the NIBP information area.

AWARNING

•This air leakage detection is different from what is described in the EN 1060-1 standard and is for users to simply detect whether there is air leakage when NIBP is inflated. If the system shows NIBP air leakage at the end of the test, please contact our company's maintenance engineer.

Air leakage detection process:

1) Connect the cuff to the NIBP air hole of the monitor.

2) Wrap the cuff around a cylinder of appropriate size.

3) Enter the "NIBP Settings" menu.

4) Turn the knob, move the cursor to the "Air Leak Detection" item, and press the knob. At this time, "Air leakage detection..." will be displayed below the NIBP parameter area on the screen, indicating that the system has started to perform air leakage detection.

5) The system automatically inflates to a pressure of 180mmHg.

6) After about 20 seconds, the system will automatically open the air release valve to indicate that the air leakage measurement is completed.

7) If there is no prompt message in the NIBP parameter area, it means that there is no air leakage in the system. If "Pump leaks..." is displayed, it means there may be a leak in the air circuit. At this time, the operator should check whether the entire connection is loose. After confirming that the connection is correct, perform the air leakage test again. If the fault message still appears, please contact the manufacturer for repair.



Figure 10-4 NIBP air leakage detection connection diagram

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Default configuration

Select this item to enter the NIBP default configuration dialog box, where you can select the system default configuration. In the dialog box, the user has two options to choose from, namely "Factory Default Configuration" and "User Default Configuration". After selecting any item and exiting the dialog box, the system will pop up a dialog box asking the user to confirm their selection.

Motice

•When the user only sets the "Patient Type" in the "Patient Information Settings" without making any selection in the "Default Configuration", the system will initially set the relevant module parameters according to the "Patient Type". And changes to the default type settings in "Default Configuration" will also change the "Patient Type" in "Patient Information Settings".

10.4 NIBP alarm information and prompt information

Physiological alarms caused by parameters exceeding the alarm limit may trigger the recorder to automatically output parameters and related measurement waveforms at the time of alarm occurrence, provided that the alarm recording switch in the relevant menu is turned on. Physiological alarms, technical alarms and prompt messages that may occur during NIBP measurement are listed in the following table:

Prompt message	Cause	Alarm level
NS is too high	NIBP systolic blood pressure measurement is	User selectable
or too low	above or below the set alarm limit	
ND is too high	NIBP diastolic blood pressure measurement is	User selectable
or too low	above or below the set alarm limit	
NM is too high	NIBP average pressure measurement value is	User selectable
or too low	higher or lower than the set alarm limit	

Physiological alarm:

Technical alarm 1 (displayed in the prompt area below the NIBP pressure value):

Prompt message	reason	Alarm level	Countermeasures
NIBP self-test error	Sensor or other hardware error in NIBP measurement module	High	Stop using the NIBP measurement function and notify biomedical engineers or our company's maintenance personnel.
NIBP communication error	Communication with NIBP measurement module failed	High	If the fault persists, stop using the NIBP function and notify biomedical engineers or our company's maintenance personnel.
Cuff is too loose or not connected	The cuff is not tied properly or there is	Low	Tie the cuff.



	no cuff		
Cuff inflation tube leaking	Damaged cuff, hose or connector	Low	Check and replace leaking parts, and if necessary, notify biomedical engineers or our company's maintenance personnel.
Air pressure error	Unable to obtain stable pressure value, such as hose tangles	Low	Check whether the hose is tangled. If the fault persists, notify the biomedical engineer or our company's maintenance personnel.
Signal too weak	The cuff is too loose or the patient's pulse is too weak	Low	Use other methods to measure blood pressure.
Pressure out of range	The measurement range exceeds the specified upper limit	High	Reset the NIBP measurement module. If the fault persists, stop using the NIBP measurement function and notify biomedical engineers or our company's maintenance personnel.
Arm movement	Affected by arm movement, too much signal noise or irregular pulse rate	Low	Ensure that the patient being tested is quiet and does not move.
Overvoltage protection	The pressure exceeds the specified safety limit	High	Measure again. If the fault persists, stop using the NIBP measurement function and notify the biomedical engineer or our company's maintenance personnel.
Signal saturation	large movement	Low	Do not allow the patient to exercise.
Pump leaking	Leak found during air leakage test	Low	Check and replace leaking parts, and if necessary, notify biomedical engineers or our company's maintenance personnel.
NIBP system error	Blood pressure pump system malfunction	High	Stop using the NIBP measurement function and notify biomedical engineers or our company's maintenance personnel.
Wrong cuff type	Cuff type does not match patient type	Low	Use appropriate cuffs.
Measurement timeout	Measurement time exceeds 120 seconds (adult/child)	High	Measure again or use another pressure measurement method.
NIBP reset error	Module reset is abnormal	High	Use the reset function again.
Measurement error	The system cannot perform	High	Check the cuff, make sure the patient is not moving during

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measurement	monitoring, and measure again.
analysis or	
calculations while	
a measurement is	
being taken.	

Prompt information (displayed in the prompt area below the NIBP pressure value):

Prompt message	Cause	Alarm level
Manual	During manual measurement	
measurement		
Continuous	During continuous measurement	
measurement		
Automatic	During automatic measurement	
measurement		
Please press the start	After selecting the measurement interval	
button	in the menu	
Measurement	During the measurement process, press	
terminated	the start key to stop the measurement.	No alarm
Calibration	Calibrating	
Calibration	The calibration process has ended	
terminated		
Air leak detection	Air leak detection in progress	
Air leak detection	Air leak detection completed	
completed		
Module reset	Reset the NIBP module after loading	
Manual reset	During NIBP reset (user trigger)	
Reset error	Reset action failed	

Chapter 11 End-Tidal Carbon Dioxide (CO2)

11.1 CO2 monitoring instructions (This function is optional, and the

instrument you are using may not be equipped with this function)

This module measures the carbon dioxide pressure of the patient's airway and can obtain the end-tidal carbon dioxide content (EtCO2), the minimum amount of carbon dioxide inhaled (InsCO2) and the airway respiratory rate, and display the CO2 pressure waveform. Equipped with automatic atmospheric pressure compensation function.

Under the conditions of indoor temperature of 23 $^{\circ}$ C, sampling gas temperature of 37 $^{\circ}$ C, and sampling relative humidity of 100%, the maximum time interval for operators to intervene in the water vapor treatment system is 120 hours, the maximum gas sampling flow rate is 60ml/min, and the minimum gas sampling flow rate is 38ml/min.

11.1.1 CO2 measurement settings:

- 1) Confirm the configured CO2 module type (mainstream or sidestream).
- 2) Connect the module to the CO2 module socket of the monitor.
- 3) Make connection settings according to different types

Mainstream CO2 settings

First insert the adapter into the mainstream CO2 module, and then string both ends of the adapter into the gas line, as shown in Figure 11-2 below.

Sidestream CO2 settings:

First, insert the dryer end of the sampling tube into the air inlet of the sidestream CO2 module. Then, according to usage needs, connect the air inlet of the sampling tube to the L-shaped tee, and finally string the tee into the gas path, as shown in Figure 11-3. If using a nasal tube to directly monitor and test patients, first insert the dryer end of the sampling tube into the inlet of the bypass CO2 module, then connect the screw end of the nasal tube to the sampling tube, and connect the other end to the patient's nostril. Adjust the tightening clip of the sampling tube to the appropriate position to prevent it from falling off. As shown in Figure 11-4.

4) The CO2 module of this device defaults to measurement mode. After inserting the module, a technical prompt of "CO2 preheating" will appear on the display until the sensor reaches the operating temperature. The preheating time is 5 minutes.

5) The values tested during the preheating process are not included in the reference. After the preheating is completed, effective and accurate measurement of CO2 begins.















diagram



Figure 11-4 Nasal oxygen tube connection
diagram

The respiratory rate can be adjusted from 0 to 150 with a step size of 5 using a respiratory rhythm fixture. The gas outlet can be connected to a mainstream adapter or a bypass sampling tube to adjust the respiratory rate setting. Different concentrations of standard gases can be changed, and the accuracy of the measured gas concentration can meet the requirement of \pm (0.43%+8% of gas concentration).

MNotice

•This equipment needs to be zeroed when the place of use or environment changes. Before zeroing, remove the sampling tube from the air connection and place it in an air environment 0.5 meters away from the original sampling port and the patient, and wait for each CO2 display value. After reaching zero, perform the zero calibration operation again.

•When the shunt RGM cannot maintain the normal flow rate, it will alarm the block.

WARNING

•Collision and vibration of the CO2 module should be avoided as much as possible.

•If the sampled gas is returned to the respiratory system, there is a risk of patient cross-infection.

•This equipment should not be used with gas supplied from an oxygen concentrator.

∕∆Careful

•Do not use the instrument in an environment containing flammable anesthetic gases.

•The instrument can only be operated by professionals who have received professional training and are familiar with this manual.

ANotice

•If the package has been opened or the accessory inside is damaged, please do not use the accessory and return it to the supplier.

•When the information "CO₂ Warming" and "CO₂ Sensor Warming" are displayed on the screen, it means that the sensor is being warmed up and started. When this message disappears from the screen, standard measurements can be taken.

•Side-flow sampling tubes and sinks are disposable items and cannot be re-sterilized and used by patients.

•If other harmful gases are used during medical treatment, users should pay attention to environmental protection issues.

11.2 CO2 Menu

ECG setup menu

Parameter setting and adjustment

Turn the knob, move the cursor to the CO_2 hot key in the parameter area on the screen, and press the knob to enter the " CO_2 Settings" menu.



Figure 11-5CO2 settings

CO2 alarm settings

(1) Alarm switch: "On" will prompt and store the alarm when there is an alarm in the CO2 parameter, "Off" will not alarm.

(2) Alarm level: There are three options, namely "high", "medium" and "low". "High" indicates the most serious alarm, followed by "medium" and "low" in severity. The change of "Alarm Level" only affects the physiological alarm level of carbon dioxide parameters (including EtCO2 upper limit, EtCO2 lower limit, InsCO2 upper limit, AwRR upper limit and AwRR lower limit). The default alarm level is "Medium".

(3) CO2 alarm high limit: used to adjust the alarm upper limit of EtCO2. When the measured value is greater than the upper alarm limit of CO2, the message "CO2 too high" appears on the screen. This message disappears when the measured value returns to normal.

(4) CO2 alarm lower limit: used to adjust the lower alarm limit of EtCO2. When the measured value is less than the lower alarm limit of CO2, the message "CO2 too low" appears on the screen. This message disappears when the measured value returns to normal.

(5) INS alarm high limit: used to adjust the alarm upper limit of InsCO2. When the

measured value is greater than the upper alarm limit of InsCO2, the message "INS too high" appears on the screen. This message disappears when the measured value returns to normal.

(6) AWRR alarm high limit: used to adjust the alarm upper limit of AwRR. When the measured value is greater than the upper alarm limit of AwRR, the message "RR too high" appears on the screen. This message disappears when the measured value returns to normal.

(7) AWRR alarm lower limit: used to adjust the alarm lower limit of AwRR. When the measured value is greater than the lower alarm limit of AwRR, the message "RR too low" appears on the screen. This message disappears when the measured value returns to normal.

Waveform speed

Used to adjust the display speed of CO2 waveform. The available options are "6.25 mm/s", "12.5 mm/s", and "25.0 mm/s".

Pressure unit

Used to change the display unit of CO2 and InsCO2 parameters. You can select "mmHg" or "kPa".

Operating mode

Used to change the working mode of carbon dioxide, you can choose "measurement" mode or "standby" mode. The default is "standby" mode. When CO2 monitoring is required, select the "Measurement" mode. "Standby" mode will turn off the air pump of the side flow module, the sensor of the mainstream module and the infrared IR source, which can reduce power consumption and extend the service life of the IR source and the entire CO2 module.

Default setting

Select this item to enter the "CO2 Default Settings" dialog box. The user can select the "Manufacturer Default Settings" or "User Default Settings" item. After selecting to exit, the system will pop up a dialog box asking the user to confirm their choice.

EtCO2 alarm upper limit: When the parameter value is higher than this limit, an upper limit alarm will be issued.

Default values: Adult: 50 mmHg Pediatric: 50 mmHg Neonate: 45 mmHg EtCO2 alarm lower limit: When the parameter value is lower than this limit, an over-lower limit alarm will be issued.

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Default values: Adult: 15 mmHg Pediatric: 20 mmHg Neonate: 30 mmHg InsCO2 alarm upper limit: When the parameter value is higher than this limit, an upper limit alarm will be issued. Default values: Adult: 4 mmHg Pediatric: 4 mmHg Neonate: 4 mmHg AwRR alarm upper limit: When the parameter value is higher than this limit, an upper limit alarm will be issued. Default values: Adult: 30 rpm Pediatric: 30 rpm Neonatal: 100 rpm AwRR alarm lower limit: When the parameter value is lower than this limit, an over-lower limit alarm will be issued. Default value: Adult: 8 rpm Pediatric: 8 rpm Newborn: 30 rpm Asphyxiation time: Options are 10S-40S Default value: 20S Operation mode: Mainstream: standby, measurement. Side flow: on standby, measuring. Default mode: standby Default mode: normal Pumping rate: 100 – 200 ml/min Default pumping rate: 100 ml/min Unit: mmHg/kPa Default: mmHg Waveform speed: 25.0/12.5/6.25 (mm/s) Default: 25.0 mm/s Waveform amplitude: low/high Default: low In addition, please refer to the "Alarm Function" chapter for the alarm function of the CO2 module, and see the "Recording Function" chapter for its recording function. For the alarm event review, trend graph and trend table related to CO2 parameters, see the

"Trends and Events" chapter.

11.3 CO2 alarm information and prompt information

When the alarm recording switch in the relevant menu is turned on, those physiological alarms caused by parameters exceeding the alarm limit will trigger the recorder to automatically output alarm parameter values and related measurement waveforms. The physiological alarms, technical alarms and prompt messages that may occur during CO2 module measurement are listed in the following table:

Prompt message	Cause	Alarm level
CO2 is too high	EtCO2 measured value is higher than the set alarm	User selectable
	high limit	
CO2 is too low	EtCO2 measured value is lower than the set alarm	User selectable
	low limit	
INS is too high	InsCO2 measured value is higher than the set	User selectable

Physiological alarm:

	alarm limit	
AWRR is too	AwRR measurement value is higher than the set	User selectable
high	alarm high limit	
AWRR is too	The AwRR measurement value is lower than the	User selectable
low	set alarm low limit	
CO2 breathing	Respiratory arrest(No breathing detected within	High
asphyxia	the set delay time)	

ANotice

- The interference gas of the gas concentration provided by this device has no impact on the quantification of gas readings and is not shared with anesthetic gases.
- The following factors may have known adverse effects on the claimed performance:
 - a. Quantitative effects of sampling gas humidity or condensation;
 - b. Leakage of sampled gas or internal exhaust;
 - c. Periodic pressure rising to 10kPa (100cmH2O);
 - d. Other sources of interference.

Chapter 11 Common Troubleshooting

Methods

Fault name	Fault description	Possible causes of failure	Solution
	Turn on but	1. The screen wire is loose;	1. Re-plug the cable plug;
Black screen	black screen appears	2. The display or KC7 motherboard is damaged.	2. Replace the display or motherboard;
	T	1. The screen wire is loose;	1. Re-fix the display cable;
White screen	Turns on but displays white screen	2. The LCD screen is damaged;	2. Replace the LCD screen;
		3. The motherboard is damaged;	3. Replace the motherboard;
Can't enter the monitoring interface	Boot and stop at LOGO or other location	1. The core board is damaged;	1. Replace the core board;
		1. Blood pressure cuff or trachea leaks;	1. The cuff selection is wrong, replace the blood pressure cuff or related connections;
Abnormal Blood blood pressure pressure measurement measuremen error with no results	Abnormal blood pressure	2. Overvoltage protection	2. Patient type and cuff selection are uncomfortable or the extension tube is bent and blocked.
	with no results	3. Large deviation in measurement values	3. The cuff is too tight or too loose, the measurement position is wrongly selected, excessive exercise, etc.
		4. Wrong air pressure	4. Replace the slow-release valve or other air-blocking parts;
ECG function not	No ECG waveform	1. ECG clutter or instability	1. The patient is noisy, the ECG electrode attachment points are not cleaned, and the electrode pads are not bonded well, etc.
working		2. The ECG lead 3/5 is wrongly selected or the ECG electrodes are not connected properly or the lead wire is	2. Select the correct lead method according to the ECG cable; attach the ECG electrodes and

		not inserted in place.	connect the lead wires.
		3. The ECG lead falls off or the wire is broken;	3. Reconnect the ECG leads or replace the lead wires;
		4. ECG clutter;	4. There is too much interference from peripheral electronic equipment;
Blood Unable to		1. The blood oxygen probe falls off;	1. Connect the blood oxygen probe to the patient's finger;
oxygen is not working properly saturation	2. The blood oxygen probe is damaged;	2. Replace the blood oxygen probe.	
	3. The blood oxygen probe plug is not inserted in place.	3. Plug the connection tightly into place	
The end-tidal	Unable to	1. The sampling line is kinked.	1. Reconnect the carbon dioxide sampling line.
carbon dioxide module is	measure end-tidal carbon	2. The drain trap is blocked.	2. Replace the water trap with a new one
not working dioxide properly.	3. The sampling line has been blocked.	3. Replace with a new sampling line.	
Pottom, not	Short bottom	1. The battery is damaged and the voltage is very low;	1. Replace the battery;
charging	Short battery life	2. The power supply is damaged and the charging power is low;	2. Replace the power board;

MNotice: Non-professionals, please do not disassemble the phone! If the fault exceeds the scope described in the above bad information or cannot be solved according to the above solutions, please contact our company's after-sales service department directly!

Chapter 12 Accessories

The supporting accessories recommended by our company are as follows:

14.1 Electrocardiogram (ECG) accessories

Name	Model	Material Number	Patient category	Illustrate	Manufacturer	Test Options
				Anti defibrillation,	Shenzhen	
	SG5143S 12.02.0051C	1	push button,	SINO-K Medical	V	
	5051155	12.02.00510	,	five-lead, 2.9m,	Technology	`
ECG				reusable	Co.,Ltd.	
cable				Anti defibrillation,	Shenzhen	
		,	push button,	SINO-K Medical		
	5051455	12.02.0037C	/	three-lead, 2.9m,	Technology	v
				reusable	Co.,Ltd.	

14.2 Blood oxygen (SPO2) accessories

Name	Model	Material Number	Patient category	Illustrate	Manufacturer	Test Options
	SP9325A	12.02.0063C	Adult	Finger Clip Sensor with 300cm Cable , reusable	Shenzhen SINO-K Medical Technology Co.,Ltd.	V
SP8325A	12.02.0058C	Adult	FingerSleeveSensor with 300cmCable , reusable	Shenzhen SINO-K Medical Technology Co.,Ltd.	\checkmark	
SpO2 Probe	SP7119A	12.02.0044C	Child	Finger Clip Sensor with 300cm Cable , reusable	Shenzhen SINO-K Medical Technology Co.,Ltd.	
	SP6119A	12.02.0045C	Child	FingerSleeveSensor with 300cmCable , reusable	Shenzhen SINO-K Medical Technology Co.,Ltd.	

Test Material Patient Name Model Illustrate Manufacturer Options Number category Shenzhen SINO-K Body cavity Medical Technology $\sqrt{}$ ST2305 12.02.0056C / sensor.3m.reusable Co.,Ltd 体温探头 Shenzhen SINO-K Body surface Medical Technology $\sqrt{}$ ST1305 12.02.0055C / sensor.3m.reusable Co.,Ltd

14.3 Temperature (TEMP) accessories

14.4 Non-Invasive Blood Pressure (NIBP) Accessories

Name	Model	Material Number	Patient category	Illustrate	Manufacturer	Test Options
NIBP	SH0908 S	12. 02. 0025C	1	Strait plug connector to pagoda connector,2.5m	Shenzhen SINO-K Medical Technology Co.,Ltd.	\checkmark
extender SH0910 S	12. 01. 0012C	1	Strait plug connector to spring joint,2.5m	Shenzhen SINO-K Medical Technology Co.,Ltd.	\checkmark	
NIBP Cuff	SC2711	12.01.0002C	Adults	Brown color, reuseable	Shenzhen SINO-K Medical Technology Co.,Ltd.	\checkmark
	SC2611	12.01.0017C	Pediatr ics	Brown color, reuseable	Shenzhen SINO-K Medical Technology	V

Co.,Ltd.	Colld
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14.5 CO2 Accessories

Name	Model	Material Number	Patient category	Illustrate	Manufacturer	Test Options
CO2 Sensor	TL1000A	12.01.0063C	/	main stream	Suzhou Troline Technology Co., Ltd.	\checkmark
	TL2000A	12.01.0064C	/	external side-strea m	Suzhou Troline Technology Co., Ltd.	\checkmark
Airway adapter	TL_Madapter	12. 02. 0061C	/	Airway adapter	Suzhou Troline Technology Co., Ltd.	\checkmark
	TL_Sadapter	12.01.0044C	/	Airway adapter	Suzhou Troline Technology Co., Ltd.	\checkmark
Sampling tube	TL_Sampline	12.01.0049C	/	externa l side-st ream	Suzhou Troline Technology Co., Ltd.	\checkmark

15.5 Othe r acces sories	Model	Material Number	Patient category	Illustrate	Manufacturer	Test Options
Na me						
7.4V4.8	ICR21700	02.10.0006C	/	SMP-02V(JST)	Shenzhen	\checkmark

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AH		connector	Xinlong Ding	
lithium		35.52wh	Technology Ltd.	
battery				

WARNING

•Please use recommended accessories. Using other accessories may damage the monitor or fail to achieve the accuracy of each parameter specification claimed in the manual.

•The listed accessories must be used in conjunction with our company's equipment. Before using the accessories, please read the instructions of the equipment and accessories to confirm the compatibility of the accessories with the product. Otherwise, the patient may be injured or the measurement accuracy may be affected.

•Disposable accessories can only be used once and reuse can lead to cross-contamination and performance degradation.

Motice

•Pay attention to the packaging label for accessories that have a safe use period. Do not use expired accessories.

•Sterilization accessories should be packaged completely and cannot be used if they are damaged or leaking.

∕∆Careful

•The use or storage environment of accessories should meet the instructions, otherwise it may not achieve the performance claimed in its specifications.

•If you find that the accessories are expired, the packaging is damaged, or the cables are damaged, please do not use it.

•Please refer to local regulations or hospital systems for the disposal of disposable accessories, expired, and scrapped accessories.

Appendix I Product Specifications

I.1 Monitor type (classified according to GB9706.1)

Standard electric shock	Class I, equipment with internal power supply
resistance type	
EMC category	Class A
Standard level of	ECG (RESP), SPO2, TEMP: type CF;
resistance to electric	NIBP and CO2: type BF.
shock	
Degree of liquid	IPX2
prevention	
Disinfection/sterilization	See Chapter 5 for details
methods	-
Way of working	continuously working
resistance to electric shock Degree of liquid prevention Disinfection/sterilization methods Way of working	NIBP and CO2: type BF. IPX2 See Chapter 5 for details continuously working

I.2 Monitor specifications

I.2.1 Monitor dimensions and weight

See the packaging markings on the outer box of each model.

I.2.2 Working environment

Tomm onstrume Dom on	Working: 5°C~40°C				
Temperature Kange	Transportation and storage temperature: (-20~+55)°C				
Uumidity Dance	Working: 15%~85% (no condensation)				
Humany Kange	Transportation and storage humidity: ≤93% (no condensation)				
	Working: Not more than 3,000) meters (9843) feet				
Altitude Range	Transportation and storage altitude: (-500-13,100) meters				
	(-1,600-43,000) feet				
	a.c.100V-a.c.240V, 50Hz/60Hz				
Electrical	Input power: <45VA				
Specifications	Fuse: T1.6AL 250V d.c.15V/2A (7~15V)				
	Breaking capacity >1.5A, over 1.6A breaking				

I.2.3 Display information

(1) Color LCD monitor, 7 inches (1024*600 pixels).

(2) Up to 7 waveform displays and one alarm indicator light (red/yellow/blue).

(3) Three sound alarm modes corresponding to the alarm status

- I.2.4 Battery
- (1)7.4V/4800m AhLithium Ion Battery.
- (2) The working time of full power in normal state is more than 150 minutes.

(3) After the first low battery alarm, it can continue to work for 10 minutes, and then it

will automatically shut down.

(4) The battery charging time is not more than 5 hours, and the AC indicator will be steady on when charging.

(5) The charging time for charging the battery from depleted to 90% capacity is 5 Hour.

(6) Expected battery life: 8 years.

I.2.5 Review

	Short trend: 2 hours, resolution 1 second or 5 seconds			
Trend review	Long-term trend: 72 hours, resolution 1 minute, 5 minutes or 10			
	minutes plus USB flash drive 15 days			
NIBP				
measurement	Review of 1000 NIBP measurement data			
review				
Alarm event	200 groups up to 1000 groups with USB disk			
review	200 groups, up to 1000 groups with OSB disk			
Waveform review	1 hour, up to 24 hours with a USB flash drive			

I.3 ECG specifications

Standards	GB0706 227_2021
Standards	0B)/00.227-2021
compliant	
	Lead mode: three-lead, five-lead
	Standard three-lead cable: RA, LA, LL
Lead configuration	Standard five-lead cable: RA, LA, LL, RL, V
	Three-lead lead mode: I, II, III
	Five-lead lead mode: I, II, III, aVR, aVL, aVF
	0.25 (2.5mm/mV), 0.5 (5mm/mV), 1 (10mm/mV), 2 (20mm/mV),
Gain	automatic
	The errors are all less than±5%
Heart rate	Adult: 15 ~ 300bpm (beats/minute)
measurement range	Children: 15 ~ 350 bpm (beats/minute)
Accuracy $\pm 1\%$ or ± 1 bpm, whichever is greater; resolution	
Heart rate accuracy	(beats/minute).
ECG signal range	0.2~8mV
Input resistance	>5 (megaohms)
D 1 14	Reference mode 0.05 Hz \sim 100Hz; Monitoring mode 0.5 Hz \sim 40Hz;
Bandwidth	Surgery mode 1Hz~20Hz
Common mode	Reference mode > 90 dB; Monitoring mode > 100 dB; Surgery mode
rejection ratio	> 100 dB
Electrode	
polarization voltage	±300mV
range	
Pacing pulse	For pacing pulses that meet the following conditions, it can be
detection	detected: Amplitude: 1 mV ~ 10mV: Width: 0.5ms ~ 2ms: Rise time:

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	10µs ~ 100µs
Pacing pulse suppression	When the pacing analysis switch is turned on, pacing pulses that meet the following conditions can be suppressed without affecting heart rate calculation; Amplitude: $\pm 2 \text{ mV} \sim \pm 700 \text{mV}$; Width: 0.1ms $\sim 2 \text{ms}$; Rise time: 10 μ s $\sim 100 \mu$ s
Baseline recovery time	<5 seconds after defibrillation
Calibration signal	1mV (peak to peak), accuracy±5%
Lead off detection current	<0.1µA
System noise	Not more than 30µV
	Calculate the average heart rate by the following method: if the last
Heart rate average	three consecutive RR intervals are greater than 1200 ms, average the 4 most recent RR intervals to calculate the heart rate; otherwise, take the 12 most recent RR intervals and subtract the maximum and minimum values and then average them to calculate the heart rate. The heart rate value displayed on the screen is refreshed every second.

I.4 Respiratory specifications

Measurement method	RA-LL, RA-LA impedance method		
Respiratory impedance detection range	(0.3~3)Ω		
Base impedance range	(200~4000)Ω		
Bandwidth	(0.1~2.5)Hz		
Descriptory rate range	Adult: 7bpm~120bpm		
Respiratory rate range	Children: 7bpm~150bpm		
Respiration rate resolution	1 bpm;		
Respiration rate accuracy	±2 bpm		
Suffocation alarm	10~40 seconds		

I.5 SPO2 specifications

Standards compliant	YY 9706.261-2023
Pulse oximeter measurement range	70~100%
Pulse Oximeter Accuracy	±3%
Pulse rate measurement range	25bpm~300bpm
Pulse rate accuracy	±3bpm

I.6 TEMP specifications

Standards compliant	YY 9706.256-2023
Applicable temperature sensor	YSI series, CYF series
Number of channels	2 channels
Measuring range	0°C∼50°C
Measurement accuracy	$25^{\circ}C \sim 45^{\circ}C$, error $\pm 0.2^{\circ}C$; other range accuracy is $\pm 0.4^{\circ}C$

I.7 NIBP specifications

Standards compliant	YY 9706.230-2023
Measurement method	pulse wave oscillation method
Operating mode	Manual measurement/automatic interval measurement/continuous measurement
Measurement interval time in automatic measurement mode	(1,2,3,4,5,10,15,30,60,90,120,180,240,480) minutes
Measurement time in continuous measurement mode	5 minutes
Pulse rate range	40 – 240 bpm
Measuring range	adult: Systolic blood pressure 40–260mmHg (5.3–36.0kPa) Average pressure 20–240mmHg (2.7–30.7kPa) Diastolic blood pressure 10–220mmHg (1.3–28.0kPa) child: Systolic blood pressure 40–230mmHg (5.3–30.7kPa) Average pressure 20–170mmHg (2.7-22.0kPa) Diastolic blood pressure 10-160mmHg (1.3–20.0kPa)
Measuring range	0mmHg~300mmHg
Display resolution	1mmHg (0.133kPa)
Average error	±5mmHg
Standard error	±8mmHg
Overvaltage protection	Adult: 300 mmHg±3 mmHg
overvoltage protection	Children: 280 mmHg±3 mmHg

I.8 CO2 specifications

Standards compliant	GB 9706.225-2022		
Measurement display range	0~20%(0~152 mmHg)		
Measurement accuracy	± (0.43%+8% of reading)		

Resolution		0.1%(1 mmHg)		
Airway respiratory rate measurement display range		0~160 brpm		
Airway respiratory rate n	neasurement accuracy	±1 brpm		
Airway respiratory rate n	neasurement resolution	1 brpm		
Rated respiratory rate		0-150 times/minute		
An overview of test methods for determining the rated respiratory rate range and the corresponding effects of changes in the accuracy of end-expiratory gas readings with respiratory rate [requirements in 2017929101; and i)]		The rated breathing rate is set at 30,60,100,140 times/minute, using 5% standard CO2 gas, and the accuracy of the end-expiratory gas reading is within the claimed standard rang		
The measurement error of CO2 may be affected by the following:		1. Use immediately after calibration zero. 2. Use during the warm-up period of the CO2 module. 3. Sampling gas flow rate is less than 38 ml/min. 4. When the respiratory rate is not at the claimed respiratory rate (0-140brpm). 5. The CO2 sensor is overheated. Note: I:E and airway respiration rate will not affect the measurement accuracy		
The minimum sampling flow rate that meets the measurement accuracy of the shunt RGM		38 ml/min ± 10 ml/mins		
CO2 sampling gas flow rate and tolerance		50ml±10ml		
Drift in measurement accuracy		Meet accuracy requirements within 6 hours		
Total response time of m	ainstream systems	<4seconds		
bypass system at rated	Total response time	≤5.0s, @50ml/mins		
flow rate	rise time	≪450ms,@50ml/mins		

I.9 Alarm specifications

Alarm level: high, medium and low alarm levels, consistent with YY 9706.108-2021.

Alarm type: physiological alarm, technical alarm

Alarm indication: red, yellow and blue alarm indicators.

Appendix II Summary of Performance

Requirements of GB 9706.227-2021

Requirements Description	Minimum/Maximum		Unit	Min./Max. value
Normal working conditions				
Ambient temperature	scope		°C	0~40
Relative humidity	scope		%	≤85% (no condensation)
Atmospheric pressure	scope		hPa	700~1060
Network (power)	scope		Hz	50±1
frequency	-			
Network (power	scope		V	100-240V
supply) voltage (rms)				
Preheat time	smallest		min	2
ECG part				
Overload protection: no	damage after loading 1V,	smallest	V	1
power frequency, differ	ential mode AC voltage			
for 10s				
Auxiliary output (if prov	ided)	none		1
No damage under short c	ircuit condition	-		
Respiration, lead-off de	tection and active noise	have		
suppression			1	1
DC current in active lead	S			
QRS wave detection	1 14 6 4			
QRS wave amplitude a	nd width range—for the			
pulse snown in Figur	e 6, comply with the			
requirements of 4.2.6			mV	0.5.5
Width (adult monitor)		scope	ma	0.5~5
Width (adult monitor)		scope	ma	/0~120
width(children)		scope	IIIS	40~120
No response to the follow	ving signals			
Amplitude (except in child working mode)		maximum	mV	0.15
Width when amplitude is 1mV (except children's		maximum	ms	10
operating mode)				
Power frequency voltage	tolerance	smallest	uV	100
Drift tolerance				
Triangular wave amplitu	de	N.A.	mV	4
QRS wave amplitude		N.A.	mV	0.5
QRS wave width		N.A.	ms	100
QRS complex repetition	rate	N.A.	bpm	80
Heart rate meter range and accuracy				1.5.000
Scope(adult monitor)		scope	bpm	15~300
Scope(child monitor)		scope	bpm	15~350
Error: or		maximum	%	±10
or (whichever is greater)		maximum	bpm	±5
Display heart rate for a minimum nominal heart	heart rate less than the rate	maximum	bpm	30
Display heart rate for sig	nals with repetition rate =	smallest	bpm	295
300 bpm (adult mode)			1	

Display heart rate for signals with repetition rate =	smallest	bpm	345
3500pm (children)			
Alarm system requirements			
Alarm limit range	11 4	1	100.200
Maximum limit (adult)	smallest	bpm	100~200
Upper limit (children)	smallest	bpm	100~250
Lower limit (adults and children)	smallest	bpm	30~100
Alarm resolution, or	smallest	bpm	1
Alarm limit error, or	maximum	%	±10
Or (whichever is greater)	maximum	bpm	±5
Start time of asystole alarm	maximum	s	10
Low heart rate alarm start time	maximum	S	10
High heart rate alarm start time	maximum	S	10
Silence alarm	Provides m	ute and resta	art settings
Alarm disabled	Display ala	rm is disable	ed
Monitor with ECG waveform display capability			
Enter dynamic range			
Input signal amplitude	maximum	mV	±5
Rate	maximum	mV/s	320
DC offset voltage	scope	mV	-300~+300
Output signal changes	maximum	%	± 10
Cannot work display (reduce the degree before	movimum	0/.	50
display)	maximum	70	30
Input impedance: signal minus (0.67Hz~40Hz)	maximum	%	20
System noise	maximum	uV	30
Multi-channel crosstalk: Channels without signals	movimum	0/.	5
are interfered by channels with signals	maximum	/0	5
Gain control and stability			
Gain selection			
All shown	smallest	mm/mV	2.5
Display permanently	N.A.	/	/
Allows continuous change of gain control, manual			
override			
Gain changes per minute	maximum	%/min	0.66
Total gain change within 1 hour	maximum	%	± 10
Time base selection and accuracy			
Time base selection			
—Display permanently	required	mm/s	25
—Non-permanent display	N.A.	mm/s	12.5, 25, 50
Time base maximum error	maximum	%	±10
Output display			
Channel width	Range	mm	Minimum10,maximum20
Aspect ratio	N.A.	s/mV	0.2, 0.4, 0.8
Input signal reconstruction accuracy			
Total system error: or	maximum	%	±20
Or (whichever is greater)	maximum	uV	±100
Frequency response			
		TT	0.67~40
Sine input	scope	Hz	(minus -3dB)
Response to 0.3mv.s impact outside the impact			/
range			
Offset	maximum	mV	0.1
Slope	maximum	mV/s	0.30

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Electrode weighting factor	smallest	%	±5	
Offset 15mm hysteresis effect	maximum	mm	0.5	
Scaling voltage	See 4.2.8.9	See 4.2.8.9 of YY1079		
Common mode rejection allows 10V line frequency noise	maximum	mV	1	
Baseline control and stability				
Recovery time after reset	maximum	S	3	
Drift rate within 10s	maximum	uV/s	10	
Baseline drift within 1 hour	maximum	uV	500	
Line drift under operating temperature	maximum	uV/°C	50	
Pacing pulse: display - when the amplitude is $2mv\sim700mv$, the width is $0.5ms\sim2ms$, the maximum rise time is $100\mu s$, ECG display when a pacing pulse of 100 pulses per minute occurs.	smallest	mV	0.2	
Sync Pulse: Time interval from R wave peak to sync output pulse, plus published amplitude, width and output impedance Z	N.A.	ms	35	
Electrosurgical interference suppression: heart rate changes compared with before interference	maximum	%	±10	

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Require	Announce		
WARNING information on whether overloading electrosurgical equipment will damage the equipment	No damage		
Breathing, lead off detection, active noise suppression	Not more than 30µA		
High T-wave suppression capability: The heart rate displays the maximum T-wave amplitude within the specified error limit. If performance is affected by low frequency response, options need to be published	Diagnosis and monitoring mode 0.9mV Surgical mode 0.3mV		
Heart Rate Average: Averaging algorithm for calculating minute heart rate	Average of eight shots, display refresh speed 1s		
Heart Rate Accuracy and Response to Arrhythmias: Heart Rate Display of Waveforms in Figure 3	They are 80bpm±5bpm, 60bpm±5bpm, 120bpm±5bpm, and 90bpm±5bpm respectively.		
The response time of the heart rate meter to changes in heart rate: 1. The number of seconds required for the heart rate meter to increase from 80bpm to 120bpm2, and to decrease from 80bpm to 40bpm	1. <10s; 2. <10s		
Start time of tachycardia alarm: Alarm time for the waveform in Figure 4	<10s		
Pace pulse suppression WARNING label: Because the heart rate meter may include pacemaker pulses in the heart rate value during cardiac arrest or arrhythmia, patients with pacemakers need to be closely monitored.	For patients with a pacemaker, the heart rate monitor may include pacemaker pulses in the heart rate value during cardiac arrest or arrhythmia. Don't rely entirely on heart rate monitor alarms		
Auditory Alarm Announcement: Sound Source Location and Sound Frequency	The speaker is placed on the rear of the product; the sound frequency is 1KHz		
Visual alarm announcement: location, color, size and flash frequency	The alarm light is on the upper left side of the from of the product; 1. The high-level alarm red ligh lights up, 11 seconds/time; 2. The intermediat alarm yellow light lights up, 25 seconds/time; 3 The low-level alarm yellow light lights up, 2 seconds/time		
Battery powered monitor: minimum operating time, battery charging time, battery consumption indication, functional description	When fully charged, it can work for 100 minutes by measuring blood pressure every half hour; the maximum battery charging time does not exceed 12 hours; there is a battery power indicator in the lower right corner of the product; the charging time to charge to 90% of the battery when the battery is consumed: KM8A and KM10A It is 4.3 hours, and other models are 3 hours. When the battery is under-voltage, the power indicator turns red and the power automatically cuts off after 10 minutes.		
Telemetry	N.A.		
Mains Isolated Monitor Transients: WARNING Messages and Methods to Reduce Interference	Connect the product to the ground correctly, treat the patient's skin correctly when attaching electrodes, and use electrodes produced by regular manufacturers.		
Special published requirements for monitors with non-permanent ECG waveform display: valid time base, aspect ratio adjustment range	Time bases are 12.5mm/s, 25 mm/s, 50 mm/s; aspect ratios are 0.2s/mV, 0.4 s/mV, 0.8 s/mV		

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Electrode Polarization: Cautionary Statement Regarding the Impact of Electrode Type on System Overload Recovery, Particularly Post-Defibrillation Recovery Time	Electrodes from the manufacturer specified by the company should be used, otherwise the recovery time after defibrillation will be affected.		
Auxiliary outputs: If auxiliary outputs are provided, provide instructions for proper connection to other equipment, paying particular attention to risk current characteristics, including bandwidth, gain, transmission delay, and handling of pacing pulses.	N.A.		
Alarm Silencing: Publishes the time the alarm will be reactivated after it has been silenced. If this time is adjustable, publish the range of its time interval	The time interval is 60s, 120s and the alarm is turned on and displayed on the screen		
Battery Disposal: Proper Guidance on Disposal of Used Batteries	Dispose of according to local end-of-life regulations for electronic products, or return to manufacturer for recycling		
Precautions for use: Description of the intended use and usable functions of the equipment; procedures for checking the controls and functions; information about the electrodes - the number of electrodes required, the polarity of the cable (if different from the standard cable); necessary for compliance with the requirements of this standard Description of any special cable characteristics; WARNINGs about using electrodes of dissimilar metals; settings necessary for child supervision	OSEN8000 series multi-parameter monitor is used in all levels of hospitals for patients to conduct electrocardiography, blood pressure, oxygen saturation, respiration, body temperature and other vital signs parameters are monitored and displayed on the screen in digital or waveform open, and can be recorded on the electrocardiogram and pulse oximetry output.; the ECG electrode uses 5 electrodes, white is RA, black is LA, red is LL, green is RL, and brown is C; the electrodes are copper, and different metal materials are not allowed. ; When the monitor is used for children, it must be set to the corresponding mode, otherwise the data may not be measured, the data may be inaccurate, and there may be safety risks to the patient.		
Application field description	Includes: in the operating room, induction of anesthesia and postoperative resuscitation, in the intensive care unit, emergency care, cardiac care, neurological care, dialysis care, geriatric care, obstetrical care, medical and surgical care.		
Service Instructions: Proper care, preventive maintenance, and repair procedures; complete electrical instructions to ensure proper on-site repair and specify acceptable repair equipment; recommended preventive maintenance frequency	If the customer requests it, maintenance instructions can be provided to the customer: the maintenance instructions include maintenance and preventive maintenance information as well as repair instructions. Includes circuit diagrams, block diagrams, wiring diagrams, and part numbers, information that ensures skilled technicians will complete the maintenance items specified in the service instructions that can be performed by a service representative.		
Pacemaker pulse suppression ability/no overshoot pacemaker pulse suppression: In the operator's manual and maintenance manual, there is no overshoot pacemaker pulse suppression ability in the following range: ± 2mV to ± 700mV amplitude, 0.1ms to 2.0ms width, overshoot less than 0.05ap [Figure 5a], and stabilization time less than 5us; The starting, rising, and falling times of the pulse shall not exceed 100us; The pulse initiation time is	When the pacing analysis switch is turned on, pacing pulses that meet the following conditions can be suppressed without affecting heart rate calculation; Amplitude: $\pm 2 \text{ mV} \sim \pm 700 \text{mV}$; Width: 0.1ms ~ 2ms; Rise time: $10 \mu \text{s} \sim 100 \mu \text{s}$		

40ms or less before the QRS wave initiation; There is an identical pulse with a time lead of 150ms to 250ms before the praince pulse mentioned above	
Decemplear mulae summersion with events	
Pacemaker pulse suppression with overshoot: Pacemaker pulse suppression with overshoot (charging) time constant between 4ms and 100ms is published in the operator's instructions and maintenance instructions. The other parameters are the same as the above except that the overshoot (charge) time constant is between 4ms and 100ms. Overshoot has Two definition methods. Is the announcement using Method A or Method B or both?	Use Method A
Suppression of fast ECG signals by pacing pulse detectors; the typical minimum input slew rate, expressed in V/sRTI, that can render the pacing pulse detector inactive must be published in the operator's instructions and service instructions	5V/sRTI
Display of pacing pulses on the auxiliary output: When there is a pacing pulse on the auxiliary output, the filtering of the pacemaker pulses and any substitutions are announced	N.A.
Pace Pulse Detector Failure: Publish operating modes or signal conditions that cause pacing pulse detection to fail or be ignored	Pacing analysis is off or pacing analysis is on but the product is operating in surgical mode

Appendix III Declaration Table of Toxic and

Harmful Substances or Elements in Products

	Toxic and harmful substances or elements						
Part Name	Lead (pb)	Hg (hg)	Cadmium (cd)	Hexavalent chromium (cr(vi))	Polybrominated biphenyls (pbb)	Polybrominated diphenyl ethers (pbde)	
Shell	0	0	0	0	0	0	
Operation panel	0	0	0	0	0	0	
РСВА	0	0	0	0	0	0	
Label	0	0	0	0	0	0	
Display	×	×	×	×	×	×	
Packaging Materials	×	×	0	0	×	×	
Connector	0	0	0	×	0	0	
Power cable	0	0	0	0	0	0	
Battery	×	0	×	0	0	0	
Accessories and sensors	×	0	0	0	0	0	

Remarks: 0: Indicates that the content of the toxic and hazardous substance in all homogeneous materials of the part is below the limit requirements specified in the SJ/T 11363-2006 standard; ×: Indicates that the toxic and hazardous substance is present in at least one homogeneous material of the part The content in the material exceeds the limit requirements specified in the SJ/T 11363-2006 standard.

When disposing of products and batteries after normal use, please comply with the laws and regulations of each local government for disposal.

The product contains certain toxic and harmful substances, which can be used with confidence within the environmental protection use period. After the environmental protection use period exceeds, it should enter the recycling system.

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Appendix IV EMC

This equipment generates and radiates radio frequency energy. This equipment can cause electromagnetic interference between other medical or non-medical equipment and radio communications. According to the statement in YY 9706.102-2021, this product belongs to the first group of emission restrictions, Class A medical equipment, and corresponding protection measures should be provided to avoid interference. However, there is no complete guarantee that electromagnetic interference will not occur under specific installation conditions.

When it is found that this equipment causes interference (it needs to be confirmed by switching the equipment on and off), the operator (or authorized maintenance personnel) can eliminate the interference according to the following measures:

- 1. Adjust or relocate the affected equipment;
- 2. Increase the distance between this device and the affected device;
- 3. Use another power source to power the device;
- 4. Consult maintenance personnel for more suggestions.

WARNING

•The use of non-original accessories, sensors, and cables may increase the device's electromagnetic emissions or reduce its electromagnetic immunity.

•KC7 can be placed on the desktop, also can be equipped with quick mounting clips, and then fixed on the diameter of the tube below 30mm (such as the hospital bed stopper, infusion bracket and other positions) on the use. It can also be plugged into MDKMed's KC16\19\22 mainframe to be used as a sub-machine..

Motice

•Before using this equipment, please make sure that the EMC requirements required by this equipment are met.

•This section will list the contents described in the YY 9706.102-2021 table. It is the user's responsibility to ensure that this equipment and its nearby equipment comply with the radio frequency interference parameters indicated in the general safety requirements.

•Do not use equipment that intentionally emits RF signals (cell phones, radio transceivers or radio-controlled products) in the vicinity of this equipment as this may cause operation to

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exceed specified values. Please turn off this type of equipment when near it. It is the operator's responsibility to instruct the patient or other persons accessing this device to fully comply with the above requirements.

•The manufacturer will not be responsible for any interference caused by the use of non-recommended internal connecting cables or by unauthorized changes or modifications to this equipment.

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Table I

Guidance and Manufacturer's StatementElectromagnetic Emissions				
[Systems and equipment] are expected to be used in the electromagnetic environment specified below. The				
purchaser or user show	purchaser or user should ensure that it is used in this electromagnetic environment.			
Launch test	Conformity Electromagnetic Environment - Guidance			
RF emissions GB 4824	Group 1	[The system or device] uses radio frequency energy only for its internal functions. Therefore, its RF emissions are very low and have a low potential to cause interference in nearby electronic equipment.		
RF emissions GB 4824	Class A	[System or equipment] Applicable to		
Harmonic emissions GB 17625.1	Class A	non-domestic use and all facilities not directly connected to the public low-voltage power supply network for domestic use.		
Voltage fluctuations/flicker emissions GB 17625.2	Conformed			

Table II

Guidance and Manufacturer's StatementElectromagnetic Immunity				
[Systems and equipment] are expected to be used in the electromagnetic environment specified below. The				
Immunity test	Immunity test IEC60601 test level Compliance level		Electromagnetic Environment - Guidance	
electrostatic discharge GB/T 17626.2	±8 kV contact discharge ±15 kV air discharge	±8 kV contact discharge ±15 kV air discharge	Floors should be wood, concrete or ceramic tiles. If the floor is covered with synthetic material, the relative humidity should be at least 30%.	
electrical burst GB/T 17626.4	±2 kV to power line ±1 kV for input/output lines	±2 kV power line	Mains power should be of a quality typically used in a commercial or hospital environment.	
surge GB/T 17626.5	±1 kV line-to-line ±2 kV line to ground	±1 kV line-to-line ±2 kV line to ground	Mains power should be of a quality typically used in a commercial or hospital environment.	
Voltage sags, short interruptions and voltage changes on power input lines GB/T 17626.11	<5% UT for 0.5 cycles (On UT, >95% dip) 40% UT for 5 cycles (On UT, 60% dip) 70% UT for 25 cycles (On UT, 30% dip) <5% UT, lasting 5s (On UT, >95% dip)	<5% UT (drop > 95% UT) 0.5 period 40% UT (drop 60% UT) 5 cycles 70% UT (drop 30% UT) 25 cycles <5% UT (drop > 95% UT) 5 seconds	Mains power should be of a quality typically used in a commercial or hospital environment. If the user of [equipment or system] requires continuous operation during a power outage, it is recommended that [equipment or system] be powered by an uninterruptible power supply or battery.	

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Power frequency magnetic field (50/60Hz) IEC 61000-4-8 GB/T17626.8	3 A/m	3 A/m	Power frequency magnetic fields should have power frequency magnetic field level characteristics typical of a typical location in a typical commercial or hospital environment.
Note: UT refers to the AC network voltage before applying the test voltage.			

Table 3

Guidance and Manufacturer's StatementElectromagnetic Immunity			
[Systems and equipment] are expected to be used in the electromagnetic environment specified below. The purchaser or user should ensure that it is used in this electromagnetic environment.			
Immunity	IEC60601 test	Compli	Electromagnetic Environment - Guidance
test	level	ance	
		level	
RF	3 Vrms	3 Vrms	Portable and mobile RF communications equipment
GB/T	150K~80MHZ		should not be placed closer than recommended to any part of
17626.6			this mobile equipment, including cables. This distance should
			be calculated by the formula corresponding to the frequency
			of the transmitter.
			Recommended isolation distance
radiofreque	3V/m		$d=1.2\sqrt{P}\sqrt{P}$
radiation GB/T	80M~2.5GHz	3V/m	d=1.2 VP VP 80MHz~ 800MHz
17626.3			d=2.3 √P √P 800MHz~2.5GHz
			In the formula:
			Paccording to the maximum rated output power of the
			transmitter provided by the transmitter manufacturer, in watts
			(W);
			dRecommended separation distance, unit is meters
			(m).
			The field strength of fixed RF transmitters is determined
			by a survey of the electromagnetic field a and should be
			lower than the compliance level in each frequency range b.
			Interference may occur in the vicinity of equipment
			marked with the following symbol.
			(((••)))
Note 1:At the 80MHz to 800MHz frequency point, the higher frequency formula is used.			
Note 2: These guidelines may not be suitable in all situations. Electromagnetic propagation is affected by			

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absorption and reflection from buildings, objects and people.

aFixed transmitters, such as base stations for wireless (cellular/cordless) telephones and land mobile radios, amateur wireless, AM and FM radio broadcasts, and television broadcasts, the field strengths of which cannot be predicted accurately theoretically. To assess the electromagnetic environment of fixed RF transmitters, a survey of the electromagnetic field should be considered. If the mobile device is exposed to measured field strengths above the applicable RF compliance levels above, the mobile device should be observed to verify normal operation. If unusual performance is observed, additional measures may be necessary, such as reorienting or relocating the mobile device.

bIn the entire frequency range of 150kHz~80MHz, the field strength should be lower than 3V/m.

Table 4

Recommended isolation distance between portable and mobile radio frequency communications equipment and [equipment or system]

[Equipment or system] is intended for use in an electromagnetic environment where radio frequency radiation disturbance is controlled. Based on the maximum rated output power of the communications equipment, the purchaser or user of [equipment or system] can prevent electromagnetic interference by maintaining the minimum distance recommended below between portable and mobile radio frequency communications equipment (transmitters) and [equipment or system]

	Calculate the isolation distance (meters) based on the frequency of the			
Maximum rated output	transmitter			
power of the	150kHz to 80MHz	80MHz to 800MHz	800MHz to 2.5GHz	
transmitter in W	d=1.2 √P √P	d=1.2√P√P	d=2.3√P√P	
0.01	0.12	0.12	0.23	
0.1	0.38	0.38	0.37	
1	1.2	1.2	2.3	
10	3.8	3.8	7.3	
100	12	12	twenty three	

For the rated maximum output power of the transmitter not listed in the above table, the recommended isolation distance d, in meters (m), can be determined by the formula in the corresponding transmitter frequency column, where P is the transmitter provided by the transmitter manufacturer. The maximum output power rating of the machine, in watts (W).

Note 1: At frequencies 80 MHz and 800 MHz, the formula for the higher frequency range applies. Note 2: These guidelines may not be suitable in all situations. Electromagnetic propagation is affected by absorption and reflection from buildings, objects and human bodies.

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