cPM 12/cPM 8

Patient Monitor

Operator's Manual

C € 0123

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- the electrical installation of the relevant room complies with the applicable national and local requirements;
- the product is used in accordance with the instructions for use.

WARNING

- Only skilled/trained clinical professionals should operate this equipment.
- It is important for the hospital or organization that uses this equipment to perform a recommended service/maintenance plan. Neglect of this may result in machine breakdown or personal injury.

Service

Mindray maintains a network of service representatives and factory-trained distributors. Prior to requesting service, perform a complete operational check of the instrument to verify proper control settings. If operational problems continue to exist, contact Mindray Service.

In North America contact the Service Department at (800) 288-2121, ext: 8116 for Technical Support or (201) 995-8000 for assistance in determining the nearest field service location.

Please include the instrument model number, the serial number, and a description of the problem with all requests for service.

Any questions regarding the warranty should be directed to your local sales or service representative.

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Preface

Manual Purpose

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

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

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

Intended Audience

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

Illustrations

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

Manual Conventions

- *Italic text* is used to quote the referenced chapters or sections.
- [] is used to enclose screen text.
- \rightarrow is used to indicate operational procedures.

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1.1 Safety Information

WARNING

• Indicates a potential hazard or unsafe practice that, if not avoided, could result in death or serious injury.

CAUTION

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

NOTE

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

1.1.1 Warnings

WARNINGS

- Before putting the system into operation, the operator must verify that the equipment, connecting cables and accessories are in correct working order and operating condition.
- To avoid risk of electric shock, this equipment must only be connected to a supply mains with protective earth. If a properly grounded power outlet is not available, operate the monitor on battery power.
- The equipment is not intended to be used within the Magnetic Resonance (MR) environment.
- To avoid an explosion hazard, do not use the equipment in the presence of oxygen-rich atmospheres, flammable anesthetics, or other flammable agents (such as gasoline).
- Before connecting the equipment to the power line, check that the voltage and frequency ratings of the power line are the same as those indicated on the equipment's label or in this manual.
- Do not open the equipment housings. All servicing or future upgrades must be carried out by Mindray trained and authorized personnel.
- Do not come into contact with patients during defibrillation. Otherwise serious injury or death could result.
- Do not rely exclusively on the audible alarm system for patient monitoring. Adjustment of alarm volume to a low level or off may result in a hazard to the patient. Remember that alarm settings should be customized according to different patient situations and always keeping the patient under close surveillance is the most reliable way for safe patient monitoring.

WARNINGS

- PATIENTS WITH A PACEMAKER On ventricular paced patients, episodes of Ventricular Tachycardia may not always be detected. Do not rely entirely upon the system's automated arrhythmia detection algorithm. Keep pacemaker patients under close surveillance.
- The physiological data and alarm messages displayed on the equipment should be reviewed by a clinician before being used for diagnostic interpretation and treatment.
- To avoid inadvertent disconnection, route all cables in a way to prevent a stumbling hazard. Wrap and secure excess cabling to reduce risk of entanglement by patients or personnel.
- Do not touch the equipment's metal parts or connectors when in contact with the patient; otherwise patient injury may result.
- Never mix patient electrode types or brands. Dissimilar metals or other incompatibilities may cause considerable baseline drift and may increase trace recovery time after defibrillation.
- To reduce the hazard of burns during high-frequency surgical procedure, ensure that the monitor's cables and transducers never come into contact with the electro-surgery unit (ESU).
- The neutral electrode of the electro-surgery unit (ESU) shall properly contact the patient. Otherwise, burns may result.
- Magnetic and electrical fields are capable of interfering with the proper performance of the equipment. For this reason make sure that all external devices operated in the vicinity of the equipment comply with the relevant EMC requirements. Mobile phone, X-ray equipment or MRI devices are a possible source of interference as they may emit higher levels of electromagnetic radiation.

1.1.2 Cautions

CAUTIONS

- Only use parts and accessories specified in this manual.
- Disposable accessories are not designed to be reused. Reuse may cause a risk of contamination and affect the measurement accuracy.
- Always install or carry the equipment properly to avoid damage caused by drop, impact, strong vibration or other mechanical force.
- Dry the equipment immediately in case of rain or water spray.
- Ensure that the patient monitor is supplied with continuous electric power during work. Sudden power failure may lead to data loss.
- This equipment is intended for single patient use.
- Store and use the equipment in specified environmental condition. The monitor and accessories may not meet the performance specification due to aging, stored or used outside the specified temperature and humidity range.
- Contact the Mindray service personnel for replacements if you find the housing is broken.

1.1.3 Notes NOTES

- Put the equipment in a location where you can easily view and operate the equipment.
- During normal use, the operator is expected to face the front of the equipment.
- The equipment uses a mains plug as a means of isolation to the mains power supply. Do not position the equipment in a place difficult to operate the mains plug.
- Remove the battery before shipping the monitor or if it will not be used for an extended period of time.
- Keep this manual in the vicinity of the equipment so that it can be conveniently referenced when needed.
- The software was developed in compliance with IEC60601-1-4. The possibility of hazards arising from software errors is minimized.
- This manual describes all features and options. Your equipment may not have all of them. They are not available in all geographies.
- At the end of its service life, the equipment, as well as its accessories, must be disposed of in compliance with the guidelines regulating the disposal of such products. If you have any questions concerning disposal of the equipment, please contact Mindray.
- When disposing of the packaging material, be sure to observe the applicable waste control regulations and keep it out of children's reach.

1.2 Equipment Symbols

Some symbols may not appear on your equipment.

\triangle	Caution		
	Refer to instruction manual/ booklet		General warning sign
⊙/Ò	Power ON/OFF (for a part of the equipment)	+	Battery indicator
\sim	Alternating current	$\stackrel{\bullet}{\frown}$	VGA output
\forall	Equipotentiality	5	Graphical record
● <u></u>	USB connector		Network connector
->0←	Zero key		Input/Output
	Check sensor		Calibrate key
$^{ } / $	Measure/standby		Inserted direction

	Gas outlet	SN	Serial number	
~~	DATE OF MANUAFACTURE		Symbol for "MANUFACTURER"	
REF	Catalog number	IPX1	Protected against vertically falling water drops per IEC 60529	
	Electrostatic sensitive devices	E.	Plastics identification symbol	
⊣♥⊦	DEFIBRILLATION-PROOF TYPE CF APPLIED PART			
۱ 🖈 ۲	DEFIBRILLATION-PROOF TYPE BF APPLIED PART			
ETL CLASSIFIED	The presence of this label indicates the machine was certified by ETL with the statement: Conforms to AAMI Std. ES 60601-1, IEC Std. 60601-1-6, IEC Std. 60601-1-8, IEC Std. 60601-2-25, IEC			
	Std. 60601-2-26, IEC Std. 60601-2-27, IEC Std. 60601-2-34, IEC Std. 60601-2-49, IEC Std. 80601-2-30, ISO Std. 80601-2-55, ISO Std. 80601-2-56, ISO Std. 80601-2-61			
LISTED US	Certified to CSA Std. C22.2 NO. 60601-1, NO. 60601-6, NO. 60601-1-8, NO. 60601-2-25, NO.			
	Intertek 60601-2-26. NO. 60601-2-27. NO. 60601-2-34. NO. 60601-2-49. NO. 80601-2-30. NO. 80601-2-55. J			
3179617	80601-2-56, NO. 80601-2-61			

2 The Basics

2.1 Intended Use

The cPM Series Patient Monitors (cPM 8 and cPM 12) are intended to be used for monitoring, displaying, reviewing, storing and alarming of multiple physiological parameters including ECG (3-lead, or 5-lead), arrhythmia detection and QT analysis, heart rate (HR), respiration (Resp), temperature (Temp), pulse oxygen saturation (SpO2), pulse rate (PR), non-invasive blood pressure (NIBP), invasive blood pressure (IBP), cardiac output (C.O.), carbon dioxide (CO2), and anesthetic gas (AG).

All the parameters can be applied to single adult, pediatric or neonatal patients with the exception of the following:

- C.O. monitoring is restricted to adult patients only;
- PAWP monitoring is not intended for neonatal patients;
- The ECG Algorithm arrhythmia detection is intended for adult and pediatric patients.
- AG monitoring is available for cPM 12 Patient Monitor only.

The monitors are to be used in healthcare facilities by clinical physicians or appropriate medical staff under the direction of physicians.

WARNING

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

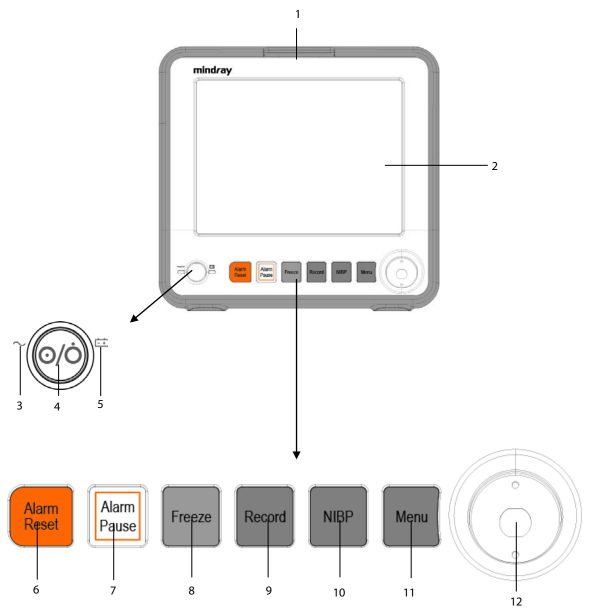
2.2 Applied Parts

The applied parts of the cPM 8/12 patient monitors are:

- ECG electrodes and leadwires,
- SpO₂ sensor
- NIBP cuff
- Temp probes
- IBP transducer
- C.O. sensor
- CO₂ sampling line/nasal sampling cannula, and water trap
- AG sampling line, water trap, and airway adapter

2.3 Main Unit

2.3.1 Front View



1. Alarm lamp

When a physiological alarm or technical alarm occurs, this lamp flashes as defined below.

- High level alarms: the lamp quickly flashes red.
- Medium level alarms: the lamp slowly flashes yellow.
- Low level alarms: the lamp lights yellow without flashing.
- 2. Display Screen
- 3. AC power LED
- 4. Power On/Off Switch
 - Pressing this switch turns the monitor on.
 - When the monitor is on, pressing and holding this switch turns the monitor off.

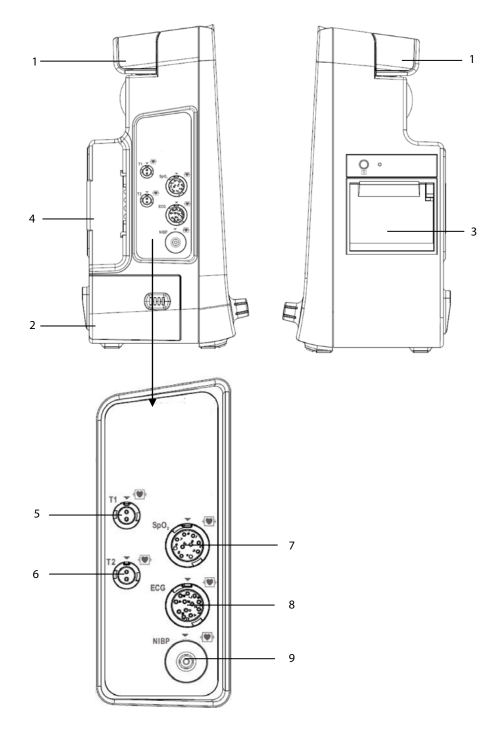
An indicator is built into this switch. It turns on when the monitor is on and turns off when the monitor is off.

- 5. Battery LED
 - On: when the battery is installed and the AC source is connected.
 - Off: when no battery is installed or the installed battery is malfunctioning, or no AC source is connected when the monitor is powered off.
 - Flashing: when the monitor operates on battery power.
- 6. Alarm Reset: Press to reset the alarms.
- 7. Alarm Pause: Press to pause or restore alarms.
- 8. Freeze: Press to freeze or unfreeze waveforms.
- 9. **Record**: When a recording task is not started, pressing this button starts recording. When a recording task is in progress, pressing this button stops recording.
- 10. **NIBP**: Press to start or stop NIBP measurements.
- 11. **Menu**

If no menu is currently displayed on the screen, pressing it enters the main menu. If a menu is currently displayed on the screen, pressing it closes that menu.

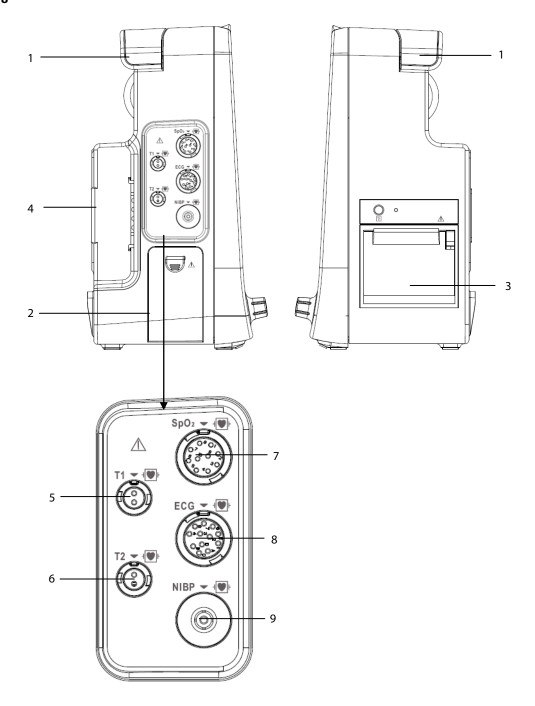
- 12. Knob
 - Rotate the knob clockwise or counter-clockwise to move the cursor.
 - Press the knob to select one item, such as accessing a menu or confirming the selection.

cPM 12



- 1. Handle
- 3. Recorder
- 5. Temp probe 1 connector
- 7. SpO₂ cable connector
- 9. NIBP cuff connector

- 2. Battery compartment
- 4. Parameter module slot
- 6. Temp probe 2 connector
- 8. ECG cable connector

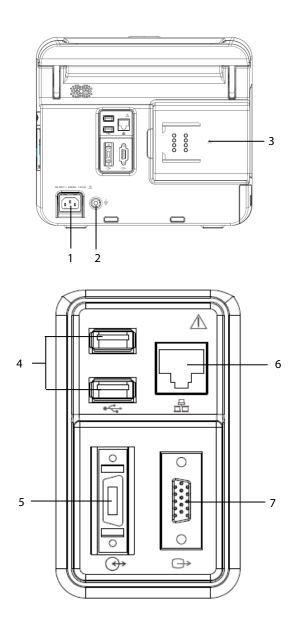


- 1. Handle
- 3. Recorder
- 5. Temp probe 1 connector
- 7. SpO₂ cable connector for
- 9. NIBP cuff connector

- 2. Battery compartment
- 4. Parameter module slot
- 6. Temp probe 2 connector
- 8. ECG cable connector

2.3.3 Rear View

cPM 12



- 1. AC Power Input
- 2. Equipotential Grounding Terminal

When using the monitor together with other devices, connect their equipotential grounding terminals together to eliminate the potential differences between them.

3. Parameter Module Slot

Used for connecting the parameter modules.

4. USB Connectors

Connects a USB drive for data or configuration transfer.

- 5. Multifunctional Connector:
 - Connects the monitor to external devices and outputs defibrillator synchronization signals, nurse call signals, analog output signals, measurement numerics, and alarm limits of parameter modules.
 - Communicates with external devices via Datascope Improved ASCII Protocol (DIAP) protocol.

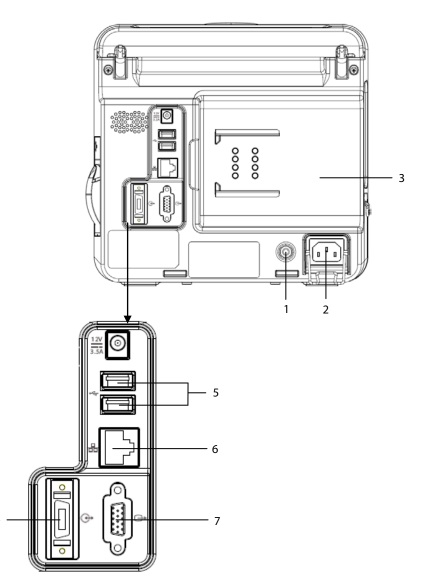
6. Network Connector

A standard RJ45 connector which connects the monitor to another monitor for remote viewing. It also connects the monitor to a PC for a system upgrade.

7. VGA Connector

Connects a secondary display, which extends the display capability of your monitor. The contents displayed on the secondary display screen matches what displays on the monitor screen.

cPM 8



1. Equipotential Grounding Terminal

When using the monitor together with other devices, connect their equipotential grounding terminals together to eliminate the potential differences between them.

- 2. AC Power Input
- 3. Parameter Module Slot

Used for connecting the parameter modules.

- 4. Multifunctional Connector
 - Connects the monitor to external devices and outputs defibrillator synchronization signals, nurse call signals, analog output signals, measurement numerics, and alarm limits of parameter modules.
 - Communicates with external devices via Datascope Improved ASCII Protocol (DIAP) protocol.
- 5. USB Connectors

Connects a USB disk for data or configuration transfer.

6. Network Connector

A standard RJ45 connector which connects the monitor tor other monitor for remote view. It also connects the monitor to PC for system upgrade.

7. VGA Connector

Connects a secondary display, which extends the display capability of your monitor. The contents displayed on the secondary display screen accords with those displayed on the monitor screen.

2.4 Modules





- 1. Sidestream CO₂ module:
- 2. IBP+C.O. + microstream CO₂ module:
- 3. IBP+C.O.+AG module (with O₂):

contains CO₂ watertrap connector, and CO₂ gas outlet.

contains IBP cable connector, C.O. cable connector, CO_2 sampling line connector, and CO_2 gas outlet.

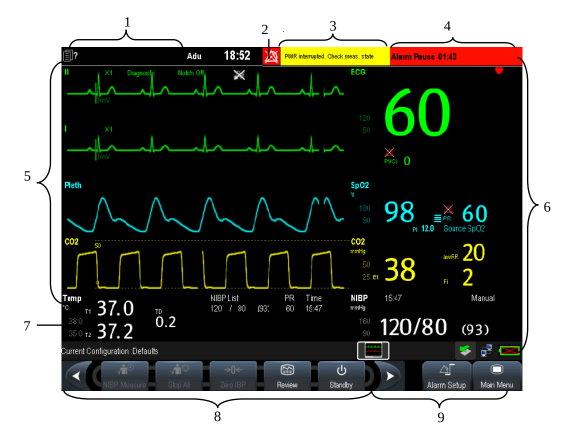
contains IBP cable connector, C.O. cable connector, AG watertrap slot, and AG gas outlet. This module is for cPM 12 only.

NOTE

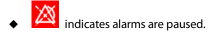
• The above modules with IBP connectors support two invasive blood pressures through an IBP extended cable (PN: 040-001029-00).

2.5 Display Screen

This patient monitor adopts a high-resolution LED screen to display patient parameters and waveforms. A typical display screen is shown below.



- 1. Patient Information Area
 - This area shows date, time, and the patient information such as department, bed number, patient name and patient category. ? indicates that no patient is admitted or the patient information is incomplete. If no patient is admitted, selecting this area enters the [Patient Setup] menu. For admitted patients, selecting this area enters the [Patient Demographics] menu.
- 2. Alarm Symbols



- indicates alarms are reset.
- indicates alarm sounds are turned off.
- indicates the system is in alarm off status.

3. Technical Alarm Area

This area shows technical alarm messages and prompt messages. When multiple messages occur, the messages scroll. Select this area and the technical alarm list displays.

4. Physiological Alarm Area

This area shows physiological alarm messages. When multiple alarms occur, the messages scroll. Select this area and the physiological alarm list displays.

5. Waveform Area

This area shows measurement waveforms. The waveform name displays at the upper left corner of the waveform. Select this area and the corresponding measurement setup menu displays.

6. Parameter Area A

This area shows measurement parameters. Each monitored parameter has a parameter window and the parameter name displays at the upper left corner. The corresponding waveform of each parameter displays in the same row in the waveform area. Select this area and the corresponding measurement setup menu displays.

7. Parameter Area B

For the digital parameters displayed in this area, their corresponding waveforms are not displayed.

8. Prompt Message Area

This area shows the current configuration name, prompt messages, network status icons, and battery status icons, etc. For details about battery status symbols, refer to chapter **24** *Batteries*.

indicates the monitor is successfully connected to a wired network.

- Indicates the monitor has failed to connect to a wired network.
- indicates a USB drive is inserted.
- 9. QuickKeys Area

This area contains QuickKeys that provide quick access to functions.

2.6 QuickKeys

A QuickKey is a configurable graphical key, located at the bottom of the main screen. They give you fast access to functions. Their availability, and the order in which they appear on your screen, depend on your monitor configuration.

To configure QuickKeys:

- 1. Select [Main Menu] \rightarrow [Maintenance >>] \rightarrow [Manage Configuration >>] \rightarrow enter the required password \rightarrow [Ok].
- 2. In the [Manage Configuration] menu, select [Edit Config.>>].
- 3. In the pop-up menu, select the desired configuration and then select [Edit].
- 4. In the pop-up menu, select [Screen Setup >>].
- 5. Select the [Select QuickKeys] tab, and then configure your desired QuickKeys and the order of them.

	Scroll left to display more QuickKeys.		Scroll right to display more QuickKeys.
	[Main Menu] QuickKey	Ċ	[Standby] QuickKey
	Enter the [Main Menu]		Enter standby mode
	[Alarm Setup] QuickKey	3	[Review] QuickKey
	Enter the [Alarm Setup] menu	2	Enter the [Review] menu
, 10	[NIBP Measure] QuickKey	, † ♡	[Stop All] QuickKey
- 7	Enter the [NIBP Measure] menu	৵∎	Stop all NIBP measurement
→0 ←	[Zero IBP] QuickKey	·*/>	[Alarm Reset] QuickKey
	Enter the [Zero IBP] menu	2×	Reset the alarm system
	[Alarm Pause] QuickKey	P	[Screens] QuickKey
	Pause or restore alarms		Enter the [Screens] menu
n ?	? [Patient Setup] QuickKey	[Manual Event] QuickKey	
'I ' <i>'</i>	Enter the [Patient Setup] menu	-71	Trigger a manual event
ፈጉ	[Realtime Print] QuickKey		[Print Setup] QuickKey
	Start the realtime print		Enter the [Print Setup] menu
Austand Austral	[Minitrends] QuickKey	linitrends] QuickKey	[Volume Setup] QuickKey
Assimt	Have a split-screen view of minitrends		Enter the [Volume Setup] menu
മി.	[Load Config.] QuickKey		[Start C.O.] QuickKey
Ð	Enter the [Load configuration] menu		Enter the C.O. measurement window
	[Calculations] QuickKey	m m	[Others] QuickKey
	Enter the [Calculations] menu	TT	Enter the [View Other Patient] window
	[Parameters] QuickKey	.	[OxyCRG] QuickKey
	Enter the [Parameters] menu		Enter the [oxyCRG] window
ிய	[NIBP STAT] QuickKey		[PAWP] QuickKey
-7 - 1	Start NIBP STAT measurement		Enter the PAWP measurement screen
2	[Night Mode] QuickKey	mmHg	[Unit Setup] QuickKey
	Enter the [Night Mode] menu		Enter the [Unit Setup] menu

The following QuickKeys are available:

~	[Privacy Mode] QuickKey Enter the privacy mode		[CPB Mode] QuickKey Enter the CPB mode
	[7-Lead ECG] QuickKey Enter the 7-lead ECG full screen	ŧ	[Discharge Patient] QuickKey Enter the [Discharge Patient] menu

NOTE

• There is space for only 10 user configurable Quickkeys on the cPM 12/cPM 8 patient monitor.

3.1 Installation

WARNING

- Connect only approved devices to this equipment. Devices connected to the equipment must meet the requirements of the applicable IEC standards (e.g. IEC 60950 safety standards for information technology equipment and IEC 60601-1 safety standards for medical electrical equipment). The system configuration must meet the requirements of the IEC 60601-1 medical electrical systems standard. Any personnel who connect devices to the equipment's signal input/output port is responsible for providing evidence that the safety certification of the devices has been performed in accordance to the IEC 60601-1. If you have any questions, please contact Mindray.
- If it is not evident from the equipment specifications whether a particular combination with other devices is hazardous, for example, due to summation of leakage currents, please consult the manufacturers or else an expert in the field, to ensure the necessary safety of patients and all devices concerned will not be impaired by the proposed combination.

CAUTION

• The equipment should be installed by authorized Mindray personnel.

NOTE

- The software equipment copyright is solely owned by Mindray. No organization or individual shall resort to modifying, copying, or exchanging it or to any other infringement on it in any form or by any means without due permission.
- Many settings in the patient monitor are password protected. It is recommended to change the default passwords and keep the passwords safe. Passwords should only be changed by authorized personnel. Contact your department manager or biomedical engineering department for the passwords used at your facility.

3.1.1 Unpacking and Checking

Before unpacking, examine the packing case carefully for signs of damage. If any damage is detected, contact the carrier or Mindray.

If the packing case is intact, open the package and remove the equipment and accessories carefully. Check all materials against the packing list and check for any mechanical damage. Contact Mindray in case of any problem.

CAUTION

- When disposing of the packaging material, be sure to observe the applicable waste control regulations and keep it out of children's reach.
- Before use, please verify whether the packages are intact, especially the packages of single use accessories. In case of any damage, do not apply it to patients.

NOTE

• Save the packing case and packaging material as they can be used if the equipment must be reshipped.

3.1.2 Environmental Requirements

The equipment operating environment must meet the requirements specified in this manual.

The equipment operating environment should be reasonably free from noises, vibration, dust, corrosive, flammable and explosive substances. To maintain good ventilation, the equipment should be at least 2 inches (5 cm) away from surrounding objects.

When the equipment is moved from one place to another, condensation may occur as a result of temperature or humidity difference. In this case, never start the system before the condensation disappears.

CAUTION

- Make sure that the equipment operating environment meets the specific requirements. Otherwise unexpected consequences, e.g. damage to the equipment, could result.
- Put the equipment in a location where you can easily see the screen, access the operating controls, and disconnect the equipment from AC power.

3.2 Getting Started

3.2.1 Connecting to Power Source

Using AC Power Source

To use the AC power source, connect one end of the power cord with the AC power input on the equipment's back panel and the other end with a wall AC mains outlet.

WARNING

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

Using a Battery

This monitor can be equipped with rechargeable lithium-ion battery. If a battery is installed, the monitor system automatically switches to the battery for power if AC power is interrupted.

NOTE

• When a battery has been stored for a long time, or the battery is depleted, recharge the battery at once. Otherwise, the low battery may not be sufficient to power the monitor if the AC power is unavailable.

Refer to Chapter 24 Batteries for detail.

Connecting Accessories

Insert the connector of accessory cables necessary for the measurements to be performed by your monitor to the corresponding connector on the monitor. For details about the parameter connectors, refer to **2.3.2** *Side View*

3.2.2 Turning Power On

Once the monitor is installed, before beginning measurements:

- 1. Check the monitor and plug-in modules for any mechanical damage, and make sure that all external cables, plug-ins and accessories are properly connected.
- 2. Check the power supply specification is met if mains power is used. Only use a power outlet that is properly grounded.
- 3. Plug the power cord into the AC power source. If you run the monitor on battery power, ensure that the battery is sufficiently charged.
- 4. Press the power on/off switch on the monitor's front panel.

The monitor will perform alarm system self-test during start-up. After pressing the power on/off button, the system shows a "**Loading...**" screen, sounds a beep and the alarm lamp simultaneously turns yellow, then red, and then turns off, followed by the start-up screen being shown. Then the start-up screen disappears. The alarm system self-test succeeds. The monitor enters the normal monitoring screen.

NOTE

• Carefully check if the system performs the self-test as described above. Contact your service personnel or Mindray if the self-test is abnormal.

WARNING

• Do not use the monitor on a patient if you suspect it is not working properly, or if it is mechanically damaged. Contact your service personnel or Mindray.

3.2.3 Starting Monitoring

- 1. Decide which measurements you want to take.
- 2. Connect the required modules, patient cables and sensors.
- 3. Make sure that the patient cables and sensors are correctly connected.
- 4. Make sure that the patient settings, such as [**Patient Cat.**], [**Paced**], etc, are appropriate for your patient.
- 5. Refer to the appropriate measurement section for details of how to perform the measurements you require.

3.3 Turning Off the Monitor

Before turning off the monitor,:

- 1. Ensure that the monitoring of the patient has been completed.
- 2. Disconnect the cables and sensors from the patient.
- 3. Make sure to save or clear the patient monitoring data as required.
- 4. Press and hold the power on/off switch for more than 2seconds to turn off the monitor. "**System is shutting down...**" is displayed on the screen and then the monitor shuts down.

CAUTION

- Press and hold the power on/off switch for 10 seconds to forcibly shut down the monitor if it could not be shut down normally. This may cause loss of patient data.
- The monitor restores the latest configuration if it restarts within 60 seconds after a power failure. The monitor restores the default configuration, rather than the latest configuration, if it restarts 120 seconds after a power failure. The monitor may load either the latest configuration or the default configuration if it restarts from 60-120 seconds after a power failure.
- Power failure may cause data corruption on the SD card. It is recommended to turn off the monitor according to the normal procedures. Do not directly unplug the power cord, unless a charged battery is installed, or remove the battery before shutting down the monitor.

NOTE

• To completely disconnect the power supply, unplug the power cord.

3.4 Using the Knob

- Rotate the knob clockwise or counterclockwise to move the cursor.
- Press the knob to select one item, such as accessing a menu or confirming the selection.

3.5 Using Keys

The monitor has three types of keys:

- Softkey: A softkey is a graphic key on the screen, giving you fast access to certain menus or functions. The monitor has two types of softkeys:
 - Parameter keys: Each parameter area can be seen as a softkey. You can enter a parameter setup menu by selecting its corresponding parameter or waveform area.
 - QuickKeys: QuickKeys are configurable graphical keys, located at the bottom of the main screen. For details, refer to the section **2.6 QuickKeys**.
- Hardkeys: A hardkey is a physical key on a monitoring device, such as the [Main Menu] hardkey on the monitor's front panel.
- Pop-up Keys: Pop-up keys are task-related keys that appear automatically on the monitor screen when needed. For example, the Confirm pop-up key appears only when you need to confirm a change.

3.6 Using the Touchscreen

Select screen items by pressing them directly on the monitor's screen. You can enable or disable touchscreen

operation by pressing and holding the [**Main Menu**] QuickKey for 3 seconds. A padlock symbol **b** displays if touchscreen operation is disabled.

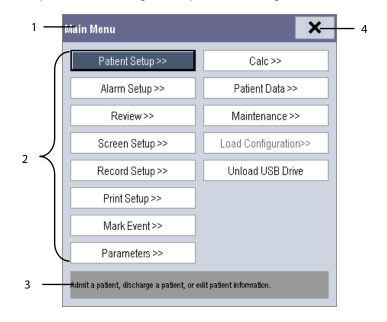
3.7 Using the On-screen Keyboard

The on-screen keyboard enables you to enter information.

- Use the ← key to delete the previously entered character.
- Use the A key to toggle between uppercase and lowercase letters.
- Select ◀ to confirm what you have entered and close the on-screen keyboard.
- Select ^{@!#} to access the symbol keyboard.
- Select ¹ to exit the symbol keyboard.

3.8 Using the Main Menu

To enter the [Main Menu], select the [Main Menu] on-screen QuickKey or the [Menu] hardkey on the monitor's front panel. Most of monitor operations and settings can be performed through this [Main Menu].



Other menus are similar to the [Main Menu] and contain the following parts:

- 1. Heading: displays the title for the current menu.
- 2. Main body: displays options, buttons, prompt messages, etc. The menu buttons with ">>" open a secondary window to reveal more options or information.
- 3. Online help area: displays help information for the highlighted menu item.
- 4. X: select to exit the current menu. If changes are made in the menu, they take effect after selecting this button.

3.9 Setting the Screen

You can enter the [Screen Setup] window as shown below by selecting [Main Menu] \rightarrow [Screen Setup] \rightarrow [Screen Layout >>]. In this window, you can arrange the positions of the parameters and waveforms. Unselected parameters or waveforms will not display.



The ECG parameter and the first ECG waveform always display in the first row. The configurable areas can be classified as Area A, Area B, and Area C.

- In Area A, you can choose to display the parameter windows and their waveforms (if one exists). Each parameter and the associated waveform are displayed in the same row.
- In Area B, you can choose to display the parameter windows. But if all parameters in area C are set to [Off], both the parameter and waveform selected in the first row of area B will be displayed.
- In Area C, you can choose to display the timer and any parameters without associated waveforms.

The screen automatically adjusts to ensure the best view based on your screen setup.

If no corresponding parameter or waveform displays after the module is inserted, perform the following inspections:

- Check the connection between the module and lead, cable, sensor, or external device.
- Enter the [Screen Setup] window for the desired display configuration by selecting [Main Menu] → [Screen Setup] → [Screen Layout >>].
- Check that the parameter is turned on in [**Parameters Switch**] window.

CAUTION

• Unallocated parameters in the [Screen Setup] window will not display. However, the monitor still sounds alarms for these parameters.

3.10 Displaying the Timer

To display the timer in the main screen:

- 1. Select [Main Menu] \rightarrow [Screen Setup>>] \rightarrow [Screen Layout >>] to access the [Screens] window.
- 2. Select [Screen Setup] tab.
- 3. In Area C, select [Timer] from the drop-down list of the desired parameter area. Refer to section **3.9 Setting the Screen**.
- 4. Select 🗙 to exit the window. The main screen will display the timer.



- Select [Start] or [Pause] to start or pause timing.
- Select [**Clear**] to clear current timing result.
- Select [Setup] to access the [Timer Setup] window, in which you can set the [Direction] to [Up] or [Down]. If you select [Down], you should set:
 - [Run Time(h:min:s)]: The available time range is 0 to 100 hours, and the default time is 5 minutes.
 - [Reminder Vol]: During the last 10 seconds of the countdown, the system issues reminder tone. The available volume range is 0 to 10. 0 means off, and 10 the maximum volume.

Timer Setup	×
Direction	Down
Run Time (h:min:s)	0 : 5 : 0
Reminder Vol	2

3.11 Changing General Settings

This chapter covers only general settings such as language, brightness, date and time, etc. Refer to measurement and other settings in their respective sections.

3.11.1 Setting up a Monitor

To install a monitor or change its location:

- 1. Select [Main Menu] \rightarrow [Maintenance >>] \rightarrow [User Maintenance >>] \rightarrow enter the required password.
- 2. In the [User Maintenance] menu, set [Monitor Name], [Department] and [Bed No.], or edit their settings.

3.11.2 Changing Language

To change the language:

- 1. Select [Main Menu] \rightarrow [Maintenance >>] \rightarrow [User Maintenance >>] \rightarrow enter the required password.
- 2. In the [User Maintenance] menu, select [Language] and then select the desired language.
- 3. Restart the monitor.

• The changed language is applied only after the monitor is restarted.

3.11.3 Adjusting the Screen Brightness

To adjust the screen brightness:

- 1. Select the [Main Menu] \rightarrow [Screen Setup >>] \rightarrow [Brightness].
- 2. Select the appropriate setting for the screen brightness. 10 is the brightest, and 1 is the dimmest.

If the monitor operates on battery power, setting a dimmer screen brightness will prolong the battery operating time. When the monitor enters Standby mode, the screen automatically changes to the dimmest setting.

3.11.4 Showing/Hiding Help Text

The monitor provides online help information. The user can display or hide the help text as needed.

- 1. Select [Main Menu]→[Screen Setup >>].
- 2. Select [Help] and select either [On] or [Off].

3.11.5 Setting the Date and Time

To set the date and time:

- 1. Select [Main Menu] \rightarrow [Maintenance >>] \rightarrow [System Time >>].
- 2. Set the date and time.
- 3. Select [Date Format] and select either [yyyy-mm-dd], [mm-dd-yyyy] or [dd-mm-yyyy].
- 4. Select [Time Format] and select either [24h] or [12h].

CAUTION

• Changing date and time affects the storage of trends and events and may cause data loss. Save or record any needed data prior to changing the date and time.

3.11.6 Adjusting Volume

Alarm Volume

To adjust the alarm volume:

- Select the [Volume Setup] QuickKey, or [Main Menu]→[Alarm Setup >>]→enter a password as required→[Others].
- 2. Select [**Alm Volume**] and then select the appropriate volume: X-10, in which X is the minimum volume, depending on the set minimum alarm volume (refer to the chapter Alarm), and 10 is the maximum volume.

Key Volume

When you press the navigation knob, the touchscreen, or the hardkeys on the panel, the monitor prompts you by making a sound of the key volume you have set.

To adjust the key volume:

- 1. Select the [Volume Setup] QuickKey, or [Main Menu]→[Screen Setup >>].
- 2. Select [Key Volume] and then select the appropriate volume. 0 means off, and 10 is the maximum volume.

QRS Volume

The QRS tone is derived from either the HR or PR, depending on which is currently selected as the alarm source in [**ECG Setup**] or [**SpO2 Setup**]. When monitoring SpO₂, there is a variable pitch tone which changes as the patient's saturation level changes. The tone pitch rises as the saturation level increases and falls as the saturation level decreases. The tone volume is user adjustable.

To adjust the QRS volume

- 1. Select the [**Volume Setup**] QuickKey, or the ECG parameter window→[**Others** >>], or the SpO₂ parameter window.
- 2. Select [**QRS Volume**] or [**Beat Vol**] and then select the appropriate volume. 0 means off, and 10 is the maximum volume.

3.12 Setting Parameters

3.12.1 Switching the Parameters On/Off

To switch the parameters on or off, select [Main Menu] \rightarrow [Screen Setup >>] \rightarrow [Screen Layout >>] \rightarrow [Parameters Switch]. When a parameter is switched off, its corresponding parameter module stops working, and its parameter value and waveform are not shown on the monitor display.

NOTE

• ECG is always selected, and you can not switch it off.

3.12.2 Accessing the Parameters Menu

Select [**Parameters** >>] from the [**Main Menu**] or select corresponding parameter area or waveform area to access a parameter setup menu.

3.13 Operating Modes

Your monitor has different operating modes. Some are password protected. This section lists the major operating modes.

3.13.1 Monitoring Mode

This is the normal, everyday working mode that you use for monitoring patients. Your monitor automatically enters the Monitoring mode after being turned on.

3.13.2 Night Mode

To avoid disturbing the patient, Night mode may be used.

To activate the Night mode:

- 1. Select the [Night Mode] QuickKey or [Main Menu] \rightarrow [Screen Setup >>] \rightarrow [Night Mode >>].
- In the pop-up menu, set the desired brightness, alarm volume, QRS volume, key volume, NIBP end tone, or whether to stop NIBP measurements or not. When [Stop NIBP] is selected, all the NIBP measurements terminate after entering the Night mode.
- 3. Select the [Enter Night Mode] button.

To cancel the Night mode:

- 1. Select the [Night Mode] QuickKey or [Main Menu]→[Screen Setup >>]→[Night Mode >>].
- 2. Select [**Ok**] in the pop-up.

CAUTION

• Before entering Night mode, confirm the brightness settings, alarm volume, QRS volume, and key volume. Consider the potential risk when the setting value is low.

3.13.3 Standby Mode

In Standby mode, you can temporarily stop patient monitoring without turning off the monitor. To enter the Standby

mode, select the [**Standby**] QuickKey

FOR YOUR NOTES

4.1 Admitting a Patient

The monitor displays physiological data and stores it in trends as soon as a patient is connected. This allows you to monitor a patient that is not admitted yet. However, it is recommended that you admit a patient so that you can clearly identify them on recordings, reports and networking devices.

To admit a patient:

- 1. Select the [Patient Setup] QuickKey, or [Main Menu]→[Patient Setup >>].
- 2. Select [**Discharge Patient**] to clear any previous patient data. If you do not erase data from the previous patient, the new patient's data is combined with the previous patient's data. The monitor makes no distinction between the old and the new patient data.
- 3. If [Discharge Patient] button appears dimmed, directly select [Admit Patient] and then select:
 - [Yes] to apply the data saved in the monitor to the new patient, or
 - [No] to clear the data saved in the monitor.
- 4. In the [**Patient Demographics**] menu, enter the demographic details, of which:
 - [Patient Cat.] allows the user to select patient size (Neo, Ped, Adu) which will determine the way your monitor processes and calculates some measurements, and what safety and alarm limits are applied for your patient.
 - [**Paced**] determines whether to show pace pulse marks on the ECG waveform. When the [**Paced**] is set to [**No**], pace pulse marks are not shown in the ECG waveform.
- 5. Select [**Ok**].

WARNING

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

4.2 Quick Admitting a Patient

Use [**Quick Admit**] only if you do not have the time or information to fully admit a patient. Complete the rest of the patient demographic details later. Otherwise, the ? symbol always displays in the patient information area.

- 1. Select the [Patient Setup] QuickKey, or [Main Menu]→[Patient Setup >>].
- 2. Select [**Quick Admit**]. If a patient has been admitted, select [**Ok**] to discharge the current patient. If no patient is admitted, you can choose either:
 - [Yes] to apply the data in your monitor to the new patient, or
 - [No] to clear any previous patient data.
- 3. Enter the Patient Category and Paced Status for the new patient, and then select [**Ok**].

4.3 Setting the Monitor Location

To set the monitor location, follow this procedure:

- 1. Select [Main Menu] \rightarrow [Maintenance >>] \rightarrow [User Maintenance >>] \rightarrow enter the required password.
- 2. Input the following location of the monitor:
 - [**Facility**]: your facility name.
 - [**Department**]: your department name.
- [Room No.]: room number.
 - [Bed No.]: bed number.

4.4 Editing Patient Information

To edit the patient information after a patient has been admitted, when the patient information is incomplete, or when you want to change the patient information:

- 1. Select the [Patient Setup] QuickKey, or [Main Menu]→[Patient Setup >>].
- 2. Select [Patient Demographics] and then make the required changes.
- 3. Select [**Ok**].

You can also input the patient's visit number in the [**Patient Demographics**] menu, but the [**Visit Number**] option needs to be enabled.

To display the [Visit Number] option in the [Patient Demographics] menu:

- 1. Select [Main Menu] \rightarrow [Maintenance >>] \rightarrow [User Maintenance >>] \rightarrow enter the required password.
- 2. Set [**Visit Number**] to [**On** >>].

4.5 Discharging a Patient

To discharge a patient:

- 1. Select the [Dicharge Patient] QuickKey, or [Main Menu] \rightarrow [Patient Setup >>] \rightarrow [Discharge Patient].
- 2. In the pop-up menu, you can either:
 - Select [Ok] to discharge the current patient, or
 - Select [Standby] then [Ok]. The monitor enters the Standby mode after discharging the current patient, or
 - Select [Cancel] to exit without discharging the patient.

If you need to print the end case report, refer to section 22.7 End Case Reports for instructions.

NOTE

After discharging an admitted patient, you can review the historical data of this patient by selecting [Main Menu]→[Patient Data >>]→[Historical Data >>]. When the SD card is full, the oldest patient data will be deleted automatically. Therefore, you cannot always view all the historical data.

4.6 Transferring Patient Data

You can transfer patient data between monitors with a USB drive without re-entering the patient demographic information. Transferring of patient data enables you to review the patient's condition history. The patient data that can be transferred includes: patient demographics, trend data, alarm events and parameter alarm limits.

From the [User Maintenance] menu, select [Others >>]. In the pop-up menu, you can set [Transferred Data Length]. The default is [4 h]. You can also set [Data Transfer Method]. The default is [Off].

WARNING

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

NOTE

 The system automatically enables the HR alarm and lethal arrhythmia alarm after transferring the patient data.

4.6.1 Transferring Data from the Monitor to a USB Drive

To transfer data from the monitor to a USB drive:

- 1. Connect a USB drive into the monitor's USB port. Select [Main Menu]→[Patient Setup >>].
- 2. Select [Transfer to Storage Medium]. In the pop-up menu, you can:
 - Select [**Ok**] to transfer the patient data, or
 - Select [**Cancel**] to exit the menu.
- 3. Wait until the following message appears: [Transfer to storage medium successful. Please remove the USB drive.].
- 4. Remove the USB drive from monitor.

4.6.2 Transferring Data from a USB Drive to the Monitor

To transfer data from a USB drive to the monitor:

- 1. Connect the USB drive to the destination monitor.
- Select [Main Menu]→[Maintenance >>]→[User Maintenance >>]→enter the required password → [Others >>]
 → [Data Transfer Method] →[USB Drive]. Then:
 - Select [**Transfer**] to transfer the patient data to the monitor, or
 - Select [Cancel Transfer] to cancel the transfer operation.
 - Select [**Unload USB Drive**] to unload the USB drive without transferring the patient data.
- 3. After you select [**Transfer**], in the pop-up menu you can select the patient data contents that need to be transferred. [**Patient Demographics**] must be selected. After [**Ok**] is selected, the monitor compares the patient information stored in both the storage medium and monitor and manages the patient data based on the following.
 - Different
 Patient:
 The monitor erases all the current patient data, transfers the patient data from the storage medium, and loads the configuration according to the patient category.
 - Same Patient: In the pop-up dialog box, you can:
 - Select [**Yes**] to merge the patient data in the monitor and storage medium.
 - Select [No] to erase all the current patient data in the monitor and transfer the patient data from the storage medium.
- 4. Wait until the message [Transfer from storage medium successful.] appears before removing the USB drive.

CAUTION

- The USB drive you use may be write-protected. In this case, make sure the USB drive is in read/write mode.
- Do not remove the storage medium during data transfer process. Otherwise, data files may be damaged.
- The normal monitoring function will be affected during data export. Do notperform any monitoring activity during data export.

5.1 Introduction

When performing continuous patient monitoring, the clinical professional often needs to adjust the monitor's settings according to the patient's condition. The collection of all these settings is called a configuration. Allowing you to configure the monitor more efficiently, the monitor provides different sets of configurations to accommodate the varying patient categories and departments. You can change the settings from a default configuration and then save it as a user configuration.

The default configurations provided for your monitor are department-oriented. You can choose either from:

- General
- OR
- ICU
- NICU
- CCU

Each department has three different sets of configurations: one for each patient category [**Neo**], [**Ped**], and [**Adu**]. For the important configuration items and their default values and user maintenance items, see *Appendix C Default Configurations*.

NOTE

• The configuration management function is password protected. The configuration management tasks must be performed by authorized personnel. Please contact your department manager or biomedical engineering department for the password used at your facility.

5.2 Entering the [Manage Configuration] Menu

To access the [Manage Configuration] menu:

- 1. Press the [Menu] hardkey on the monitor's front panel to enter the [Main Menu].
- 2. Select [Maintenance >>]→[Manage Configuration >>]. Enter the required password and then select [Ok].

Manage Configuration	×	
Donartmont	Select Default Config. >>	
Department:	Select Delduit Connig. ~~	
	Save Current Settings As>>	
General	Edit Config.>>	
	Delete Config.>>	
Change Department >>	Export Config. >>	
	Import Config.>>	
	M IT D IV	
	Modify Password >>	
Change the department.		

5.3 Changing Department

If the current department configuration is not the one you want to view, you can select [**Change Department** >>] in the [**Manage Configuration**] menu and then choose the one you want for viewing as shown below.

Select Department			
General			
D OR			
D NICU			
CCU			
Ok	Cancel		
Select a department for configuration management.			

NOTE

• Changing the department will delete all current user configurations.

5.4 Setting Default Configuration

The monitor will load the pre-set default configuration in the following cases:

- The monitor restarts after being switched off for more than 120 seconds.
- A patient is admitted.
- A patient is discharged.
- Patient data is cleared.
- Patient category is changed.

To set the default configuration:

- 1. Select [Select Default Config. >>] in the [Manage Configuration] menu.
- 2. In the [Select Default Config.] menu, select [Load the Latest Config.] or [Load Specified Config.].

When you select [**Load Specified Config.**], the restored configuration is subject to the patient category (adult, pediatric or neonate). This configuration can be either factory configuration or a saved user configuration. As an example, select [**Default Adu Config.**] and then select [**Defaults**] or user configuration(s).

When you select [Load the Latest Config], the latest configuration is loaded when the monitor is started or a patient is admitted.

NOTE

• To identify which configuration is restored when the monitor starts, enter the main screen to check the prompt information at the lower part of the screen (displayed for about 10 seconds).

5.5 Saving Current Settings

Current settings can be saved as a user configuration. Up to 5 user configurations can be saved.

To save current settings:

- 1. Select [Save Current Settings As >>] in the [Manage Configuration] menu.
- 2. In the pop-up dialog box, enter the configuration name and then select [**Ok**]. The current settings are saved as a user configuration with the name you entered.

5.6 Editing Configurations

To edit an existing configuration:

1. Select [Edit Config. >>] in the [Manage Configuration] menu. The following menu appears.

Edit Config.	
Defaults(Adu)	
Defaults(Ped)	
Defaults(Neo)	
	Config. on USB drive>>
Edit	Back
Select a configuration to edit.	

 The pop-up menu shows the existing monitor configurations. Selecting [Config. on USB drive >>] will show the existing configurations on the USB drive. Select the desired configuration and then select the [Edit] button. The following menu appears.

Edit ConfigDefaults				
Patient Cat.		Adu		
	Alarm Setup>>			
	Screen Setup >>			
	Parameters >>			
Save	Save as		Back	
Edit alarm settings of this configuration.				

- 3. Select [Alarm Setup >>], [Screen Setup >>] or [Parameters >>] to enter the corresponding menu in which settings can be changed. The changed alarm setup items will be marked in red.
- 4. You can select [**Save**] or [**Save as**] to save the changed configuration. Select [**Save**] to overwrite the original configuration. Select [**Save as**] to save the changed configuration as another file with another name.

5.7 Deleting a Configuration

To delete a configuration:

- 1. Select [**Delete Config.** >>] in the [**Manage Configuration**] menu.
- The pop-up menu shows the existing monitor's user configurations. Selecting [Config. on USB drive >>] will show the existing user configurations on the USB drive. Select the user configurations you want to delete and then select [Delete].
- 3. Select [Yes] in the pop-up.

5.8 Loading a Configuration

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

To load a configuration,

- 1. Select [Load Configuration >>] from the [Main Menu]. The pop-up menu shows the existing configurations on the monitor.
- 2. Selecting [**Config. on USB drive** >>] to show the existing configurations on the USB drive.
- 3. Select a desired configuration.
- 4. Select [View] to see the configuration details. In the pop-up menu, you can select [Alarm Setup >>], [Screen Setup >>] or [Parameter >>] to view the corresponding contents. The alarm setup items which are different than those currently used are marked in red.
- 5. Select [Load].

NOTE

• The monitor may configure some settings by default when you load a configuration of different software version with the current configuration.

5.9 Restoring the Latest Configuration Automatically

During operation, you may make changes to some settings. However, these changes may not be saved as a user configuration. To prevent the changes from being lost in case of a sudden power failure, the monitor stores the configuration in real time. The saved configuration is the latest configuration.

The monitor restores the latest configuration if it restarts within 60 seconds after the power failure. And it will restore the default configuration rather than the latest configuration if it restarts 120 seconds after the power failure. The monitor loads either the latest configuration or the default configuration if it restarts from 60-120 seconds after the power failure.

FOR YOUR NOTES

6.1 Configuring Your Screens

You can configure your monitor's screens by setting:

- Waveform line size
- The color in which each measurement's numerics and waveforms are displayed
- The parameter to monitor.

CAUTION

• Changing some settings may be hazardous. Therefore, those settings are password-protected and can be modified by authorized personnel only. Once a change is made, notify those who use the monitor.

6.1.1 Changing the Waveform Line Size

To change the waveform line size:

- 1. Select [Main Menu] \rightarrow [Maintenance >>] \rightarrow [User Maintenance >>] \rightarrow enter the required password.
- 2. Select [Others >>].
- 3. Select [Wave Line] and select [Thick], [Med] or [Thin].

6.1.2 Changing Measurement Colors

To change the measurement colors:

- 1. Select [Main Menu] \rightarrow [Screen Setup >>] \rightarrow [Measurement Color Setup >>].
- 2. Select the color box next to your desired parameter and then select a color from the pop-up menu.

6.1.3 Changing Screen Layout

Select the [Screens] QuickKey, or [Main Menu] \rightarrow [Screen Setup >>] \rightarrow [Screen Layout >>] to enter the [Screens] menu.

- You can choose the desired screen type in the [**Choose Screen**] window.
- You can select the parameters and waveforms you want to view in the [Screen Setup] window. For details, please refer to the section 3.9 Setting the Screen.
- You can select the parameters you want to view on the big numerics screen in the [Big Numerics Screen Setup] window.
- You can switch the connected parameter modules on or off in the [Parameters Switch] window. If a parameter module is switched off, parameter values and waveforms will not display on the screen.

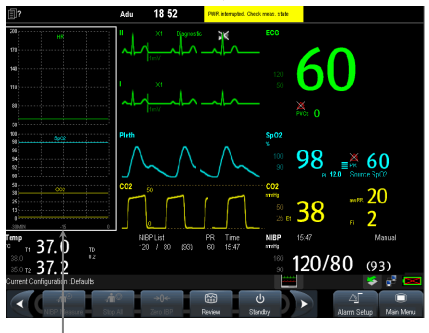
6.2 Viewing Minitrends

6.2.1 Having a Split-Screen View of Minitrends

You can split the normal screen so that the left hand side continuously shows graphic minitrends beside waveforms as shown in the figure below.

To have a split-screen view of minitrends:

- Select the [Minitrends] QuickKey, or
- Select the [Screens] QuickKey→[Choose Screen] → [Minitrends Screen] → X, or
- Select [Main Menu]→[Screen Setup >>]→[Screen Layout >>]→[Choose Screen]→ [Minitrends Screen]→X.



Minitrends View

The split-screen view provides minitrends for multiple parameters. In each field, the label and scale are respectively displayed at the top and left. The time is displayed at the bottom of the minitrends view as shown below.

200 [NIBP	
120		· · · · · · · · · · · · · · · · · · ·
40		
→ -2H	-1	0

6.2.2 Setting Minitrends

Select the minitrends area. From the pop-up [Minitrend Setup] menu:

■ Select the parameters to be displayed, or

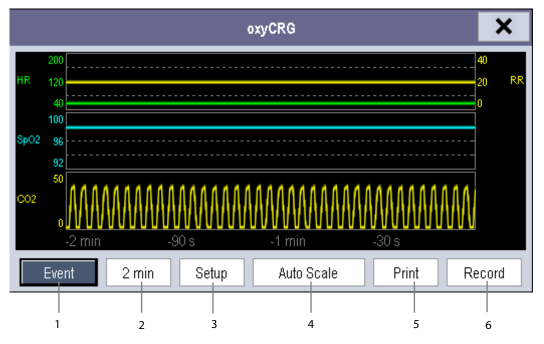
Time

■ Select [Minitrend Length] and then select the appropriate setting.

6.3 Viewing OxyCRG

To have a split screen view of oxyCRG:

- Select the [OxyCRG] QuickKey;
- Select the [Screens] QuickKey→[Choose Screen]→ [OxyCRG Screen]→ \times ; or,
- Select [Main Menu] → [Screen Setup >>] → [Screen Layout >>] → [Choose Screen] → [OxyCRG Screen] → X.



The split-screen view covers the lower part of the waveform area and shows HR trend, SpO_2 trend, RR trend, and a compressed waveform (Resp or CO_2 waveform). At the bottom, there are controls:

1. Event

You can enter the [Review] menu by selecting the [Event] button.

2. Trend length list box

In the trend length list box, you can select [1 min], [2 min], [4 min], or [8 min].

3. Setup

Select the [**Setup**] button to enter the [**Setup**] menu, in which you can select the parameters for display, the time length to be saved before and after an event, and the scale of the graphic trends and waveform. The trend area can display two parameter trends, e.g. HR trend and RR trend, simultaneously.

4. Auto Scale

Select the [Auto Scale] button, and the system automatically adjusts the scaling.

5. Print

Select [Print] to print the real-time OxyCRG to a printer.

6. Record

Select [Record] to print the currently displayed OxyCRG trends to the recorder.

6.4 Viewing Other Patients

6.4.1 Care Group

This function lets you:

- View information on the monitor screen from another bed in the same Care Group.
- Be notified of physiological and technical alarm conditions at the other beds in the same Care Group.

To have a Care Group:

- 1. Open the [**View Other Patient**] window by:
 - Selecting [Others] QuickKey, or
 - Selecting [Screens] QuickKey \rightarrow [Choose Screen] \rightarrow [View Others Screen] \rightarrow X, or
 - ♦ Selecting [Main Menu]→[Screen Setup >>]→[Screen Layout >>]→[Choose Screen]→[View Others Screen]→X.
- 2. Select [Setup] in the [View Other Patient] window.
- 3. Select the desired monitors from the [**Connected Monitor List**], and then select the X button. The selected monitors constitute a Care Group.

6.4.2 Viewing the Care Group Overview Bar



The Care Group overview bar is located at the bottom of the [**View Other Patient**] window. In the overview bar, the department and bed label for any Care Group beds are displayed. For telemetry, # is displayed before the department label. The color in which a Care Group bed appears matches its status:

- Red: indicates the bed is having high-level physiological alarms or the telemetry is in alarm, such as nurse call or event.
- Yellow: indicates the bed is having medium-level or low-level physiological alarms, or medium-level technical alarms.
- Blue: indicates the bed is having low-level technical alarms.
- Light grey: indicates the bed fails to be networked.
- Dark grey: indicates the bed is in the Standby mode.

You can view a Care Group bed's alarms by selecting it from the Care Group, and you can select the [View This Patient] button to view the bed in the [View Other Patient] window.

For more details about Care Group alarms, refer to the chapter 24 Batteries.

6.4.3 Understanding the View Other Patient Window

When you first open the [View Other Patient] window, the monitor automatically selects a device from the network to display in the [View Other Patient] window.



The [View Other Patient] window covers the lower part of the waveform area and consists of:

- 1. Information Area: shows the patient information (including department, bed number, patient name, etc.), network status symbol.
- View Area: shows physiological waveforms and parameters. You can switch a waveform area to a parameter area by selecting your desired waveform area and then selecting [Switch to Parameter Area], or switch a parameter area to a waveform area by selecting your desired parameter area and then selecting [Switch to Waveform Area].
- 3. Care Group Overview Bar. See section 6.4.2 Viewing the Care Group Overview Bar for information.
- 4. Message Area: shows physiological, technical and prompt messages from the currently viewed monitor. It also shows the alarm given by the device such as nurse call or event. By selecting this area, you can enter the [Alarm Information List] to view all physiological, technical and prompt messages coming from the currently viewed patient.
- [Alarm Reset] button: resets alarms from the currently viewed bed. Refer to section 7.8 Resetting Alarms.
 When the [Reset Other Bed's Alarms] is disabled, no button will appear here.

Additionally, you can change a waveform or parameter for viewing:

- To change a waveform for viewing, select the waveform segment where you want a new waveform to appear and then select the waveform you want from the pop-up menu.
- To change a parameter for viewing, select the parameter window where you want a new parameter to appear and then select the parameter you want from the pop-up menu.

WARNING

- The data presented in the [View Other Patient] window has a delay. Do not rely on this window for realtime data.
- The 👿 icon indicates network disconnection. You cannot view other monitors when this icon is displayed.

6.5 Understanding the Big Numerics Screen

To enter the big numerics screen:

- 1. Select the [Screens] QuickKey, or [Main Menu] \rightarrow [Screen Setup >>] \rightarrow [Screen Layout >>].
- 2. Select [**Big Numerics**] \rightarrow \times .



You can select your desired parameters to display in this screen: select the [Screens] QuickKey \rightarrow [Big Numerics Screen Setup] and then select the parameters you want to view. For parameters having a waveform, the waveform will also be displayed.

Alarms, triggered by an abnormal vital sign or technical issue with the monitor, as visually and audibly indicated to the user.

WARNING

• A potential hazard can exist if different alarm presets are used for the same or similar equipment in any single area, e.g. an intensive care unit or cardiac operating room.

For troubleshooting specific alarms, see appendix **D** Alarm Messages.

7.1 Alarm Categories

The monitor's alarms can be classified into two categories: physiological alarms and technical alarms.

1. Physiological alarms

Physiological alarms, also called patient status alarms, are triggered by a monitored parameter value that violates set alarm limits or an abnormal patient condition. Physiological alarm messages are displayed in the physiological alarm area.

2. Technical alarms

Technical alarms, also called system status alarms, are triggered by a device malfunction or a patient data distortion due to improper operation or mechanical problems. Technical alarm messages are displayed in the technical alarm area.

Apart from the physiological and technical alarm messages, the monitor will show some messages telling the system status or patient status. Messages of this kind are included in the prompt message category and usually displayed in the prompt information area. Some prompt messages that indicate the arrhythmia events are displayed in the physiological alarm area. For some measurements, their related prompt messages are displayed in their respective parameter windows.

7.2 Alarm Levels

The monitor's alarms can be classified into three severity categories: high level, medium level and low level.
--

	Physiological alarms	Technical alarms
High level	Indicate that the patient is in a life	Indicate a severe device malfunction or an improper operation
	threatening situation, such as Asystole,	which could make it possible that the monitor cannot detect
	Vfib/Vtac and so forth, and emergency	critical patient status and thus threaten the patient's life.
	treatment is required.	
Medium	Indicate that the patient's vital signs	Indicate a device malfunction or an improper operation, which
level	appear abnormal and immediate	may not threaten the patient's life, but may compromise the
	treatment is required.	monitoring of vital physiological parameters.
Low level	Indicate that the patient's vital signs	Indicate a device malfunction or an improper operation which
	appear abnormal and immediate	may compromise a certain monitoring function, but will not
	treatment may be required.	threaten the patient's life.

7.3 Alarm Indicators

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

- Alarm lamp
- Alarm message
- Flashing numerics
- Audible alarm tones

7.3.1 Alarm Lamp

If a technical alarm or physiological alarm occurs, the alarm lamp will flash. The color and flashing frequency match the alarm level as follows:

- High level alarms: the lamp quickly flashes red.
- Medium level alarms: the lamp slowly flashes yellow.

*

Low level alarms: the lamp lights yellow without flashing.

7.3.2 Alarm Messages

When an alarm occurs, an alarm message will appear in the technical or physiological alarm area. For physiological alarms, the asterisk symbols (*) before the alarm message match the alarm level as follows:

- High level alarms: ***
- Medium level alarms: **
- Low level alarms:

Additionally, the alarm message uses different background color to match the alarm level:

- High level alarms: red
- Medium level alarms: yellow
- Low level alarms: yellow

You can view the alarm messages by selecting the physiological or technical alarm area.

7.3.3 Flashing Numeric

If an alarm triggered by an alarm limit violation occurs, the numeric of the measurement in alarm will flash every second, and the corresponding alarm limit will also flash at the same frequency indicating the high or low alarm limit is violated.

7.3.4 Audible Alarm Tones

The alarm tone is distinct from heart beat tone, keystroke tone and pulse tone in frequency. This monitor has three alarm tones and patterns: ISO, Mode 1 and Mode 2. For each pattern, the alarm tones identify the alarm levels as follows:

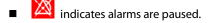
- ISO pattern:
 - ◆ High level alarms: triple+double+triple+double beep
 - Medium level alarms: triple beep
 - Low level alarms: single beep
- Mode 1:
 - High level alarms: high-pitched single beep
 - Medium level alarms: double beep
 - ◆ Low level alarms: low-pitched single beep
- Mode 2:
 - High level alarms: high-pitched triple beep
 - Medium level alarms: double beep
 - ◆ Low level alarms: low-pitched single beep

NOTE

- When multiple alarms of different levels occur simultaneously, the monitor will select the highest level alarm, light the alarm lamp and give alarm sounds accordingly, while all the alarm messages are displayed circularly on the screen.
- Some physiological alarms, such as Asystole, are exclusive. They have identical alarm tones and alarm lights
 with normal high level physiological alarms, but their alarm messages are displayed exclusively. That is,
 when an exclusive physiological alarm and a normal high level physiological alarm are triggered
 simultaneously, only exclusive physiological alarm messages will be displayed.

7.3.5 Alarm Status Symbols

Apart from the aforementioned alarm indicators, the monitor still uses the following symbols telling the alarm status:



- indicates alarm is reset.
- indicates the alarm sound is turned off.
- Indicates individual measurement alarms are turned off or the system is in alarm off status.

7.4 Alarm Tone Configuration

7.4.1 Setting the Minimum Alarm Volume

To set the minimum alarm volume:

- 1. Select [Main Menu] \rightarrow [Maintenance >>] \rightarrow [User Maintenance >>] \rightarrow enter the required password.
- 2. Select [Alarm Setup >>] to enter the [Alarm Setup] menu.
- 3. Select [Minimum Alarm Volume] and then select a value between 0 and 10.

The minimum alarm volume refers to the minimum value you can set for the alarm volume, which is not affected by user or factory default configurations. The minimum alarm volume setting remains unchanged when the monitor shuts down and restarts.

7.4.2 Changing the Alarm Volume

To change the alarm volume:

- Select the [Volume Setup] QuickKey or the [Alarm Setup] QuickKey→[Others], or [Main Menu]→[Alarm Setup >>]→[Others].
- 2. Select the appropriate volume from [**Alm Volume**]: X to 10, in which X is the minimum volume, depending on the set minimum alarm volume, and 10 the maximum volume.
- 3. Select [High Alarm Volume] to set the high priority alarm volume as [Alm Volume+0], [Alm Volume+1] or [Alm Volume+2].
- 4. Select [Reminder Vol] to set the volume of the reminder tone as [High], [Med] or [Low].

When alarm volume is set to 0, the alarm sound is turned off and a 🔀 symbol appears in the alarm symbols area of the screen.

7.4.3 Setting the Interval Between Alarm Sounds

You cannot change the interval between alarm tones if you choose mode 1 or 2 as your desired alarm tone pattern. For these two patterns, the interval between alarm tones identifies the alarm levels as follows:

- Mode 1:
 - Interval between high level alarm tones: continuously
 - Interval between medium level alarm tones: 5 s
 - Interval between low level alarm tones: 20 s
- Mode 2:
 - Interval between high level alarm tones:
 1 s
 - Interval between medium level alarm tones: 5 s
 - Interval between low level alarm tones: 20 s

If you choose the ISO pattern, you can change the interval between alarm tones. To change the interval between alarm tones:

- 1. Select [Main Menu] \rightarrow [Maintenance >>] \rightarrow [User Maintenance >>] \rightarrow enter the required password.
- 2. Select [Alarm Setup >>] to enter the [Alarm Setup] menu.
- 3. Select [High Alarm Interval (s)], [Med Alarm Interval (s)] and [Low Alarm Interval (s)] in order and then select the appropriate settings.

WARNING

- When the alarm sound is switched off, the monitor will give no audible alarm tones even if a new alarm occurs. Therefore the user should be very careful about whether to switch off the alarm sound or not.
- Do not rely exclusively on the audible alarm system for monitoring. Adjustment of alarm volume to a low level may result in a hazard to the patient. Always keep the patient under close surveillance.

7.4.4 Changing the Alarm Tone Pattern

To change the alarm tone pattern:

- 1. Select [Main Menu] \rightarrow [Maintenance >>] \rightarrow [User Maintenance >>] \rightarrow enter the required password.
- 2. Select [Alarm Setup >>] to enter the [Alarm Setup] menu.
- 3. Select [Alarm Sound] and then select [ISO], [Mode 1] or [Mode 2].

NOTE

• User or factory default configurations have no impact on the alarm tone pattern setup. The alarm tone pattern remains unchanged after the monitor restarts.

7.4.5 Setting the Reminder Tones

When the reminder tone is switched on and a physiological or technical alarm condition exists, the monitor gives a periodic reminder tone in the following cases:

- When the alarm volume is set to zero, a high, medium, low, or technical audible alarm indication will be generated at the set reminder interval until the alarm condition is cleared.
- When the alarm condition has been acknowledged (e.g. pressing the Alarm Reset button), a high, medium, or low audible alarm indication will be generated at the set reminder interval until the alarm condition is cleared.
- When the alarm system has been permanently paused (e.g. pressing the Alarm Pause button with the Alarm Pause Time set to Permanent), an Alarm Paused audible alarm indication will be generated at the set reminder interval until the alarm condition is cleared or the alarm system re-enabled.

To turn on the reminder tones:

- 1. Select [Main Menu] \rightarrow [Maintenance >>] \rightarrow [User Maintenance >>] \rightarrow enter the required password.
- 2. Select [Alarm Setup >>] to enter the [Alarm Setup] menu.

 Set the [Reminder Tones] to [On], [Off] or [Re-alarm]. When [Re-alarm] is selected, the acknowledged physiological alarms and technical alarms marked with "√" will be re-generated after the [Reminder Interval] if the alarm condition persists.

To set the interval between reminder tones, select [Reminder Interval] and then select [1min], [2min] or [3min].

In addition, you can set the volume of alarm reminder tones. To set the volume of alarm reminder tones, select [Main Menu] \rightarrow [Alarm Setup >>] \rightarrow [Others] or the [Alarm Setup] QuickKey \rightarrow [Others]. Then, select [Reminder Vol] and then select [High], [Medium] or [Low].

7.5 Understanding the Alarm Setup Menu

Select the [Alarm Setup] QuickKey or [Main Menu]→[Alarm Setup >>] to enter the [Alarm Setup], where you can:

- Set alarm properties for all parameters.
- Change arrhythmia alarm settings.
- Set the threshold for some arrhythmia alarms.
- Change other settings.

Refer to the chapter **8** Monitoring ECG for how to change arrhythmia alarm settings and how to set the threshold for some arrhythmia alarms.

7.5.1 Setting Alarm Properties for All Parameters

In the [Main Menu], select [Alarm Setup >>] \rightarrow [Parameters]. You can review and set alarm limits, alarm switches, alarm level and alarm recordings for all parameters.

When a measurement alarm occurs, automatic recording of all the measurement numerics and related waveforms is possible when the measurement's [**On/Off**] and [**Record**] are set on.

WARNING

- Make sure that the alarm limit settings are appropriate for your patient before monitoring.
- Setting alarm limits to extreme values may cause the alarm system to become ineffective. For example, high oxygen levels may predispose a premature infant to retrolental fibroplasia. If this is a consideration, do NOT set the high alarm limit to 100%, which is equivalent to switching the alarm off.
- When monitoring patients that are not continuously attended by a clinical operator, properly configure the alarm system and adjust alarm settings as per the patient's condition.

7.5.2 Adjusting Alarm Limits Automatically

The monitor can automatically adjust alarm limits according to the measured vital signs. When auto limits are selected, the monitor calculates safe auto limits based on the latest measured values.

To get accurate auto alarm limits, you need to collect a set of measured vital signs as a baseline. Then, in the [Main Menu], select [Alarm Setup >>] \rightarrow [Parameters] \rightarrow [Auto Limits] \rightarrow [Ok]. The monitor will create new alarm limits based on the measured values.

Before applying these automatically created alarm limits, confirm if they are appropriate for your patient in the [Alarm Setup Menu]. If not, you can adjust them manually. These alarm limits will remain unchanged until you select auto limits again or adjust them manually.

		Low alarm limit		High alarm limit			
Module	Parameter	Adult/ pediatric	Neonate	Adult/ pediatric	Neonate	Auto alarm limits range	
ECG	HR/PR	(HR × 0.8) or 40 bpm (whichever is greater)	(HR – 30) or 90 bpm (whichever is greater)		bpm (whichever is	Adult/pediatric: 35 to 240 Neonate: 55 to 225	
Resp	RR	5	(RR – 10) or 30 rpm	(RR × 1.5) or 30 rpm (whichever is	(RR + 25) or 85 rpm	Adult/pediatric: 6 to 55 Neonate: 10 to 90	
SpO ₂	SpO ₂	Same as the	Same as the default alarm limit	Same as the	Same as the default	Same as the measurement range	
	NIBP-S	(SYS × 0.68 + 10) mmHg	(SYS – 15) or 45 mmHg (whichever is greater)	(SYS × 0.86 + 38) mmHg	-	Adult: 45 to 270 Pediatric: 45 to 185 Neonate: 35 to 115	
NIBP	NIBP-D	(Dia × 0.68 + 6) mmHg	(Dia – 15) or 20 mmHg (whichever is greater)	(Dia × 0.86 + 32) mmHg	mmHg (whichever	Adult: 25 to 225 Pediatric: 25 to 150 Neonate: 20 to 90	
	NIBP-M	(Mean × 0.68 + 8) mmHg	(Mean – 15) or 35 mmHg (whichever is greater)	(Mean × 0.86 + 35) mmHg	mmHg (whichever	Adult: 30 to 245 Pediatric: 30 to 180 Neonate: 25 to 105	
	T1	(T1 – 0.5)°C	(T1 – 0.5) °C	(T1 + 0.5)°C	(T1 + 0.5)°C	1 to 49°C	
Tomor	T2	(T2 – 0.5)°C	(T2 – 0.5) °C	(T2 + 0.5)℃	(T2 + 0.5)℃	1 to 49°C	
Temp	TD	Same as the default alarm limit	Same as the default alarm limit		Same as the default alarm limit	Same as the measurement range	
IBP: ART/ Ao/ UAP/	IBP-S	(SYS × 0.68 + 10) mmHg	(SYS – 15) or 45 mmHg (whichever is greater)	(SYS × 0.86 + 38) mmHg	-	Adult: 45 to 270 Pediatric: 45 to 185 Neonate: 35 to 115	
BAP/ FAP/ LV/	IBP-D	(Dia × 0.68 + 6) mmHg	(Dia – 15) or 20 mmHg (whichever is greater)	(Dia × 0.86 + 32) mmHg	mmHg (whichever	Adult: 25 to 225 Pediatric: 25 to 150 Neonate: 20 to 90	
P1-P4 (Arterial pressure)	IBP-M	(Mean × 0.68 + 8) mmHg	(Mean – 15) or 35 mmHg (whichever is greater)	(Mean × 0.86 + 35) mmHg	mmHg (whichever	Adult: 30 to 245 Pediatric: 30 to 180 Neonate: 25 to 105	
	IBP-S	SYS × 0.75	SYS × 0.75	SYS × 1.25	SYS × 1.25		
РА	IBP-D	Dia × 0.75	Dia × 0.75	Dia × 1.25	Dia × 1.25	3 to 120 mmHg	
	IBP-M	Mean × 0.75	Mean × 0.75	Mean × 1.25	Mean × 1.25		

The monitor calculates the auto limits based on the following guidelines:

		Low alarm limit		High alarm limit		
Module	Parameter	Adult/		Adult/		Auto alarm limits
		pediatric	Neonate	pediatric	Neonate	range
IBP:						
CVP/						
ICP/						
LAP/						
RAP/	IBP-M	Mean × 0.75	Mean × 0.75	Mean × 1.25	Mean × 1.25	3 to 40mmHg
UVP/						
P1-P4						
(Venous						
pressure)						
		0 to 32 mmHg:	0 to 32 mmHg:	0 to 32 mmHg:	0 to 32 mmHg:	
		remains the same	remains the same	remains the same	remains the same	
		32 to 35 mmHg:	32 to 35 mmHg:	32 to 35 mmHg:	32 to 35 mmHg:	
		29 mmHg	29 mmHg	_	41 mmHg	
		35 to 45 mmHg:	35 to 45 mmHg:	35 to 45 mmHg:	35 to 45 mmHg:	Same as the
	EtCO ₂	(etCO ₂ -6) mmHg	(etCO ₂ -6) mmHg	$(etCO_2+6)$ mmHg	5	measurement range
CO ₂		45 to 48 mmHg:	45 to 48 mmHg:	_	45 to 48 mmHg:	
		39 mmHg	39 mmHg	-	51 mmHg	
		> 48 mmHg:	> 48 mmHg:	> 48 mmHg:	> 48 mmHg:	
		remains the same	remains the same	remains the same		
	FiCO ₂	N/A			Same as the default	Same as the
				default alarm limit		measurement range
	awRR	awRR $ imes$ 0.5 or 6	(awRR – 10) or 30			Adult/pediatric: 6 to
		rpm (whichever is	rpm (whichever is	rpm (whichever is	rpm (whichever is	55
		greater)	greater)	smaller)	smaller)	Neonate:10 to 90
	EtCO ₂ (AG)	Same as CO ₂ module				
	FiCO ₂ (AG)				r	
		awRR $ imes$ 0.5 or 6	awRR – 10 or 30	awRR $ imes$ 1.5 or 30	awRR + 25 or 85	Adult/pediatric: 6 to
	awRR	rpm (whichever is	rpm (whichever is	rpm (whichever is	rpm (whichever is	55
AG		greater)	greater)	smaller)	smaller)	Neonate: 10 to 90
	FiAA/	Same as the	Same as the	Same as the	Same as the default	Same as the
	EtAA	default alarm limit	default alarm limit	default alarm limit	alarm limit	measurement range
	FiO ₂ /	Same as the	Same as the	Same as the	Same as the default	Same as the
	EtCO ₂	default alarm limit	default alarm limit	default alarm limit	alarm limit	measurement range
	FiN ₂ O/	Same as the	Same as the	Same as the	Same as the default	Same as the
	EtN ₂ O	default alarm limit	default alarm limit	default alarm limit	alarm limit	measurement range
C.O.	BT	Adult:		Adult:		Same as the
		(BT – 1)°C	N/A	(BT – 1)°C	N/A	measurement range

7.5.3 Setting Alarm Delay Time

You can set the alarm delay time for alarms of continuously measured parameters. If the alarm condition is resolved within the delay time, the monitor will not sound the alarm.

To set the alarm delay time:

- 1. Select [Main Menu]→[Maintenance >>]→[User Maintenance >>]. Enter the required password and then select [Ok].
- 2. Select [Alarm Setup >>] \rightarrow [Alarm Delay].

Alarm delay is not applied to the following physiological alarms:

- Apnea
- Arrhythmia alarms
- ECG Weak Signal
- Resp Artifact
- SpO₂ Desat
- No Pulse
- Nellcor SpO₂ over alarm limits
- FiO₂ Shortage
- Measurements of noncontinuous parameters over alarm limits
- HR over alarm limits
- Anesthetic Mixture's MAC > 3

You can set [Apnea Delay] separately.

7.5.4 Setting SpO₂ Technical Alarm Delay

You can set the [**Tech. Alarm Delay**] in the [**Others**] tab of the [**Alarm Setup**] menu. The options are [**Off**], [**5** s], [**10** s] and [**15** s]. The delay is effective to the following technical alarms: SpO₂ Sensor Off, SpO₂ Too Much Light, SpO₂ Low Signal and SpO₂ Interference.

7.5.5 Setting Recording Length

You can change the length of the recorded waveforms. In the [**Others**] window of the [**Alarm Setup**] menu, select [**Recording Length**] and then select [**8 s**], [**16 s**] or [**32 s**]:

- **8 s**]: 4 seconds respectively before and after the alarm or manual event trigger moment.
- **If s**: 8 seconds respectively before and after the alarm or manual event trigger moment.
- **32** s]: 16 seconds respectively before and after the alarm or manual event trigger moment.

7.5.6 Entering CPB Mode (Cardiopulmonary Bypass Mode)

When performing CPB, you can put the monitor in CPB mode in order to reduce unnecessary alarms. The CPB mode is activated only if you set the department to [**OR**]. To set the department to [**OR**]:

- 1. Press the [Menu] hardkey on the monitor's front panel to enter [Main Menu].
- 2. Select [Maintenance >>]→[Manage Configuration >>]. Enter the required password and then select [Ok].
- 3. Select [Change Department >>] \rightarrow [OR].

In the CPB mode, all the physiological alarms, technical alarms and prompt alarms are switched off. In the CPB mode, [**CPB Mode**] is displayed in the physiological alarm area with a red background color.

To enter CPB mode:

- 1. Select the [CPB Mode] Quickkey or select [Enter CPB Mode] in the [Others] window of the [Alarm Setup] menu.
- 2. Then select [**Ok**] in the pop-up dialog box.

7.6 Pausing Alarms

If you want to temporarily prevent alarms from sounding, you can pause alarms by pressing the [Alarm Pause] hardkey on the monitor's front panel. When alarms are paused:

- No alarm lamps flash and no alarms are sounded.
- No numerics or alarm limits flash.
- No alarm messages are shown.
- The remaining pause time is displayed in the physiological alarm area.
- The alarms paused symbol A is displayed in the sound symbol area.

If the time interval of the monitor's last shutdown from this start-up is greater than 2 minutes, the monitor enters into the alarm paused status as soon as it is turned on. The alarm pause time is fixed to be 2 minutes.

When the alarm pause time expires, the alarm paused status is automatically cancelled and the alarm tone will sound. You can also cancel the alarm paused status by pressing the [**Alarm Pause**] hardkey.

You can set the alarm pause time to [1 min], [2 min], [3 min], [5 min], [10 min], [15 min], or [Permanent]. The default alarm pause time is 2 minutes.

- 1. Select [Main Menu] \rightarrow [Maintenance >>] \rightarrow [User Maintenance >>] \rightarrow enter the required password.
- 2. Select [Alarm Setup >>] \rightarrow [Alarm Pause Time] and then select the appropriate setting from the pop-up list.

7.7 Switching Off All Alarms

If [Alarm Pause Time] is set to [Permanent], the monitor enters into the alarm off status after the [Alarm Pause] hardkey is pressed. During the alarm off status:

- For physiological alarms, no alarm lamps flash and no alarms are sounded.
- For physiological alarms, no numerics or alarm limits flash.
- No physiological alarm messages are shown.
- [Alarm Off] is displayed in the physiological alarm area with a red background.
- For technical alarms, no alarms are sounded.
- The alarm off symbol is displayed in the alarm symbol area.

You can cancel the alarm off status by pressing the [Alarm Pause] hardkey.

WARNING

• Pausing or switching off alarms may result in a hazard to the patient.

7.8 Resetting Alarms

By selecting the 2 QuickKey, you can reset the alarm system to acknowledge the on-going alarms and enable the alarm system to respond to a subsequent alarm condition.

For physiological alarms, except the NIBP-related alarms, when the alarm system is reset:

- The alarm sound is silenced.
- A $\sqrt{}$ appears before the alarm message, indicating that the alarm is acknowledged.
- The icon 2 appears in the alarm symbol area.
- The parameter numeric and alarm limits still flash.

The indication of alarm lamp for the physiological alarm depends on the alarm light setting.

- When [Alarm Light on Alarm Reset] is set to [On], the alarm lamp remains flashing.
- When [Alarm Light on Alarm Reset] is set to [Off], the alarm lamp stops flashing.

To set [Alarm Light on Alarm Reset]:

- 1. Select [Main Menu] \rightarrow [Maintenance >>] \rightarrow [User Maintenance >>] \rightarrow enter the required password.
- 2. Select [Alarm Setup >>] to enter the [Alarm Setup] menu.
- 3. Select [Alarm Light on Alarm Reset], and then select [On] or [Off].

The default setting for [Alarm Light on Alarm Reset] is [On].

Technical alarms give different alarm indicators when the alarm system is reset:

- For some technical alarms, including the NIBP-related alarms, a √ appears before the alarm message and appears in the alarm symbol area, indicating that the alarm is acknowledged. The indication of the alarm lamp depends on the alarm light setting.
- Some technical alarms are changed to the prompt messages.
- Some technical alarms are cleared. The monitor gives no alarm indications.

For details about the indications of technical alarms when the alarm system is reset, refer to appendix **D.2 Technical Alarm Messages**.

7.9 Latching Alarms

The latching setting for physiological alarms defines how alarm indicators behave if you do not reset the alarms.

- If you do not "latch" the physiological alarms, their alarm indications disappear when the alarm condition ends.
- If you "latch" physiological alarms, all visual and audible alarm indications remain until you reset the alarms. For latched alarms the time when the alarm is last triggered is displayed beside the alarm message.

You can separately latch the visual indications or simultaneously latch the visual and the audible indications.

- When the visual indications are latched, the visual indications, including alarm lamp, alarm message and its background remains when the alarm condition ends.
- When the audible indications are latched, the monitor issues alarm sounds when the alarm condition ends.

To latch a physiological alarm:

- 1. Select [Main Menu] \rightarrow [Maintenance >>] \rightarrow [User Maintenance >>] \rightarrow enter the required password.
- 2. Select [Alarm Setup >>]→[Latching Alarms>>].
- 3. Select how you want to latch the alarms.

Latching Alarms		×		
High 🗖 Latc	hing Visual Signal hing Visual Signal hing Visual Signal	 Latching Audible Signal Latching Audible Signal Latching Audible Signal 		
Low 🗖 Late	hing Visual Signal	Latching Audible Signal		
VTac Latching Lock	Enable	Defaults		
Enable/disable latching visual signal of Lethal level alarms.				

The rules for latching the alarms are:

- You can separately select [Latching Visual Signal].
- Selecting [Latching Audible Signal] simultaneously latches the visual signal.
- Selecting alarms of lower priority simultaneously latches the alarms of higher priority.
- The lethal alarms are latched by default. High, medium, and low priority alarms are unlatched by default.

NOTE

- Changing of alarm priority may affect the latching status of corresponding alarms. Please determine if you need to adjust the latching status for the specific alarm when you have changed its alarm priority.
- When the alarm system is reset, the latched physiological alarms are cleared.

You can separately configure the latching of the Vtac alarm. To do so, follow this procedure:

- 1. Select [Main Menu] \rightarrow [Maintenance >>] \rightarrow [User Maintenance >>] \rightarrow enter the required password.
- 2. Select [Alarm Setup >>]→[Latching Alarms>>].
- 3. Set [Vtac Latching Lock] to [Disable].
- 4. Select [Alarm Setup >>] quick key to enter [Alarm Setup] menu.
- 5. Select [Others] tab.
- 6. Set [**Vtac Latching**] as desired.

NOTE

• If the visual signal of lethal alarms is not latched, [Vtac Latching Lock] is not configurable and is automatically set to [Enable].

7.10 Testing Alarms

When the monitor starts up, a self-test is performed. In this case the alarm lamp is lit in yellow and red respectively, and the system gives a beep. This indicates that the visible and audible alarm indicators are functioning correctly.

For further testing of individual measurement alarms, perform the measurement on yourself (for example, SpO_2 or CO_2) or use a simulator. Then adjust alarm limits and check that appropriate alarm behavior is observed.

7.11 Using Care Group Alarms

7.11.1 Care Group Auto Alarms

When a Care Group is set up on your monitor, a flashing symbol will appear beside the QuickKeys area if any monitor in your Care Group, which is not currently viewed by your monitor, is alarming. The alarm symbol is shown as below.



When a monitor in the Care Group is disconnected, the flashing symbol is shown as below.



The department and bed label of the alarming monitor appear on the symbol. You can enter the View Other Patient window by pressing this symbol.

7.11.2 Setting Care Group Alert Tone

When a monitor in the Care Group issues an alarm, your monitor prompts you by sounding an alert tone. To set the alert tone:

- 1. In the [Main Menu], select [Screen Setup >>] \rightarrow [Screen Layout >>] \rightarrow [Choose Screen].
- 2. In the [Choose Screen] window, select [View Others Screen].
- 3. In the view other patient window, select [Setup>>], and set [Alert Tone] to [Repeat], [Once] or [Off].

7.11.3 Resetting Care Group Alarms

You can reset the alarm of the currently viewed bed in the View Other Patient window. This function can be set in the **[Alarm Setup]** menu from the **[User Maintenance]** menu only.

To enable this function:

- Select [Main Menu]→[Maintenance>>]→[User Maintenance>>]→enter the required password→[Alarm Setup>>].
- 2. In the [Alarm Setup] menu, set [Reset Other Bed's Alarms] to [On].

When your monitor is viewed by other monitors and [**Alarm Reset By Other Bed**] is switched on, alarms on your monitor can be reset by other monitors.

WARNING

• Resetting care group alarms may cause a potential hazard.

8.1 Introduction

The electrocardiogram (ECG) measures the electrical activity of the heart and displays it on the monitor as a waveform and a numeric. This section also discusses arrhythmia monitoring. The ECG algorithm enables 3-lead and 5-lead ECG monitoring and extended arrhythmia analysis.

8.2 Safety

WARNING

- Use only ECG electrodes and cables specified by the manufacturer.
- Make sure the conductive parts of electrodes and associated connectors for applied parts, including the neutral electrode, should not contact any other conductive parts including earth.
- Periodically inspect the electrode application site to ensure skin quality. If the skin quality changes, replace the electrodes or change the application site.
- Use defibrillation-proof ECG cables during defibrillation.
- Do not touch the patient, or table, or instruments during defibrillation.
- The neutral electrode of the HF surgical unit shall properly contact the patient. Otherwise, burns may result.
- The equipment is not intended for direct cardiac application.

CAUTION

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

NOTE

• After defibrillation, the screen display recovers within 10 seconds if the correct electrodes are used and applied in accordance with the manufacturer's instructions for use.

8.3 Preparing to Monitor ECG

8.3.1 Preparing the Patient and Placing the Electrodes

To prepare the patient and place the electrodes:

- 1. Prepare the patient's skin. Proper skin preparation is necessary for good signal quality at the electrode site, as the skin is a poor conductor of electricity. To properly prepare the skin, choose flat areas and then:
 - Shave hair from skin at chosen sites.
 - Gently rub skin surface at sites to remove dead skin cells.
 - Thoroughly cleanse the site with a mild soap and water solution. We do not recommend using ether or pure alcohol, because this dries the skin and increases the resistance.
 - Dry the skin completely before applying the electrodes.
- 2. Attach the clips or snaps to the electrodes before placing them.
- 3. Place the electrodes on the patient.
- 4. Attach the electrode cable to the patient cable and then plug the patient cable into the ECG connector.

8.3.2 Choosing AHA or IEC Lead Placement

To choose AHA or IEC lead placement:

- 1. Select the ECG parameter window or waveform area to enter the [ECG Setup] menu.
- 2. Select [Others]→[Lead Set] and then select [3-lead], [5-lead], or [Auto] according to the applied electrodes.
- 3. Select [Main Menu] \rightarrow [Maintenance >>] \rightarrow [User Maintenance >>] \rightarrow enter the required password.
- 4. Select [**Others** >>]→[**ECG Standard**] and then select [**AHA**] or [**IEC**] according to the hospital standard.

8.3.3 ECG Lead Placements

The electrode placement illustrations in this chapter adopt the AHA standard.

3-Leadwire Electrode Placement

Following is an electrode configuration when using 3 leadwires:

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

7

5-Leadwire Electrode Placement

Following is an electrode configuration when using 5 leadwires:

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

The chest (V) electrode can be placed on one of the following positions:

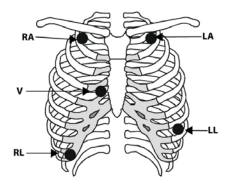
- V1 placement: on the fourth intercostal space at the right sternal border.
- V2 placement: on the fourth intercostal space at the left sternal border.
- V3 placement: midway between the V2 and V4 electrode positions.
- V4 placement: on the fifth intercostal space at the left midclavicular line.
- V5 placement: on the left anterior axillary line, horizontal with the V4 electrode position.
- V6 placement: on the left midaxillary line, horizontal with the V4 electrode position.
- V3R-V6R placement: on the right side of the chest in positions corresponding to those on the left.
- VE placement: over the xiphoid process.
- V7 placement: on posterior chest at the left posterior axillary line in the fifth intercostal space.
- V7R placement: on posterior chest at the right posterior axillary line in the fifth intercostal space.

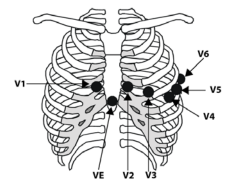
Lead Placement for Surgical Patients

The surgical site should be taken into consideration when placing electrodes on a surgical patient. e.g. for open-chest surgery, the chest electrodes can be placed on the lateral chest or back. To reduce artifacts and interference from electrosurgical units, you can place the limb electrodes close to the shoulders and lower abdomen and the chest electrodes on the left side of the mid-chest. Do not place the electrodes on the upper arm. Otherwise, the ECG waveform will be very small.

WARNING

- When using electrosurgical units (ESU), patient leads should be placed in a position that is equal distance from the Electrosurgery electrotome and the grounding plate to avoid burns to the patient. Never entangle the ESU cable and the ECG cable together.
- When using electrosurgical units (ESU), never place ECG electrodes near to the grounding plate of the ESU, as this can cause a lot of interference on the ECG signal.





8.3.4 Checking Paced Status

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

To change the paced status, you can select either:

- the patient information area, or
- [Main Menu]→[Patient Setup]→[Patient Demographics], or,
- the ECG parameter window or waveform area→[**Others** >>],

and then, select [Paced] from the pop-up menu and then select [Yes] or [No].

If you do not set the paced status, the monitor issues a prompt tone when pace pulse is detected. At the same time, the paced symbol flashes and the message [**Please confirm the pace of patient**] appears in the ECG waveform area. Then, please check and set the paced status of the patient.

WARNING

- For paced patients, you must set [Paced] to [Yes]. If it is incorrectly set to [No], the monitor could mistake a pace pulse for a QRS and fail to alarm when the ECG signal is too weak.
- On ventricular paced patients, episodes of Ventricular Tachycardia may not always be detected.
- False low heart rate indicators or false Asystole calls may result with certain pacemakers because of pacemaker artifact such as electrical overshoot of the pacemaker overlapping the true QRS complexes.
- Do not rely entirely upon the system's automated arrhythmia detection algorithm. Keep pacemaker patients under close surveillance.
- For non-paced patients, you must set [Paced] to [No].
- The auto pacer recognition function is not applicable to pediatric and neonatal patients.

8.4 Understanding the ECG Display

Your display may be configured to look slightly different.



- 1. Lead label of the displayed wave
- 2. ECG gain
- 3. ECG filter label
- 4. Notch filter status

Besides, when a paced signal has been detected, the pace pulse marks "|" are shown on the ECG wave if the [**Paced**] has been set to [**Yes**].



- 1. Current heart rate alarm limits
- 2. Current heart rate
- 3. Heart beat symbol

8.5 Changing ECG Settings

8.5.1 Accessing ECG Menus

By selecting the ECG parameter window or waveform area, you can access the [ECG Setup] menu.

8.5.2 Choosing the Alarm Source

In most cases the HR and PR numerics are identical. In order to avoid simultaneous alarms on HR and PR, the monitor uses either HR or PR as its active alarm source. To change the alarm source, select [**Alm Source**] in the [**ECG Setup**] menu and then select either:

- [HR]: if you want the HR to be the alarm source for HR/PR.
- [**PR**]: if you want the PR to be the alarm source for HR/PR.
- [Auto]: If the [Alm Source] is set to [Auto], the monitor will use the heart rate from the ECG measurements as the alarm source whenever a valid heart rate is available. If the heart rate becomes unavailable, for example the ECG module is turned off or becomes disconnected, the monitor will automatically switch to PR as the alarm source.

8.5.3 Changing ECG Wave Settings

In the [ECG Setup] menu:

- You can select [ECG], [ECG1], or [ECG2] to select a lead to view. The ECG lead (s) selected is used for analysis (beat detection, beat classification, and V-Fib detection). The waveform of selected lead should have the following characteristics:
 - The QRS should be either completely above or below the baseline and it should not be biphasic.
 - The QRS should be tall and narrow.
 - The P-waves and T-waves should be less than 0.2mV.
- If the wave is too small or clipped, you can change its size by selecting an appropriate [Gain] setting. If you select [Auto] from [Gain], the monitor will automatically adjust the size of the ECG waves. In normal screen, only the selected ECG wave's size is adjusted. In other screens, all ECG waves' size is adjusted simultaneously.
- You can change the wave sweep speed by selecting [**Sweep**] and then selecting the appropriate setting.

NOTE

• Changing the display gain on the monitor does not affect the amplitude of signal that is used by the algorithm for beat detection.

8.5.4 Changing the ECG Filter Settings

The ECG filter setting defines how ECG waves are smoothed. To change the filter setting, select [Filter] from [ECG Setup] and then select the appropriate setting.

- [Monitor]: Use under normal measurement conditions.
- [Diagnostic]: Use when diagnostic quality is required. The unfiltered ECG wave is displayed so that change such as R-wave notching is visible.
- [Surgery]: Use when the signal is distorted by high frequency or low frequency interference. High frequency interference usually results in large amplitude spikes making the ECG signal look irregular. Low frequency interference usually leads to wandering or rough baseline. In the operating room, the surgery filter reduces artifacts and interference from electrosurgical units. Under normal measurement conditions, selecting [Surgery] may suppress the QRS complexes too much and then interfere with ECG analysis.

CAUTION

• The [Diagnostic] filter is recommended when monitoring a patient in an environment with slight interference only.

8.5.5 Setting the ECG Lead Set

You can set the [Lead Set] by selecting [ECG Setup] \rightarrow [Others>>]. You can set the [Lead Set] as [Auto] if the auto lead detection function is available.

8.5.6 Choosing an ECG Display Screen

When monitoring with a 5-lead set, you can select the [Screens] Quickkey. In the [Choose Screen] window, choose the screen type as:

- [Normal Screen]: The ECG waveform area shows 2 ECG waveforms.
- [ECG 7-Lead Full-Screen]: The whole waveform area shows 7 ECG waveforms only.
- [ECG 7-Lead Half-Screen]: The upper half part of the whole waveform area displays 7 ECG waveforms.

When the screen type is set to [Normal Screen], cascaded ECG waveforms can be displayed. To cascade ECG waveforms:

- 1. Select the [Screens] Quickkey \rightarrow [Screen Setup].
- 2. Select [ECG1 Casc.] in the second row. A cascaded waveform is displayed in two waveform positions.

8.5.7 Setting the Notch Filter

The notch filter removes the line frequency interference. Only when [**Filter**] is set to [**Diagnostic**], the [**Notch Filter**] is adjustable.

To set the notch filter:

- 1. Select the ECG parameter window or waveform area to enter its setup menu. Then select [Others >>].
- 2. Set [Notch Filter] to:
 - [Strong]: when waveform interference is strong (such as spikes).
 - [Weak]: when waveform interference is weak.
 - [**Off**]: to turn the notch filter off.

Set notch frequency according to the electric power frequency of your country. To set notch filter frequency:

- 1. When [Notch Filter] is turned on, select [Main Menu]→[Maintenance >>]→[User Maintenance >>]→enter the required password.
- 2. Select [Others >>]→[Notch Freq.] and then select [50Hz] or [60Hz] according to the power line frequency.

8.5.8 Changing the Pacer Reject Settings

Select [ECG Setup] \rightarrow [Others>>] \rightarrow [Pacer Reject], and then select [On] or [Off]. When [Paced] is set to [Yes]:

- When [**Pacer Reject**] is switched on, the pace pulses are not counted as extra QRS complexes.
- The pace pulse marks "|" are shown on the ECG wave when pace pulses are detected.

When [**Paced**] is set to [**No**], the pace markers are not shown on the ECG wave, and the options of [**Pacer Reject**] are inactivated.

8.5.9 Enabling Smart Lead Off

When the smart lead off function is set on and there is a "lead off" in the lead that has an ECG waveform in filter mode and notch status, if another lead is available, this available lead automatically becomes that lead. The system will re-calculate HR and analyze and detect arrhythmia. When the "lead off" condition is corrected, the leads are automatically switched back.

To switch on/off the smart lead off function, select [**Others** >>] from the [**ECG Setup**] menu; select [**Smart Lead Off**] and then select [**On**] or [**Off**] from the pop-up menu.

8.5.10 Setting the Alarm Level for ECG Lead Off Alarms

Select [Alarm Setup >>] from the [User Maintenance] menu. You can set [ECGLeadOff Lev.] from the pop-up menu.

8.5.11 Adjusting QRS Volume

QRS sounds are produced based on the alarm source.

To adjust the QRS volume, select [**Others** >>] from the [**ECG Setup**] menu; select [**QRS Volume**] from the pop-up menu and select the appropriate setting. When a valid SpO_2 measured value is available, the system adjusts the pitch tone of QRS sound based on the SpO_2 value.

8.5.12 Adjusting the Minimum QRS Detection Threshold

To avoid false asystole alarms when the R wave amplitude is low and missed asystole alarms during ventricular standstill (tall P waves, but no QRS), a means to manually adjust the minimum QRS detection threshold is provided.

To adjust the QRS threshold,

- 1. In the [ECG Setup] menu, set [Filter] to [Monitor].
- 2. Select [Others >>]→[Minimum QRS Threshold >>] to enter the [Minimum QRS Threshold] menu.
- 3. Select the up or down arrow to adjust the QRS threshold. Selecting [**Default**] resets the QRS threshold to the default value (0.16 mV).
- 4. Select [**Confirm**] to make the changes effective.

CAUTION

- The setting of QRS threshold can affect the sensitivity of arrhythmia, QT/QTc detection, and heart rate calculation.
- If QRS amplitude is low, the monitor might not be able to calculate heart rate and false asystole may occur.

NOTE

• The minimum QRS detection threshold can only be adjusted when the ECG filter is set to [Monitor].

8.5.13 About the Defibrillator Synchronization

A defibrillator synchronization pulse (100 ms, +5V) is output through the Multifunctional connector every time the monitor detects an R-wave.

To use the defibrillator synchronization function, connect the monitor and the defibrillator with a synchronization cable.

WARNING

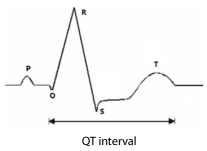
• Improper use of a defibrillator may cause injury to the patient. The user should determine whether to perform defibrillation or not according to the patient's condition.

8.6 QT/QTc Interval Monitoring

The QT interval is defined as the time between the beginning of the Q-wave and the end of the T-wave. It measures the total duration of the depolarization (QRS duration) and repolarization (ST-T) phases of the ventricles. QT interval monitoring can assist in the detection of long QT syndrome.

The QT interval has an inverse relationship to heart rate. As heart rate increases, the QT interval shortens, while at lower heart rates QT interval gets longer. Several formulas are available to correct QT interval for heart rate. The heart rate corrected QT interval is abbreviated as QTc.

QT/QTc Interval Monitoring is intended for adult, pediatric, and neonate patients.



CAUTION

• For QT interval monitoring to be effective, basic or enhanced arrhythmia monitoring should be turned on.

8.6.1 QT/QTc Monitoring Limitations

Some conditions may make it difficult to achieve reliable QT monitoring, for example:

- R-wave amplitudes are too low
- The presence of frequent ventricular ectopic beats
- Unstable RR intervals
- P-waves tending to encroach on the end of the Adrevious T-wave at high heart rates
- T-waves are very flat or not well defined
- The end of the T-wave is difficult to delineate because of the presence of U-waves
- QTc measurements are not stable
- In the presence of noise, asystole, ventricular fibrillation, and ECG lead off

For these cases you should select a lead with good T-wave amplitude and no visible flutter activity, and without a predominant U-wave or P-wave.

Some conditions such as left or right bundle branch block or hypertrophy can lead to a widened QRS complex. If a long QTc is observed you should verify it to ensure that it is not caused by QRS widening.

Because normal beats followed by ventricular beats are not included in the analysis, no QT measurement will be generated in the presence of a bigeminy rhythm.

If the heart rate is extremely high (over 150bpm for adults and over 180bpm for pediatrics and neonates), QT will not be measured. When the heart rate changes, it can take several minutes for the QT interval to stabilize. For reliable QTc calculation it is important to avoid the region where the heart rate is changing.

WARNING

• QT/QTc measurements should always be verified by a qualified clinician.

8.6.2 Enabling QT/QTc Monitoring

The QT monitoring function is disabled by default. Before you start QT monitoring, enable the QT function.

To enable QT/QTc monitoring:

- 1. In the [ECG Setup] menu, select [QT Analysis>>] to enter the [QT Analysis] menu.
- 2. Set [QT Analysis] to [On].

8.6.3 Displaying QT/QTc Parameters and Waveform

To display QT numerics and waveform:

- Select the [Screens] QuickKey or select [Main Menu]→[Screen Setup >>]→[Screen Layout >> to enter the [Screens] window.
- 2. Select the [Screen Setup] tab.
- 3. Select the parameter area where you want to display the QT parameters and select [QT].

The following picture shows the QT numeric area. Your monitor screen may look slightly different:



- 1: Parameter label
- 2: QTc alarm limit (if QTc alarm is off, the alarm off symbol is displayed)
- 3: QTc value
- 4: ΔQTc value (the difference between the current and reference QTc values. If ΔQTc alarm is off, the alarm off symbol is displayed on the right.)
- 5: QT value
- 6: QT-HR value

NOTE

• QTc values are calculated based on the QT-HR, not the ECG HR.

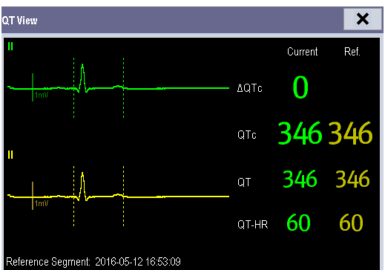
8.6.4 Accessing the QT View

QT View shows the current and reference QT parameter values and waveforms.

To accss the QT View:

- 1. Select the QT parameter area or waveform area to enter the [**QT Analysis**] menu.
- 2. Select [**QT View>>**].

The following picture shows the QT view.



- The current waveform is shown in the upper half in green.
- The reference waveform is shown below in yellow.
- The start of QRS complex and the end of the T wave are marked with vertical lines.
- In some conditions, no QT measurement can be calculated. Then the cause of failed QT measurement is shown at the bottom of the QT numerics area. Additionally the message "Cannot Analyze QT" is shown in the technical alarm area.

In the QT View, you can:

- Select the arrows besides [**View Leads**] to switch the displayed lead.
- Select [**Save Ref.**] to save the current QT parameters and waveforms as reference. If no reference has been set for this patient within the first five minutes after getting valid QT values, the monitor will automatically set a reference. If you set a new reference, the previous reference will be discarded.

CAUTION

• Updating QTc reference affects ΔQTc value and alarm.

8.6.5 Changing QT Settings 8.6.5.1 Setting QT Alarm Properties

To set QT alarm properties,

- 1. Select the [Alarm Setup] QuickKey, or select [Alarm Setup>>] from the [QT Analysis] menu.
- 2. Set QTc and Δ QTc alarm properties.

8.6.5.2 Selecting the QT Analysis Lead

You can select one lead or all leads for QT calculation. To select the QT analysis lead:

- 1. Select the QT parameter area or waveform area to enter the [QT Analysis] menu.
- 2. Set [Analysis Lead]. [All] is selected by default. This means all leads are used for QT calculation.

8.6.5.3 Changing QTc Formula

The monitor uses as a default the Hodges correction formula to correct the QT interval for heart rate. To change the QTc formula:

- 1. Select the QT parameter area or waveform area to enter the [QT Analysis] menu.
- 2. Set [QTc Formula].
 - Hodges: $QTc = QT + 1.75 \times (HeartRate 60)$

Bazett:
$$QTc = QT \times \left(\frac{HeartRate}{60}\right)^{\frac{1}{2}}$$

$$QTc = QT \times \left(\frac{HeartRate}{60}\right)^{\frac{1}{3}}$$

$$QTc = QT + 154 \times \left(1 - \frac{60}{HeartRate}\right)$$

Framingham:

8.7 About Arrhythmia Monitoring

Arrhythmia analysis provides information about the patient's cardiac parameters like heart rate, PVC rate, and rhythm.

WARNING

- Arrhythmia analysis program is intended to detect ventricular arrhythmias and atrial fibrillation. It is not designed to detect all the atrial or supraventricular arrhythmias. It may incorrectly identify the presence or absence of an arrhythmia. Therefore, a physician must analyze the arrhythmia information with other clinical findings.
- The arrhythmia detection is not intended for neonatal patients.
- Heart-rate reading may be affected by cardiac arrhythmias. Do not rely entirely on rate meter alarms when monitoring patients with arrhythmias. Always keep these patients under close surveillance.
- Atrial fibrillation (Afib) detection function is not intended for pediatric and neonatal patients.

8.7.1 Understanding the Arrhythmia Events

Arrhythmia message	Description	Category
Acustolo	No QRS detected within the set time threshold in absence of ventricular	
Asystole	fibrillation or chaotic signal.	
Vfib/Vtac	A fibrillatory wave for 6 consecutive seconds.	
VIID/VTaC	A dominant rhythm of adjacent Vs and a HR > the V-Tac HR limit.	Lethal arrhythmia
Vtac	The consecutive PVCs \geq Vtac PVCs limit, and the HR \geq the Vtac rate limit.	
Vant Brady	The consecutive PVCs \geq the Vbrd threshold and the ventricular HR < the Vbrd	
Vent. Brady	Rate threshold.	
Extreme Tachy	The heart rate is equal to or greater than the extreme tachycardia limit.	
Extreme Brady	The heart rate is equal to or less than the extreme bradycardia limit.	
PVCs	PVCs/min exceeds high limit	
Da sor Not Da sin s	No pace pulse detected for 1.75 x average R-to-R intervals following a QRS	
Pacer Not Pacing	complex (for paced patients only).	
De eeu Net Construis	No QRS complex detected for 300 milliseconds following a pace pulse (for	
Pacer Not Capture	paced patients only).	
PVC	C One PVC detected in normal heartbeats.	
Couplet	plet Paired PVCs detected in normal heartbeats.	
Run PVCs	More than 2 consecutive PVCs within the last minute.	
Bigeminy	eminy A dominant rhythm of N, V, N, V, N, V.	
Trigeminy		
R on T		
	No beat detected for 1.75 x average R-R interval for HR <120, or No beat for 1	Nonlethal
Missed Beats	second with HR > 120 (for non-paced patients only), or No beat detected for	arrhythmia
	more than the set pause threshold.	
Brady	The average heart rate is equal to or less than the bradycardia limit.	
Tachy	The average heart rate is equal to or greater than the tachycardia limit.	
	The consecutive PVCs \geq the Vbrd PVCs limit, and the HR \geq Vbrd Rate limit but <	
Vent. Rhythm	the Vtac Rate limit.	
Multif. PVC	if. PVC Multiform PVCs detected in Multif. PVC's Window (which is adjustable).	
	The consecutive PVCs < the Vtac PVCs limit but > 2, and $HR \ge$ the Vtac Rate	1
Nonsus. Vtac	limit.	
Pause	No QRS detected within the set time threshold of pause.	
Irr. Rhythm	Consistently irregular rhythm.]
Afib	Absence of P-waves and/or variable RR intervals between normal beats.	

8.7.2 Changing Arrhythmia Alarm Settings

To change arrhythmia alarm settings, select the ECG parameter area or waveform area \rightarrow [ECG Setup] \rightarrow [Arrh. Analysis >>]. In the pop-up menu, you can set the [Alm Lev] to [High], [Med], [Low] or [Message], or switch on lethal arrhythmia analysis alarms only or switch on/off all arrhythmia analysis alarms. In the [Alarm Setup] menu from the [User Maintenance] menu, you can enable/disable turning off lethal arrhythmia analysis alarms.

WARNING

• If you switch off all arrhythmia analysis alarms, the monitor cannot give any arrhythmia analysis alarm. Always keep the patient under close surveillance.

NOTE

• The priority of lethal arrhythmia alarms is always high. It is unchangeable.

8.7.3 Changing Arrhythmia Threshold Settings

Select the ECG parameter window or waveform area \rightarrow [**Arrh. Analysis** >>] \rightarrow [**Arrh. Threshold**], and you can then change threshold settings for some arrhythmia alarms. When an arrhythmia violates its threshold, an alarm will be triggered. The asystole delay time relates to ECG relearning. When HR is less than 30 bpm, it is recommended to set the asystole delay time to 10 seconds.

Arrh. event	Range	Default	Step	Unit
PVCs High	1 to 100	10	1	/min
Asys. Delay	3 to 10	5	1	s
Tachy High		Adult: 120	5	bpm
Tachy High	60 to 300	Pediatric: 160		
Productions	15 to 120	Adult: 50	E	hom
Brady Low	15 10 120	Pediatric: 75	5	bpm
Extromo Tachy	120.4- 200	Adult: 160	5	bpm
Extreme Tachy	120 to 300	Pediatric: 180		
Extreme Brady	15 to 60	Adult: 35	5	haas
Extreme brady	13 10 00	Pediatric: 50	5	bpm
Multif. PVC's Window	3 to 31	15	1	/min
Vtac Rate	100 to 200	Adult, pediatric: 130	5	bpm
Vtac PVCs	3 to 99	6	1	/min
Pause Time	1.5, 2.0,2.5	2	/	S
Vbrd PVCs	3 to 99	5	1	/min
Vbrd Rate	15 to 60	40	5	bpm

8.7.4 Setting the Extended Arrhythmia

The following arrhythmia events are defined as extended arrhythmia:

- Extreme Tachy
- Extreme Brady
- Vent. Brady
- Nonsus. Vtac
- Multif. PVC
- Irr. Rhythm
- Pause
- Afib

You can select [Main Menu] \rightarrow [Maintenance >>] \rightarrow [User Maintenance >>] \rightarrow enter the required password \rightarrow select [Alarm Setup >>], and set [Extended Arrh.] to [Enable] or [Disable]. When [Extended Arrh.] is set to [Disable], the monitor does not analyze the extended arrhythmia events, and corresponding alarms are not given.

8.7.5 Reviewing Arrhythmia Events

Please refer to section 19.4 Reviewing Events.

8.8 ECG Relearning

8.8.1 Initiating an ECG Relearning Manually

During ECG monitoring, you may need to initiate an ECG relearning when the patient's ECG template changes dramatically. A change in the ECG template could result in:

- Incorrect arrhythmia alarms, and/or
- Inaccurate heart rate

ECG relearning allows the monitor to learn the new ECG template to correct arrhythmia alarms and HR value. To initiate relearning manually, select the ECG parameter window or waveform area \rightarrow [**Relearn**]. When the monitor is learning, the message [**ECG Learning**] is displayed in the technical alarm area.

WARNING

• Initiate ECG relearning only during periods of normal rhythm and when the ECG signal is relatively noise-free. If ECG learning takes place during ventricular rhythm, the ectopics may be incorrectly learned as the normal QRS complex. This may result in missed detection of subsequent events of V-Tach and V-Fib.

8.8.2 Automatic ECG Relearning

ECG relearning is initiated automatically whenever:

- The ECG lead or lead label is changed.
- The ECG lead is re-connected.
- A new patient is admitted.
- After the ECG calibration is completed, and [Stop Calibrating ECG] is selected.
- A switch happens between the options of screen type during 5-lead ECG monitoring.
- The paced status of the patient is changed.

8.9 Troubleshooting

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

CAUTION

• Never try to disassemble the equipment or supplied accessories. There are no internal user-serviceable parts.

Symptoms	Possible Cause	Correction Action
Noisy ECG traces	Loose or dry electrodes	Apply fresh and moist electrodes.
1. Allen, Hust	Defective electrode wires	Replace wires if necessary.
Minhanne	Patient cable or leads are routed too	Move the patient cable or leads away from the
•	close to other electrical devices	electrical device.
Excessive Electro-surgical	Wrong ECG cable used	Use ESU-proof ECG cables. For details, refer to
Interference		section 27.1 ECG Accessories.
Muscle Noise	Inadequate skin preparation prior to	Repeat skin preparation as described in section
	application of electrode, tremors,	8.3.1 Preparing the Patient and Placing the
	tense subject, and/or poor electrode	<i>Electrodes</i> and re-place the electrodes.
فسأن بنساك بلساك يقسأ وخساك واساك باساك ش	placement	Apply fresh, moist electrodes.
arah, anah, anah, masih anah, anah, anah, ana		Avoid muscular areas.
Intermittent Signal	Connections not tight and/or properly	Check that the cables are properly connected.
	secured	
	Electrodes dry or loose	Repeat skin preparation as described in section
		8.3.1 Preparing the Patient and Placing the
		<i>Electrodes</i> and apply fresh and moist electrodes.
	Cable or lead wires damaged	Change cable and lead wires.
Excessive alarms: heart rate,	Electrodes dry	Repeat skin preparation as described in section
lead fault		8.3.1 Preparing the Patient and Placing the
		<i>Electrodes</i> and apply fresh, moist electrodes.
	Excessive patient movement or	Reposition the electrodes.
	muscle tremor	Replace fresh and moist electrodes if necessary.

Symptoms	Possible Cause	Correction Action
Low Amplitude ECG Signal	Gain set too low	Set the gain as required. For details, refer to section
		8.5.3 Changing ECG Wave Settings.
	Electrodes dry / old	Apply fresh and moist electrodes.
	Skin improperly prepared	Repeat skin preparation as described in section
		8.3.1 Preparing the Patient and Placing the
		Electrodes.
	This could be the patient's normal QRS	Verify with another well-functioning monitor.
	complex	
	Electrode could be positioned over a	Move ECG patches away from the bone or muscle
	bone or muscle mass	mass.
No ECG Waveform	Gain set too low	Set the gain as required. For details, refer to section
		8.5.3 Changing ECG Wave Settings.
	Lead wires and patient cable not fully	Check that the leadwires and patient cables are
	or properly inserted	properly connected.
	Cable or lead wires damaged	Change cable and lead wires.
Base Line Wander	Patient moving excessively	Secure leadwires and cable to patient.
, I	Electrodes dry or loose	Repeat skin preparation as described in section
which land		8.3.1 Preparing the Patient and Placing the
A lama		<i>Electrodes</i> and apply fresh and moist electrodes.
- lon	ECG Filter set to Diagnostic mode	Set ECG Filter to "Monitor" mode.

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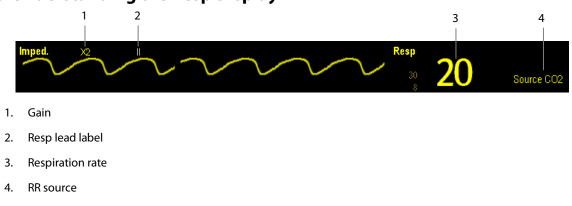
9.1 Introduction

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

9.2 Safety Information

WARNING

- When monitoring the patient's respiration, do not use ESU-proof ECG cables.
- If you do not set the detection level for the respiration correctly in manual detection mode, it may not be possible for the monitor to detect apnea. If you set the detection level too low, the monitor is more likely to detect cardiac activity, and to falsely interpret cardiac activity as respiratory activity in the case of apnea.
- The respiration measurement does not recognize the cause of apneas. It only indicates an alarm if no breath is detected when a pre-configured time has elapsed since the last detected breath. Therefore, it cannot be used for diagnostic purpose.
- If operating under conditions according to the EMC Standard EN 60601-1-2 (Radiated Immunity 3V/m), field strengths above 1V/m may cause erroneous measurements at various frequencies. Therefore it is recommended to avoid the use of electrically radiating equipment in close proximity to the respiration measurement unit.



9.3 Understanding the Resp Display

By selecting the waveform area or parameter area, you can enter the [Resp Setup] menu.

NOTE

• Respiration monitoring is not for use on the patients who are very active, as this will cause false alarms.

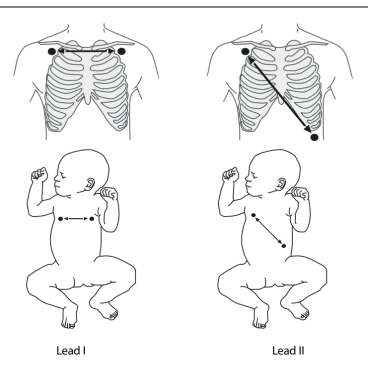
9.4 Placing Resp Electrodes

As the skin is a poor conductor of electricity, preparing the skin is necessary for a good respiration signal. You can refer to the ECG section for how to prepare the skin. For details, refer to section **8.3.1** *Preparing the Patient and Placing the Electrodes*.

As the respiration measurement adopts the standard ECG electrode placement, you can use different ECG cables (3-lead, or 5-lead). Since the respiration signal is measured between two ECG electrodes, if a standard ECG electrode placement is applied, the two electrodes should be RA and LA of ECG Lead I, or RA and LL of ECG Lead II.

NOTE

• To optimize the respiration waveform, place the RA and LA electrodes horizontally when monitoring respiration with ECG Lead I; place the RA and LL electrodes diagonally when monitoring respiration with ECG Lead II.



9.4.1 Optimizing Lead Placement for Resp

If you want to measure Resp and you are already measuring ECG, you may need to optimize the placement of the two electrodes between which Resp will be measured. Repositioning ECG electrodes from standard positions results in changes in the ECG waveform and may influence arrhythmia interpretation.

9.4.2 Cardiac Overlay

Cardiac activity that affects the Resp waveform is called cardiac overlay. It happens when the Resp electrodes pick up impedance changes caused by the rhythmic blood flow. Correct electrodes placement can help to reduce cardiac overlay: avoid the liver area and the ventricles of the heart in the line between the respiratory electrodes. This is particularly important for neonates.

9.4.3 Abdominal Breathing

Some patients with restricted movement breathe mainly abdominally. In these cases, you may need to place the left leg electrode on the left abdomen at the point of maximum abdominal expansion to optimise the respiratory wave.

9.4.4 Lateral Chest Expansion

In clinical applications, some patients (especially neonates) expand their chests laterally, causing a negative intrathoracic pressure. In these cases, it is better to place the two respiration electrodes in the right midaxillary and the left lateral chest areas at the patient's maximum point of the breathing movement to optimise the respiratory waveform.

9.5 Choosing the Respiration Lead

In the [Resp Setup] menu, set [Resp Lead] to [I], [II] or [Auto].

9.6 Changing the Apnea Alarm Delay

The apnea alarm is a high-level alarm used to detect apneas. You can set the apnea alarm delay time after which the monitor alarms if the patient stops breathing.

In the [**Resp Setup**] menu, select [**Apnea Delay**] and then select the appropriate setting. The [**Apnea Delay**] of Resp, CO₂, and AG module keeps consistent with each other.

9.7 Changing Resp Detection Mode

In the [Resp Setup] menu, select [Detection Mode] and then select [Auto] or [Manual].

In auto detection mode, the monitor adjusts the detection level automatically, depending on the wave height and the presence of cardiac artifact. Note that in auto detection mode, the detection level (a dotted line) is not displayed on the waveform.

Use auto detection mode for situations where:

- The respiration rate is not close to the heart rate.
- Breathing is spontaneous, with or without continuous positive airway pressure (CPAP).
- Patients are ventilated, except patients with intermittent mandatory ventilation (IMV).
- In manual detection mode, you adjust the dotted detection level line to the desired level by selecting [Upper Line] or [Lower Line] and then selecting or selecting or selection level will not adapt automatically to different respiration depths. It is important to remember that if the depth of breathing changes, you may need to change the detection level.

Use manual detection mode for situations where:

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

In Auto Detection Mode, if you are monitoring Resp and ECG is switched off, the monitor cannot compare the ECG and Resp rates to detect cardiac overlay. The respiration detection level is automatically set higher to prevent the detection of cardiac overlay as respiration.

In Manual Detection Mode, cardiac overlay can in certain situations trigger the respiration counter. This may lead to a false indication of a high respiration or an undetected apnea condition. If you suspect that cardiac overlay is being registered as breathing activity, raise the detection level above the zone of cardiac overlay. If the Resp wave is so small that raising the detection level is not possible, you may need to optimize the electrode placement as described in section **9.4.4 Lateral Chest Expansion**.

9.8 Changing Resp Wave Settings

WARNING

• When monitoring in manual detection mode, make sure to check the respiration detection level after you have increased or decreased the size of the respiration wave.

In the [**Resp Setup**] menu, you can:

- Select [Gain] and then select an appropriate setting. The bigger the gain is, the larger the wave amplitude is.
- Select [Sweep] and then select an appropriate setting. The faster the wave sweeps, the wider the wave is.

9.9 Setting Respiration Rate Source

To set RR source:

- 1. Enter the [Resp Setup] menu.
- 2. Select [RR Source] and then select a source or [Auto] from the dropdown list.

The dropdown list displays the currently available RR source. When you select [**Auto**], the system will automatically select the RR source according to the priority. When the current RR source does not have valid measurement, the system will automatically switch the [**RR Source**] to [**Auto**]. RR source switches back to impedance respiration if you press the [**Alarm Reset**] key during an apnea alarm.

The priority of RR source is (from high to low): CO_2 measurement, and impedance respiration measurement. The [**RR Source**] settings of Resp, CO_2 , and AG module are linked.

The RR source options and description are shown in the table below.

Option	Description	
Auto	RR source is automatically selected according to the priority.	
CO ₂	RR source is from CO_2 measurement.	
ECG	RR source is from impedance respiration measurement.	

9.10 Setting Alarm Properties

Select [**Alarm Setup** >>] from the [**Resp Setup**] menu. In the pop-up menu, you can set alarm properties for this parameter.

FOR YOUR NOTES

10.1 Introduction

The pulse numeric counts the arterial pulsations that result from the mechanical activity of the heart. You can display a pulse from any measured SpO₂ or any arterial pressure (see chapter **14 Monitoring IBP**). The displayed pulse numeric is color-coded to match its source.



- 1. PR: detected beats per minute.
- 2. PR Source

NOTE

• A functional tester or SpO₂ simulator can be used to determine pulse rate accuracy.

10.2 Setting the PR Source

The current pulse source is displayed in the PR parameter area. The pulse rate chosen as pulse source:

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

To set which pulse rate as PR source:

- 1. Enter the [**SpO2 Setup**] menu.
- 2. Select [**PR Source**] and then select a label or [**Auto**] from the pop-up menu.

The pop-up menu displays the currently available PR sources from top to bottom by priority. When you select [**Auto**], the system will automatically select the first option as the PR source from the pop-up menu. When the current PR source is unavailable, the system will automatically switch [**PR Source**] to [**Auto**]. When you select [**IBP**], the system will automatically select the first pressure label as the PR source from the pop-up menu.

10.3 Selecting the Active Alarm Source

In most cases the HR and pulse numerics are identical. In order to avoid simultaneous alarms on HR and PR, the monitor uses either HR or PR as its active alarm source. To change the alarm source, select [**Alm Source**] in the [**ECG Setup**] or [**SpO2 Setup**] menu and then select either:

- [**HR**]: The monitor will use the HR as the alarm source for HR/pulse.
- [**PR**]: The monitor will use the PR as the alarm source for HR/pulse.
- [Auto]: If the [Alm Source] is set to [Auto], the monitor will use the heart rate from the ECG measurement as the alarm source whenever the ECG measurement is switched on and a valid heart rate is available. If the heart rate becomes unavailable, for example if leads becomes disconnected, and a pulse source is switch on and available, the monitor will automatically switch to Pulse as the alarm source. When the Leads Off condition is corrected, the monitor will automatically switch back to the heart rate as the alarm source.

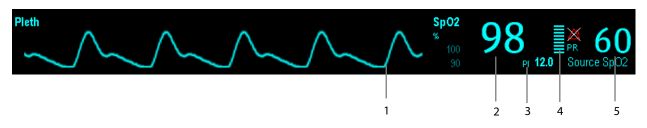
10.4 QRS Tone

When PR is used as the alarm source, the PR source will be used as a source for the QRS tone. You can change the QRS volume by adjusting [**Beat Vol**] in the [**SpO2 Setup**] menu. When a valid SpO₂ value exists, the system will adjust the pitch tone of QRS volume according to the SpO₂ value.

11.1 Introduction

 SpO_2 monitoring is a non-invasive technique used to measure the amount of oxygenated haemoglobin and pulse rate by measuring the absorption of selected wavelengths of light. The light generated in the probe passes through the tissue and is converted into electrical signals by the photodetector in the probe. The SpO_2 module processes the electrical signal and displays a waveform and digital values for SpO_2 and pulse rate.

This device is calibrated to display functional oxygen saturation. It provides the following measurements:



- 1. Pleth waveform (Pleth): visual indication of patient's pulse. The waveform is not normalized.
- 2. Oxygen saturation of arterial blood (SpO₂): percentage of oxygenated hemoglobin in relation to the sum of oxyhemoglobin and deoxyhemoglobin.
- 3. Perfusion index (PI): gives the numerical value for the pulsatile portion of the measured signal caused by arterial pulsation. PI is an indicator of the pulsatile strength. You can also use it to assess the quality of SpO₂ measurement.
 - Above 1 is optimal
 - Between 0.3 and 1 is acceptable
 - Below 0.3 indicates low perfusion;. When PI is below 0.3, a question mark (?) is displayed to the right of the SpO₂ value, indicating that the SpO₂ value may be inaccurate. R eposition the SpO₂ sensor or find a better site. If low perfusion persists, choose another method to measure oxygen saturation if possible.

PI is available for Masimo SpO₂ module. For Masimo SpO₂ module, PI value can be displayed under the PR value in larger characters if [**PI Zoom**] is enabled.

- 4. Perfusion indicator: the pulsatile portion of the measured signal caused by arterial pulsation.
- 5. Pulse rate (derived from the pleth wave): detected pulsations per minute.

NOTE

• A functional tester or SpO₂ simulator cannot be used to assess the accuracy of a SpO₂ module or a SpO₂ sensor.

11.2 Safety

WARNING

- Use only SpO₂ sensors specified in this manual. Follow the SpO₂ sensor's instructions for use and adhere to all warnings and cautions.
- When a trend toward patient deoxygenation is indicated, blood samples should be analyzed by a laboratory co-oximeter to completely understand the patient's condition.
- The equipment is not intended for use in an MRI environment.
- Prolonged continuous monitoring may increase the risk of undesirable changes in skin characteristics, such as irritation, reddening, blistering or burns. Inspect the sensor site every two hours and move the sensor if the skin quality changes. Change the application site every four hours. For neonates, or patients with poor peripheral blood circulation or sensitive skin, inspect the sensor site more frequently.

11.3 Identifying SpO₂ Modules

To identify which SpO_2 module is incorporated into your monitor, see the color of the SpO_2 connector and the company logo located at the monitor. The color of the cable connector matches the company as shown below:

- Masimo SpO₂ module: a purple connector with a logo of Masimo SET.
- Nellcor SpO_2 module: a grey connector with a logo of Nellcor.

The connectors for these three SpO₂ sensors are mutually exclusive.

11.4 Applying the Sensor

To apply the sensor:

- 1. Select an appropriate sensor according to the module type, patient category and weight.
- 2. Remove colored nail polish from the application site.
- 3. Apply the sensor to the patient.
- 4. Select an appropriate adapter cable according to the connector type and plug this cable into the SpO₂ connector.
- 5. Connect the sensor cable to the adapter cable.

WARNING

• If the sensor is too tight because the application site is too large or becomes too large due to edema, excessive pressure for prolonged periods may result in venous congestion distal from the application site, leading to interstitial edema and tissue ischemia.

11.5 Changing SpO₂ Settings

11.5.1 Accessing SpO₂ Menus

By selecting the SpO₂ parameter window or waveform area, you can access the [SpO2 Setup] menu.

11.5.2 Adjusting the Desat Alarm

The desat alarm is a high level alarm notifying you of potentially life threatening drops in oxygen saturation. Select [**Alarm Setup** >>] from the [**SpO2 Setup**] menu. From the pop-up menu, you can set low alarm limit, alarm switch, and alarm recording for [**Desat**]. When the SpO₂ value is below the desat alarm limit and desat alarm switch is set on, the message [**SpO2 Desat**] is displayed.

11.5.3 Setting SpO₂ Sensitivity

For Masimo SpO₂ module, you can set [Sensitivity] to [Normal] or [Maximum] in the [SpO2 Setup] menu. When the [Sensitivity] is set to [Maximum], the monitor is more sensitive to minor signals. When monitoring critically ill patients whose pulsations are very weak, it is strongly recommended that the sensitivity is set to [Maximum]. When monitoring neonatal or non-critically ill patients who tend to move a lot, noise or invalid signals may be caused. In this case, it is recommended that the sensitivity is set to [Normal] so that the interference caused by motion can be filtered and therefore the measurement stability can be ensured.

11.5.4 Changing Averaging Time

The SpO₂ value displayed on the monitor screen is the average of data collected within a specific time. The shorter the averaging time is, the quicker the monitor responds to changes in the patient's oxygen saturation level. Contrarily, the longer the averaging time is, the slower the monitor responds to changes in the patient's oxygen saturation level, but the measurement accuracy will be improved. For critically ill patients, selecting shorter averaging time will help understanding the patient's state.

To set the averaging time for Masimo SpO₂ module:

- ,1. Select [Averaging] in the [SpO2 Setup] menu.
- 2. Then select [2-4 s], [4-6 s], [8 s], [10 s], [12 s], [14 s] or [16 s].

11.5.5 Monitoring SpO₂ and NIBP Simultaneously

When monitoring SpO₂ and NIBP on the same limb simultaneously, you can switch [**NIBP Simul**] on in the [**SpO2 Setup**] menu to lock the SpO₂ alarm status until the NIBP measurement ends. If you switch [**NIBP Simul**] off, low perfusion caused by NIBP measurement may lead to inaccurate SpO₂ readings and therefore cause false physiological alarms.

11.5.6 Sat-Seconds Alarm Management

With traditional alarm management, high and low alarm limits are set for monitoring oxygen saturation. During monitoring, as soon as an alarm limit is violated, an audible alarm immediately sounds. When the patient % SpO₂ fluctuates near an alarm limit, the alarm sounds each time the limit is violated. Such frequent alarms can be distracting. Nellcor's Sat-Seconds alarm management technique is used to reduce these nuisance alarms.

The Sat-Seconds feature is available with the Nellcor SpO₂ module to decrease the likelihood of false alarms caused by motion artifacts. To set the Sat-Seconds limit, select [**Sat-Seconds**] in the [**SpO2 Setup**] menu and then select the appropriate setting.

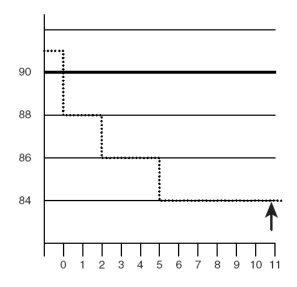
With Sat-Seconds alarm management, high and low alarm limits are set in the same way as traditional alarm management. A Sat-Seconds limit is also set. The Sat-Seconds limit controls the amount of time that SpO₂ saturation may be outside the set limits before an alarm sounds. The method of calculation is as follows: the number of percentage points that the SpO₂ saturation falls outside the alarm limit is multiplied by the number of seconds that it remains outside the limit. This can be stated as the equation:

Sat-Seconds= Points × Seconds

Only when the Sat-Seconds limit is reached, the monitor gives a Sat-Seconds alarm. For example, the figure below demonstrates the alarm response time with a Sat-Seconds limit set at 50 and a low SpO_2 limit set at 90%. In this example, the patient % SpO_2 drops to 88% (2 points) and remains there for 2 seconds. Then it drops to 86% (4 points) for 3 seconds, and then to 84% (6 points) for 6 seconds. The resulting Sat-Seconds are:

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

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



Saturation levels may fluctuate rather than remaining steady for a period of several seconds. Often, the patient % SpO₂ may fluctuate above and below an alarm limit, re-entering the non-alarm range several times. During such fluctuation, the monitor integrates the number of %SpO₂ points, both positive and negative, until either the Sat-Seconds limit is reached, or the patient %SpO₂ re-enters the non-alarm range and remains there.

NOTE

• The "SpO2 Too Low" or "SpO2 Too High" alarm is presented in the case that SpO₂ value violates the alarm limits for 3 times within one minute even if the setting of Sat-Seconds is not reached.

11.5.7 Changing the Speed of the Pleth Wave

In the [**SpO2 Setup**] menu, select [**Sweep**] and then select the appropriate setting. The faster the waveform sweeps, the wider the waveform is.

11.5.8 Zooming PI Value

For Masimo SpO₂ module, you can display PI value in larger characters for better view. To zoom in the display of PI value, set [**PI Zoom**] to [**Yes**] from the [**SpO2 Setup**] menu.

11.5.9 Setting the Alarm Level for SpO₂ Sensor Off Alarm

Select [**Alarm Setup** >>] from the [**User Maintenance**] menu. You can set the [**SpO2SensorOff Lev.**] in the pop-up menu.

11.5.10 Setting the SpO₂ Tone Mode

Select [Others >>] from the [User Maintenance] menu. In the pop-up menu, you can set [SpO2 Tone] as [Mode 1] or [Mode 2].

WARNING

• The same SpO₂ tone mode shall be used for the same monitors in a single area.

11.6 Measurement Limitations

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

- Ambient light
- Physical movement (patient and imposed motion)
- Diagnostic testing
- Low perfusion
- Electromagnetic interference, such as MRI environment

- Electrosurgical units
- Dysfunctional haemoglobin, such as carboxyhemoglobin (COHb)and methemoglobin (MetHb)
- Presence of certain dyes, such as methylene and indigo carmine
- Inappropriate positioning of the SpO₂ sensor, or use of incorrect SpO₂ sensor.
- Drop of arterial blood flow to immeaurable level caused by shock, anemia, low temperature or vasoconstrictor.

11.7 Masimo Information



Masimo Patents

This posting serves as notice under 35 U.S.C.§287(a) for Masimo patents: http://www.masimo.com/patents.htm.

No Implied License

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

11.8 Nellcor Information



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

11.9 Troubleshooting

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

NOTE

• Never try to disassemble the equipment or supplied accessories. There are no internal user-serviceable parts.

Symptoms	Possible Cause	Correction Action
Dashes "" display in place of	Measurement is invalid.	Check that the sensor is properly applied. Change
numerics.		the application site if necessary.
Do not see SpO ₂ parameter	Parameter not configured to display.	Switch the SpO ₂ monitoring function on as
tiles in display.		described in section 3.12.1 Switching the
		Parameters On/Off.
Unable to obtain SpO ₂ reading	Patient has poor perfusion	Switch limbs/notify physician
	Sensor not on patient	Check if the "SPO2 Sensor Off" alarm is reported.
		If so, reapply the sensor.
		If not, contact the service personnel.
	Cables loose/not connected	Check if the sensor and cable are properly
		connected.
	Ambient light	Check if the " SpO2 Too Much Light " alarm is
		reported. If so, move the sensor to a place with
		lower level of ambient light or cover the sensor to
		minimize the ambient light.
No SpO ₂ waveform	Waveform not selected to display	Switch the SpO ₂ monitoring function on as
		described in 3.12.1 Switching the Parameters
		On/Off.
	Cable or sensor not plugged in	Check that the cable is properly connected and
		sensor securely applied.
Low amplitude SpO ₂ signal	SpO ₂ sensor on same limb as cuff	Check that the sensor is properly applied. Change
		the application site if necessary.
	Patient has poor perfusion	Change the application site.

FOR YOUR NOTES

12.1 Introduction

The patient monitor uses the oscillometric method for measuring the non-invasive blood pressure (NIBP). This measurement can be used for adults, pediatrics and neonates.

Automatic non-invasive blood pressure monitoring uses the oscillometric method of measurement. To understand how this method works, we'll compare it to the auscultative method. With auscultation, the clinician listens to the blood pressure and determines the systolic and diastolic pressures. The mean pressure can then be calculated with reference to these pressures as long as the arterial pressure curve is normal.

Since the monitor cannot hear the blood pressure, it measures cuff pressure oscillation amplitudes. Oscillations are caused by blood pressure pulses against the cuff. The oscillation with the greatest amplitude is the mean pressure. This is the most accurate parameter measured by the oscillometric method. Once the mean pressure is determined, the systolic and diastolic pressures are calculated with reference to the mean.

Simply stated, auscultation measures systolic and diastolic pressures and the mean pressure is calculated. The oscillometric method measures the mean pressure and determines the systolic and diastolic pressures.

As specified by IEC 80601-2-30, NIBP measurement can be performed during electro-surgery and discharge of defibrillator.

NIBP diagnostic significance must be decided by the doctor who performs the measurement.

NOTE

 Blood pressure measurements determined with this device are equivalent to those obtained by a trained observer using the cuff/stethoscope auscultatory method or an intra-arterial blood pressure measurement device, within the limits prescribed by the American National Standard, Manual, electronic, or automated sphygmomanometers.

WARNING

- Be sure to select the correct patient category setting for your patient before measurement. Do not apply the higher adult settings for pediatric or neonatal patients. Otherwise it may present a safety hazard.
- Do not measure NIBP on patients with sickle-cell disease or on the limb where skin damage has occurred or is expected.
- Use clinical judgment to determine whether to perform frequent unattended blood pressure measurements on patients with severe blood clotting disorders because of the risk of hematoma in the limb fitted with the cuff.
- Do not use the NIBP cuff on a limb with an intravenous infusion or arterial catheter in place. This could cause tissue damage around the catheter when the infusion is slowed or blocked during cuff inflation.
- NIBP reading can be affected by the measurement site, the position of the PATIENT, exercise, or the patient's physiologic condition. If you doubt the NIBP measurements, determine the patient's vital signs by alternative means and then verify that the monitor is working correctly.
- Do not apply cuff on the arm on the side of a mastectomy.
- Do not modify or replace connectors of the NIBP air hose except with mindray-approved connectors. Use neonatal and infant cuffs with CM1901 hoses only. Use pediatric/adult cuffs with CM1903 hoses only.
- Never connect intra-arterial or intra-venous lines, or any other incompatible connectors to the NIBP hose. This can cause serious in jury or death.
- Avoid placing cuff on patient in a manner that can lead to a hose becoming kinked (hose kinking may cause inaccurate readings).

12.3 Measurement Limitations

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

The measurement may be inaccurate or impossible:

- If a regular arterial pressure pulse is hard to detect
- With excessive and continuous patient movement such as shivering or convulsions
- With cardiac arrhythmias
- Rapid blood pressure changes
- Severe shock or hypothermia that reduces blood flow to the peripheries
- Obesity, where a thick layer of fat surrounding a limb dampens the oscillations coming from the artery

12.4 Measurement Methods

There are three methods of measuring NIBP:

- Manual: measurement on demand.
- Auto: continually repeated measurements at set intervals.
- STAT: continual rapid series of measurements over a five minute period, then return to the previous mode.

12.5 Setting Up the NIBP Measurement

12.5.1 Preparing the Patient

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

- Comfortably seated;
- Legs uncrossed;
- Feet flat on the floor;
- Back and arm supported; and,
- Middle of the cuff at the level of the right atrium of the heart.

NOTE

- It is recommended that the patient relax as much as possible before performing measurement and that the patient does not talk during measurement.
- It is recommended that 5 min should elapse before the first reading is taken.
- The operator shall not touch cuff and tubing during NIBP measurement.

12.5.2 Preparing to Measure NIBP

To prepare for the NIBP measurement:

- 1. Power on the monitor.
- Verify that the patient category is correct. If not, select the [Patient Setup] QuickKey → [Patient Demographics]
 → [Patient Cat.] and set the patient category to [Adu], [Ped] or [Neo].
- 3. Plug the air tubing into the NIBP connector on the patient monitor.
- 4. Select a correct sized cuff and then apply it as follows:
 - Determine the patient's limb circumference.
 - Select an appropriate cuff by referring to the limb circumference marked on the cuff. The width of the cuff should be 40% (50% for neonates) of the limb circumference, or 2/3 of the upper arm's length. The inflatable part of the cuff should be long enough to encircle at least 50% to 80% of the limb.
 - Apply the cuff to the patient's upper arm or leg and make sure the Φ marking on the cuff matches the artery location. Do not wrap the cuff too tightly around the limb. Otherwise, it may cause discoloration, and ischemia of the extremities. Make sure that the cuff edge falls within the marked range. If it does not, use a larger or smaller cuff that will fit better.

5. Connect the cuff to the air tubing. Avoid compression or restriction of pressure tubes. Air must pass unrestricted through the tubing.

WARNING

• Continuous CUFF pressure due to connection tubing kinking may cause blood flow interference and resulting harmful injury to the patient.

NOTE

• Equipment use is restricted to one patient at a time.

12.5.3 Starting and Stopping Measurements

Select the [**NIBP Measure**] QuickKey and you can start the desired measurement from the popup menu. You can select [**Stop All**] QuickKey to stop all NIBP measurements. You can also start and stop measurements by using the [**NIBP**] hardkey on the monitor's front panel.

12.5.4 Correcting the Measurement if Limb is not at Heart Level

The cuff should be applied to a limb at the same level as the patient's heart. If the limb is not at the heart level, to the displayed value:

- Add 0.75 mmHg (0.10 kPa) for each centimetre higher, or
- Deduct 0.75 mmHg (0.10 kPa) for each centimeter lower.

12.5.5 Enabling NIBP Auto Cycling and Setting the Interval

To enable the NIBP auto cycling and set the interval:

- 1. Select the NIBP parameter window to enter the [**NIBP Setup**] menu.
- 2. Select [Interval] and then select a desired time interval. Selecting [Manual] switches to manual mode.
- 3. Start a measurement manually. The monitor will then automatically repeat NIBP measurements at the set time interval.

12.5.6 Enabling Measurement on Clock

In auto measurement mode, if the clock is enabled, the monitor automatically synchronizes NIBP automatic measurements with the real time clock. For example, if [Interval] is set to [20 min], and you start NIBP auto measurement at 14: 03, the next measurement will be taken at 14: 20, and then at 14:40, 15:00, and so on.

To enable measurement on clock:

- 1. Select the NIBP parameter window to enter the [**NIBP Setup**] menu.
- 2. Set [Clock] to [On].

• Measurement on clock is effective only when NIBP Interval is set to [5min] or an option greater than 5 min.

12.5.7 Starting a STAT Measurement

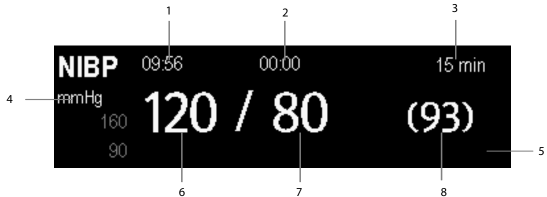
To start a STAT measurement:

- 1. Select the NIBP parameter window to enter the [NIBP Setup] menu.
- 2. Select [NIBP STAT]. The STAT mode initiates 5 minutes of continuous, sequential, automatic NIBP measurements.

WARNING

• Continuous non-invasive blood pressure measurements may cause purpura, ischemia and neuropathy in the limb with the cuff. Inspect the application site regularly to ensure skin quality and inspect the cuffed limb for normal color, warmth and sensitivity. If any abnormity occurs, move the cuff to another site or stop the blood pressure measurements immediately.

12.6 Understanding the NIBP Numerics



The NIBP display shows numerics only as below. Your display may be configured to look slightly different.

- 1. Time of last measurement
- 2. Time remaining to next measurement
- 3. Measurement mode
- 4. Unit of pressure: mmHg or kPa
- 5. Prompt message area: shows NIBP-related prompt messages
- 6. Systolic pressure
- 7. Diastolic pressure
- 8. Mean pressure obtained after the measurement and cuff pressure obtained during the measurement

If the NIBP measurement exceeds the measurement range, "---" will be displayed.

12.7 Changing NIBP Settings

By selecting the NIBP parameter window, you can enter the [**NIBP Setup**] menu.

12.7.1 Setting the Initial Cuff Inflation Pressure

You can set the initial cuff inflation pressure manually. In the [**NIBP Setup**] menu, select [**Initial Pressure**] and then select the appropriate setting.

NOTE

• For known hypertensive patients, you need to set initial cuff pressure to a higher value to reduce the measurement time.

12.7.2 Setting NIBP Alarm Properties

Select [Alarm Setup >>] from the [NIBP Setup] menu. You can set the alarm properties for this parameter in the popup menu.

12.7.3 Displaying NIBP List

Select [**Screens**] QuickKey→[**Screen Setup**]. You can set [**NIBP List**] to be displayed on the screen, where the most recent NIBP measurements display. Then, multiple sets of most recent NIBP measurements will be displayed. And the displayed PR is derived from NIBP.

NIBP	Lis	st	PR	Time	
120	7	80	(93)	60	16:53
120	7	80	(93)	60	15:52

You can not display NIBP list in some screens such as the big numerics screen.

12.7.4 Setting the Pressure Unit

Select [Unit Setup >>] from the [User Maintenance] menu. In the popup menu, select [Press. Unit] and then select [mmHg] or [kPa].

12.7.5 Switching On NIBP End Tone

The monitor can issue a reminder tone at the completion of NIBP measurement. The NIBP End Tone is off by default. You can switch it on by accessing the [**NIBP Setup**] menu.

12.8 Assisting Venous Puncture

You can use the NIBP cuff to cause sub-diastolic pressure to block the venous blood vessel and therefore help venous puncture.

- 1. Select [VeniPuncture >>] from the [NIBP Setup] menu. In the popup menu, verify that the [Cuff Press.] value is appropriate. Change it if necessary.
- 2. Select [VeniPuncture].
- 3. Puncture vein and draw blood sample.
- 4. Select the [**NIBP**] hardkey on the monitor's front panel, or the [**Stop All**] QuickKey to deflate the cuff. The cuff deflates automatically after a set time if you do not deflate it.

During measurement, the NIBP display shows the inflation pressure of the cuff and the remaining time in venous puncture mode.

FOR YOUR NOTES

13.1 Introduction

The equipment is used to monitor skin temperature and core temperature. You can simultaneously monitor two temperature sites using the monitor.

13.2 Safety

WARNING					
• Verify that the probe detection program	m works correctly before monitoring. Remove the temperature probe				
cable from the T1 or T2 connector, and	the monitor can display the message [T1 Sensor Off] or [T2 Sensor				
Off] and give alarm tones correctly.					

13.3 Making a Temp Measurement

The equipment is intended for monitoring skin temperature and core temperature.

- 1. Select an appropriate probe for your patient according to patient type and measuring site.
- 2. If you are using a disposable probe, connect the probe to the temperature cable.
- 3 Plug the probe or temperature cable to the temperature connector.
- 4. Attach the probe to the patient correctly.
- 5. Check that the alarm settings are appropriate for this patient.

13.4 Understanding the Temp Display

The temperature monitoring is displayed on the monitor as three numerics: T1, T2 and TD. By selecting this area, you can enter the [Alarm Setup] menu.



13.5 Setting the Temperature Unit

Select [**Unit Setup** >>] from the [**User Maintenance**] menu. In the pop-up menu, select [**Temp Unit**] and then select [**°C**] or [**°F**].

14.1 Introduction

The cPM 12 monitor can monitor 2 invasive blood pressures (using an external module). The monitor can display the systolic, diastolic and mean pressures and a waveform for each pressure.

14.2 Safety

WARNING

- Use only pressure transducers specified in this manual. Never reuse disposable pressure transducers.
- Make sure that the applied parts never contact other conductive parts.
- To reduce the hazard of burns during high-frequency surgical procedure, ensure that the monitor's cables and transducers never come into contact with the high-frequency surgical units.
- When using accessories, their operating temperature should be taken into consideration. For details, refer to instructions for use of accessories.
- The neutral electrode of the electro-surgery unit (ESU) shall properly contact the patient. Otherwise, burns may result.

14.3 Zeroing the Transducer

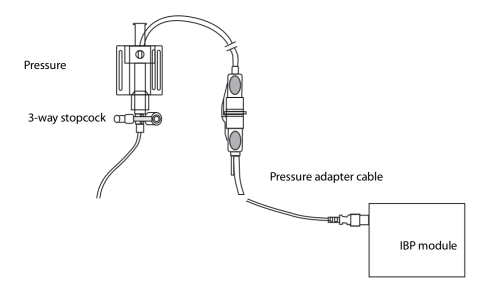
To avoid inaccurate pressure readings, the monitor requires a valid zero. Zero the transducer in accordance with your hospital policy (at least once per day).

Zero whenever:

- A new transducer or adapter cable is used.
- You reconnect the transducer cable to the module.
- The monitor restarts.
- You doubt the readings.

To zero the transducer:

1. Turn off the stopcock to the patient.



- 2. Vent the transducer to the atmospheric pressure by turning on the stopcock to the air.
- In the setup menu for the pressure (e.g. Art), select [Art Zero >>]→[Zero]. During zero calibration, the [Zero] button appears dimmed. It recovers after the zero calibration is completed. To zero all IBP channels, select [Zero IBP] hotkey, and then select [Zero All Channels] in the pop-up menu.
- 4. After the zero calibration is completed, close the stopcock to the air and open the stopcock to the patient.

NOTE

• Your hospital policy may recommend that the ICP transducer is zeroed less frequently than other transducers.

14.4 Setting Up the Pressure Measurement

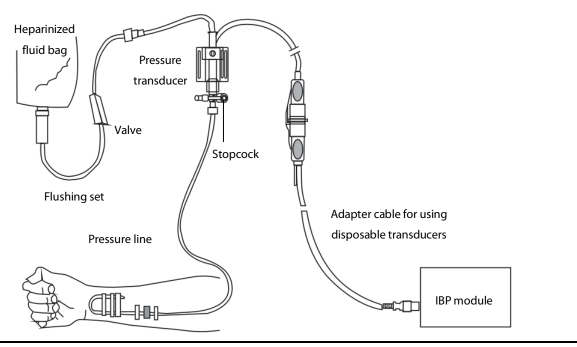
To set up the pressure measurement:

- 1. Plug the pressure cable into the IBP connector.
- 2. Prepare the flush solution.
- 3. Flush the system to exhaust all air from the tubing. Ensure that the transducer and stopcocks are free of air bubbles.

WARNING

• If air bubbles appear in the tubing system, flush the system with the infusion solution again. Air bubble may lead to wrong pressure reading.

- 4. Connect the pressure line to the patient catheter.
- 5. Position the transducer so that it is level with the heart, approximately at the level of the midaxillary line.
- 6. Select the appropriate label.
- 7. Zero the transducer. After a successful zeroing, turn off the stopcock to the air and turn on the stopcock to the patient.

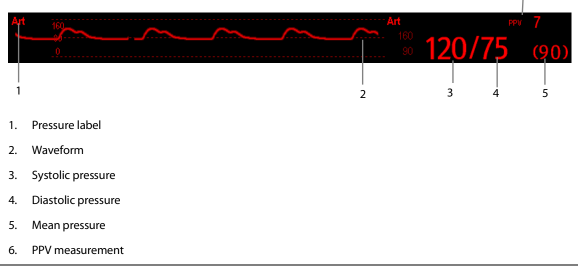


WARNING

• If measuring intracranial pressure (ICP) with a sitting patient, level the transducer with the top of the patient's ear. Incorrect leveling may give incorrect values.

14.5 Understanding the IBP Display

The IBP measurement is displayed on the monitor as a waveform and numeric pressures. The figure below shows the waveform and numerics for the Art pressure. For different pressures, this display may be slightly different.



6

For some pressures, the parameter window may show the mean pressure only. For different pressures, their defaults unit may be different. If the Art and ICP pressures are measured simultaneously, the ICP parameter area will display numeric CPP, which is obtained by subtracting ICP from the Art mean.

14.6 Changing IBP Settings

14.6.1 Changing a Pressure for Monitoring

To change a pressure for monitoring:

- 1. Select the pressure you want to change to enter its setup menu. In the menu, there is a figure showing the current IBP measurement connector.
- 2. Select [Label] and then select your desired label from the list. Labels already used cannot be selected.

Label	Description	Label	Description
PA	Pulmonary artery pressure	CVP	Central venous pressure
Ao	Aortic pressure	LAP	Left atrial pressure
UAP	AP Umbilical arterial pressure		Right atrial pressure
BAP	AP Brachial arterial pressure		Intracranial pressure
FAP	Femoral arterial pressure	UVP	Umbilical venous pressure
Art	Arterial blood pressure	LV	Left ventricular pressure
P1 to P4	Non-specific pressure label		

NOTE

• When two pressures are detected having the same label, the monitor changes one pressure label to a currently unused one.

14.6.2 Setting Alarm Properties

Select [**Alarm Setup** >>] from the parameter setup menu. You can set alarm properties for this parameter in the pop-up menu.

14.6.3 Changing Averaging Time

The IBP value displayed on the monitor screen is the average of data collected within a specific time. The shorter the averaging time is, the quicker the monitor responds to changes in the patient's blood pressure. Contrarily, the longer the averaging time is, the slower the monitor responds to changes in the patient's blood pressure, but the measurement accuracy will be improved. For critically ill patients, selecting shorter averaging time will help understanding the patient's state.

To set the averaging time, in the parameter setup menu, select [**Sensitivity**] and then select [**High**], [**Med**] or [**Low**], the corresponding averaging time is about 1 s, 8 s and 12 s respectively.

14.6.4 Setting Up the IBP Wave

In the setup menu for the pressure, you can:

- Select [Sweep] and then select the appropriate setting. The faster the wave sweeps, the wider the wave is.
- Select [Scale] and then select the appropriate setting. If [Auto] is selected, the size of the pressure's waveform will be adjusted automatically.
- Select [Filter] and then select the desired option.

14.6.5 Setting the Pressure Unit

Select [Unit Setup >>] from the [User Maintenance] menu. In the pop-up menu, select [Press. Unit] and then select [mmHg] or [kPa]. Select [CVP Unit] and then select [mmHg], [cmH2O] or [kPa].

14.6.6 Enabling PPV Measurement and Setting PPV Source

PPV indicates pulse pressure variation. To enable PPV measurement, set [PPV Measurement] to [On].

You can select PPV source by enabling the PPV measurement.

WARNING

- This monitor can calculate PPV from beat-to-beat values of any arterial pulsatile pressure. The circumstances under which the calculation of a PPV value is clinically meaningful, appropriate and reliable must be determined by a physician.
- The clinical value of the derived PPV information must be determined by a physician. According to recent scientific literature, the clinical relevance of PPV information is restricted to sedated patients receiving controlled mechanical ventilation and mainly free from cardiac arrhythmia.
- PPV calculation may lead to inaccurate values in the following situations:
 - at respiration rates below 8 rpm
 - during ventilation with tidal volumes lower than 8 ml/kg
 - for patients with acute right ventricular dysfunction ("cor pulmonale").
- The PPV measurement has been validated only for adult patients.

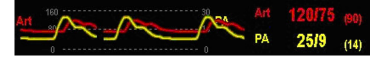
14.7 Overlapping IBP Waveforms

The IBP waveforms can be displayed together. To combine IBP waveforms,

- 1. Select [Main Menu]→[Screen Setup>>]→[Screen Layout>>] to access the [Screens] window.
- 2. Select the [Screen Setup] tab.
- 3. In Area A, select [**IBP Overlap**] from the drop-down list, and then select the IBP waves to be overlapped on the left side of the same line.



4. Select imes to save the setting and exit the window. The main screen will display the overlapped IBP waves.



Selecting the overlapped IBP waveforms on the main screen opens the [**Overlapping Waveform Setup**] menu, where you can:

- Set [Left Scale] and [Right Scale] and then set the scales for the overlapped waveforms. The left scale is for Art, LV, Ao, FAP, BAP, UAP, and the arterial waveforms of P1~P4; the right scale is for CVP, ICP, LAP, RAP, UVP, and the venous waveforms of P1~P4.
- Set [**CVP Scale**] individually if the CVP waveform is combined and CVP unit is different from IBP unit.
- Set [**PA Scale**] individually if PA waveform is combined.
- Set [Gridlines] to [On] or [Off] to show gridlines or not in the overlapped waveform area.
- Select [**Sweep**] and then set the sweep speed for the overlapped waveforms.
- Select [Filter] and then set the filter for the overlapped waveforms.

You can also change above settings from corresponding IBP setup menu.

Note

• The CVP scale changes simultaneously with right scale. The unit of CVP scale is consistent with CVP parameter unit.

14.8 Measuring PAWP

Pulmonary Artery Wedge Pressure (PAWP) values, used to assess cardiac function, are affected by fluid status, myocardial contractility, and valve and pulmonary circulation integrity.

Obtain the measurement by introducing a balloon-tipped pulmonary artery flotation catheter into the pulmonary artery. When the catheter is in one of the smaller pulmonary arteries, the inflated balloon occludes the artery allowing the monitor to record changes in the intrathoracic pressures that occur throughout the respiration cycle.

The pulmonary wedge pressure is the left ventricular end diastolic pressure when the airway pressure and valve function are normal. The most accurate PAWP values are obtained at the end of the respiration cycle when the intrathoracic pressure is fairly constant and the artifact caused by respiration is minimal.

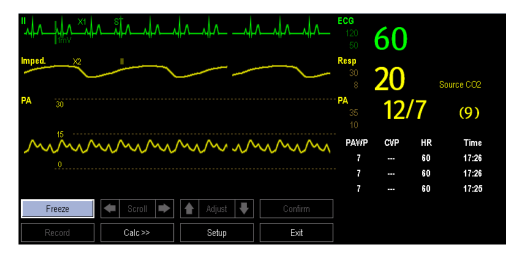
WARNING

• PAWP monitoring is not intended for neonatal patients.

14.8.1 Preparing to Measure PAWP

To prepare for the PAWP measurement:

- Prepare the same accessories as in the C.O. measurement. Connect the parts such as catheter, syringe, etc. following the C.O. measurement steps and use the balloon inflation port. Refer to section 15.4 Setting Up the C.O. Measurement for more information.
- 2. Connect the PAWP cable into the IBP connector on the monitor. Since PAWP is measured on PA, selecting [**PA**] as the IBP label is recommended.
- 3. Select the PA parameter window or waveform area to enter its setup menu. Then, select [**PAWP**] to enter the PAWP measurement window. You can also enter the PAWP measurement window from the P1-P4 parameter window.



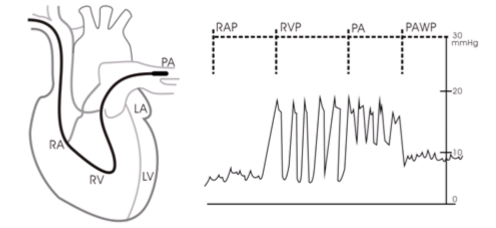
NOTE

• After entering the PAWP measurement window, the monitor will turn off the PA alarm automatically.

14.8.2 Setting Up the PAWP Measurement

To setup the PAWP measurement:

1. Wedge the flotation catheter into the pulmonary artery. Then inflate the balloon and pay attention to the PA waveform changes on the screen.



- 2. After obtaining a stable PAWP waveform, press the [Freeze] key to freeze the waveform and deflate the balloon.
- 3. You can adjust the PAWP scale to an appropriate position by adjusting A or solution. Press the [**Confirm**] key to save one PAWP measurement.
- 4. If you need to start a new measurement, select [Next Measure].

WARNING

- Prolonged inflation can cause pulmonary hemorrhage, infarction or both. Inflate the balloon for the minimum time necessary to get an accurate measurement.
- If the PAWP is greater than the PA (systolic), deflate the balloon and report the incident in accordance with hospital policy. Because the pulmonary artery could be accidentally ruptured, and the PAWP value derived will not reflect the patient's hemodynamic state, but will merely reflect the pressure in the catheter or balloon.

14.8.3 Understanding the PAWP Setup Menu

Select [Setup] to enter the [PAWP Setup] menu. In this menu, you can:

- Select an ECG lead wave as the first reference wave.
- Select a respiration wave as the second reference wave.
- Select a sweep speed for the displayed waveform.
- Change the size of the PA waveform by adjusting the scale height.

14.9 Troubleshooting

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

CAUTION

• Never try to disassemble the equipment or supplied accessories. There are no internal user-serviceable parts.

Symptoms	Possible Cause	Correction Action	
Damped invasive	Air bubbles in tubing	Eliminate air from tubing as described in section	
waveform		14.4 Setting Up the Pressure Measurement.	
	Kinked catheter	Change the position of catheter.	
	Blood in tubing	Pressurize the solution bag to 300 mmHg. For	
		details, refer to the instructions for use of the	
		solution bag.	
IBP not displayed/no IBP	Improper setup	Check display setup in monitor setup.	
waveform	Cable not plugged in	Check that the cables are properly connected.	
	Transducer not connected.	Check that the transducer is properly connected.	
	Stopcock turned improperly.	Check that the stopcock is turned to the correct	
		position.	
	Transducer not zeroed	Check and zero the transducer as described in	
		section 14.3 Zeroing the Transducer.	
Dashes "" display in place	The measured result is invalid or out of	Change to a pulsatile label.	
of numerics.	range.		
	IBP might be set to non-pulsatile labels		
	like CVP, LA, RA, and ICP.		
Abnormally high or low	Transducer too High or too Low.	Adjust the position of the transducer and make	
readings		sure that it is level with the heart, approximately at	
		the level of the midaxillary line.	
		Zero the transducer as described in 14.3 Zeroing	
		the Transducer.	
Unable to Zero	Stopcock not open to atmosphere.	Check the transducer and make sure the stopcock	
		is turned to the air.	
PAWP button disabled	One IBP channel must be labeled PA	Label an IBP channel as PA. (Also Label an IBP	
		channel as P1/P2/P3/P4, it will automatically	
		change to PA)	

FOR YOUR NOTES

15.1 Introduction

The cardiac output (C.O.) measurement invasively measures cardiac output and other hemodynamic parameters using the right heart (atria) thermodilution method. A cold solution of known volume and temperature is injected into the right atrium through the proximal port of a pulmonary artery (PA) catheter. The cold solution mixes with the blood in the right ventricle and the change in blood temperature is measured with a thermistor at the distal end of the catheter in the pulmonary artery. The temperature change is displayed as a curve in the C.O. split screen, and the monitor calculates the C.O. value from this curve. The C.O. value is inversely proportional to the area under the curve. As cardiac output varies continuously, a series of measurements must be carried out to achieve a reliable C.O. average value. Always use the average of multiple thermodilution measurements for therapy decisions. The monitor is capable of storing 6 measurements.

WARNING

• C.O. monitoring is restricted to adult patients only.

15.2 Understanding the C.O. Display

The C.O. measurement is displayed on the monitor as numeric C.O., C.I. and TB in the C.O. parameter window as shown below. To enter the [C.O. Setup] menu, select the C.O. parameter window.



- 1. Cardiac output
- 2. Time at which the C.O. average is calculated
- 3. Cardiac index
- 4. Blood temperature

15.3 Influencing Factors

The factors that affect cardiac output are:

- temperature of injectate solution,
- volume of injectate solution,
- patient's baseline blood temperature,
- patient's inspiratory/expiratory cycle,
- placement of catheter with relation to proximity of lung field,
- the catheter itself,
- the patient rhythm and hemodynamic status, and
- any other rapid IV solutions which are infused while the C.O. measurement is being performed.

Followings are some technical suggestions to obtain accurate C.O.:

- Injectate solution must be cooler than the patient's blood.
- Inject solution rapidly and smoothly.
- Inject at end of expiration.

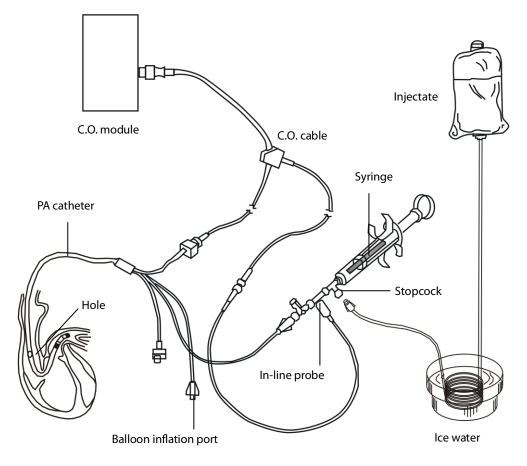
15.4 Setting Up the C.O. Measurement

WARNING

• Use only accessories specified in this manual. Make sure that the accessories never come into contact with conductive parts.

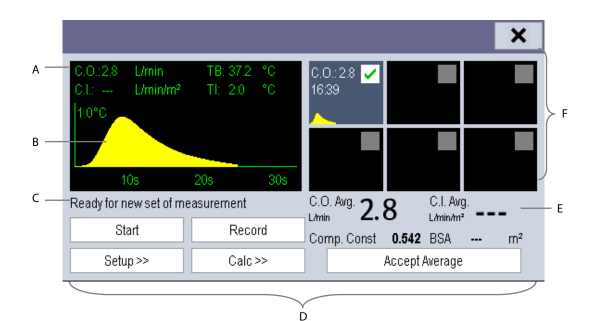
To set up the C.O. measurement:

- 1. Connect the C.O. cable to the C.O. connector on the monitor.
- 2. Interconnect the C.O. module, catheter and syringe as shown below. Make sure that:
 - The module is securely inserted.
 - The PA catheter is in place in the patient.
 - The C.O. cable is properly connected to the module.



NOTE

- The above picture illustrates when using a TI sensor (PN 6000-10-02079). The connection may be different if other TI sensors are used.
- 3. Select the C.O. parameter window to enter the [**C.O. Setup**] menu. Check if the height and weight are appropriate for your patient. Change if necessary.
- 4. In the [**C.O. Setup**] menu:
 - Check that the correct computation constant is entered. Refer to the Instruction for Use of pulmonary artery catheter to determine the [Comp. Const] according to the entered injectate volume and temperature. To change the computation constant, select [Comp. Const] and then enter the correct value. When a new catheter is used, the computation constant should be adjusted in accordance with the manufacturer's instructions for use.
 - Set the [Auto TI] to [Manual] or [Auto]. If you select [Auto], the system automatically detects the injectate temperature, and the [Manual TI(°C)] is disabled. If you select [Manual], you need to enter the injectate temperature at [Manual TI(°C)] manually.
 - Set the [Measuring Mode] to [Manual] or [Auto]. In [Auto] mode, the monitor automatically takes C.O. measurement after establishing a baseline blood temperature. In [Manual] mode, you need to click the [Start] button in the C.O. measurements window when the monitor is ready for new C.O. measurement.
- 4. Select [Enter C.O. Screen] to enter the C.O. measurements window.



- A. Currently measured numeric
- B. Currently measured C.O. curve
- C. Prompt message area
- D. Buttons
- E. Averaged values
- F. Measurement windows
- 6. Proceed as follows.
 - In [Manual] measure mode, select the [Start] button and then inject the solution quickly when you see the message [Ready for new set of measurement]. As shown in the figure above, during the measurement, the currently measured thermodilution curve is displayed. At the end of the measurement, the thermodilution curve is transferred to one of the 6 measurement windows and the monitor prompts you to wait for a certain period of time before starting a new measurement.
 - In [Auto] measure mode, the monitor consecutively takes C.O. measurements automatically without the need for pressing the [Start] button between two measurements. A new thermodilution measurement is possible as soon as the message [Inject now!] is displayed on the screen. The monitor automatically detects further thermodilution measurements.
- 7. Consecutively take 3 to 5 single measurements as instructed by Step 6.

A maximum of 6 measurements can be stored. If you perform more than six measurements without rejecting any, the oldest will automatically be deleted when a seventh curve is stored. Select from the 6 measurement curves and the system will automatically calculate and display the averaged C.O. and C.I. values. Then select the [Accept Average] button to accept and store the averaged values.

When injecting, the stopcock to the PA catheter is open and the stopcock to the injectate solution is closed. After the measurement is completed, turn off the stopcock to the PA catheter and turn on the stopcock to the injectate solution, and then draw the injectate solution into the injectate syringe.

In the buttons area, you can:

- Select [Start] to start a C.O. measurement.
- Select [**Stop**] to stop the current measurement.
- Select [Cancel] during a measurement to cancel the measurement. Selecting it after a measurement deletes the measured results.
- Select [Record] to print out the curves selected for average calculation, numerics and averaged values by the recorder.
- Select [Setup >>] to access the [C.O. Setup] menu.
- Select [Calc >>]→[Hemodynamic >>] to access the [Hemodynamic Calculation] menu.

The system can automatically adjust the X-axis scale range to 30 s or 60 s and Y-axis scale range to 0.5°C, 1.0°C, or 2.0°C.

CAUTION

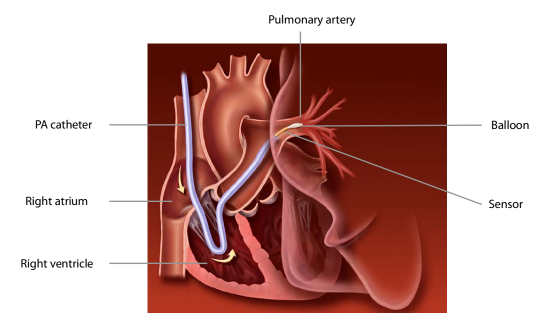
• Starting a measurement without blood temperature being stable may cause measuring failure.

NOTE

- During a cardiac output measurement, blood temperature alarms are inactive.
- Please refer to the Instructions for Use of the pulmonary artery catheter to determine the [Comp. Const] and the volume of injectate.

15.5 Measuring the Blood Temperature

As shown below, the blood temperature is measured with a temperature sensor at the distal end of the catheter in the pulmonary artery. During C.O. measurements, blood temperature alarms are suppressed to avoid false alarms. They will automatically recover as soon as the C.O. measurements are completed.



15.6 Changing C.O. Settings

15.6.1 Setting the Temperature Unit

Select [**Unit Setup** >>] from the [**User Maintenance**] menu. In the pop-up menu, select [**Temp Unit**] and then select [°**C**] or [°**F**].

15.6.2 Setting Alarm Properties

Select [Alarm Setup >>] from the [C.O. Setup] menu. You can set alarm properties for this parameter in the pop-up menu.

16.1 Introduction

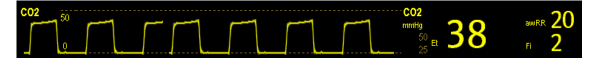
 CO_2 monitoring is a continuous, non-invasive technique for determining the concentration of CO_2 in the patient's airway by measuring the absorption of infrared (IR) light of specific wavelengths. CO_2 has its own absorption characteristic and the amount of light passing the gas probe depends on the concentration of the measured CO_2 . When a specific band of IR light passes through respiratory gas samples, some of IR light will be absorbed by the CO_2 molecules. The amount of IR light transmitted after it has been passed through the respiratory gas sample is measured with a photodetector. From the amount of IR light measured, the concentration of CO_2 is calculated.

Sidestream/Microstream measurement samples expired patient gas at a constant sample flow from the patient's airway and analyzes it with a CO_2 sensor built into the CO_2 module.

The sidestream and microstream CO₂ measurement can be used, with specified accessories, with intubated and non-intubated adult, pediatric, and neonatal patients. With intubated patients, a sample of the respiratory gas is drawn from the patient's breathing circuit through an airway adapter and a gas sampling line. With non-intubated patients, the gas sample is drawn through a nasal cannula.

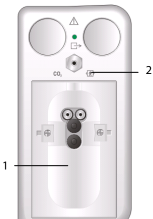
The measurement provides:

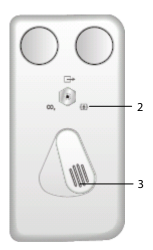
- 1. A CO₂ waveform.
- 2. End tidal CO₂ value (EtCO₂): the CO₂ value measured at the end of the expiration phase.
- 3. Fraction of inspired CO_2 (FiCO₂): the smallest CO_2 value measured during inspiration.
- 4. Airway respiration rate (awRR): the number of breaths per minute, calculated from the CO₂ waveform.



16.2 CO₂ Module

This monitor uses an external module to perform CO₂ monitoring.





- 1. CO₂ watertrap seat
- 2. Gas outlet
- 3. Connector for sampling line

16.3 Preparing to Measure CO₂

WARNING

- Check that the alarm limit settings are appropriate before taking measurement.
- Eliminate the exhausted gas before performing the measurement.

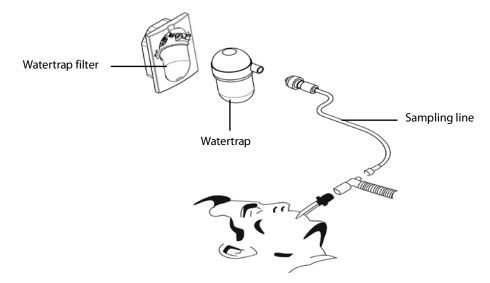
NOTE

• Perform the measurement in a well-ventilated environment.

16.3.1 Making a Sidestream CO₂ Measurement

To make a sidestream CO2 measurement:

1. Attach the watertrap to the module and then connect the CO₂ components as shown below.



- 2. The CO₂ module needs time to warm up to reach operating temperature. The message [**CO2 Sensor Warmup**] is displayed during warm-up.
- 3. After warm-up is finished, you can perform CO₂ measurements.

CAUTION

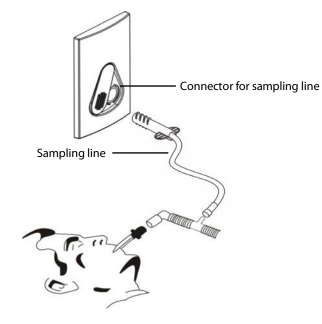
- The watertrap collects water drops condensed in the sampling line and therefore prevents them from entering the module. To avoid blocking the airway, empty the watertrap container whenever half full. Dispose of accumulated fluids in accordance with hospital policy or your local regulations.
- The watertrap has a filter preventing bacterium, water and secretions from entering the module. After long-term use, dust or other substances may compromise the performance of the filter or even block the airway. In this case, replace the watertrap. It is recommended that you replace the watertrap once a month, or when the watertrap is found leaky, damaged or contaminated.
- Do not apply adult watertrap to the neonate patient. Otherwise, patient injury could result.

NOTE

• To extend the life of the watertrap and module, disconnect the watertrap and set the CO₂ module operating mode to the Standby mode when CO₂ monitoring is not required.

16.3.2 Making a Microstream CO₂ Measurement

Connect the sampling line to the module and then connect the CO_2 components as shown below. After warm-up is finished, you can perform CO_2 measurements.



16.4 Changing CO₂ Settings

16.4.1 Accessing CO₂ Menus

By selecting the CO₂ parameter window, you can access the [**CO2 Setup**] menu.

16.4.2 Entering the Standby Mode

The Standby mode of the CO₂ module relates to the Standby mode of the monitor as follows:

- If the monitor enters the Standby mode, the CO₂ module will also enter the Standby mode.
- If the monitor exits the Standby mode, the CO₂ module will also exit the Standby mode.
- If the CO₂ module enters or exits the Standby mode, it will not affect the monitor.

To enter or exit the Standby mode manually:

- 1. Select [Operating Mode] in the [CO2 Setup] menu
- 2. Select [Standby] or [Measure].

When you set the sidestream CO_2 module to the Standby mode, the CO_2 gas sample intake pump automatically sets the sample flow rate to zero. When exiting the Standby mode, the CO_2 module continues to work at the previously set sample flow rate.

For the sidestream CO_2 module, you can set a delay time after which, the CO_2 module will enter the Standby mode if no breath is detected.

For the microstream CO_2 module, you can also set a period of time after which the CO_2 module enters the Standby mode if no breath is detected, or since the CO_2 module was powered on, or since the CO_2 module was switched to the measuring mode, or the automatic Standby time is re-set.

To set the Standby time:

- 1. Select the CO₂ parameter window to access the [**CO2 Setup**] menu:
- 2. Select [Auto Standby] and then select the appropriate setting.

16.4.3 Setting the CO₂ Unit

To set the CO₂ unit:

- 1. Select [Main Menu] \rightarrow [Maintenance >>] \rightarrow [User Maintenance] \rightarrow enter the required password \rightarrow [Unit Setup >>].
- 2. In the Unit Setup menu, select [CO2 Unit] and then select [mmHg], [%] or [kPa].

16.4.4 Setting up Gas Compensations

WARNING

• Make sure to use the appropriate compensations. Inappropriate compensations may cause inaccurate measurement values and result in misdiagnosis.

For the sidestream CO₂ module:

- 1. Select the CO₂ parameter window to access the [**CO2 Setup**].
- 2. According to the actual condition, set the concentration required for the following compensations:
 - ♦ [O2 Compen]
 - [N2O Compen]
 - [Des Compen]

For the microstream CO₂ module, gas compensations are not required.

16.4.5 Setting up Humidity Compensation

Sidestream and microstream CO_2 modules are configured to compensate CO_2 readings for either Body Temperature and Pressure, Saturated Gas (BTPS), to account for humidity in the patient's breath, or Ambient Temperature and Pressure, Dry Gas (ATPD).

- 1. ATPD: $P_{co2}(mmHg) = CO_2(vol\%) \times P_{amb}/100$
- 2. BTPS: $P_{co2}(mmHg) = CO_2(vol\%) \times (P_{amb} 47)/100$

Where, P_{CO2} = partial pressure, vol% = CO₂ concentration, P_{amb} = ambient pressure, and unit is mmHg.

For the sidestream and microstream CO_2 module, you can set the humidity compensation on or off according to the actual condition. To set the humidity compensation:

- 1. Select the CO₂ parameter window to access the [**CO2 Setup**] menu, and then select [**BTPS Compen**].
- 2. Select either [**On**] for BTPS or [**Off**] for ATPD, depending on which compensation applies.

16.4.6 Setting the Apnea Alarm Delay

To set the apnea alarm delay:

- 1. Select the CO₂ parameter window to access the [**CO2 Setup**] menu.
- 2. Select [Apnea Delay] and then select the appropriate setting.

The monitor will alarm if the patient has stopped breathing for longer than the selected apnea time. The setting of [**Apnea Delay**] takes effect simultaneously to the Resp, CO₂, and AG modules.

WARNING

• The respiration measurement does not recognize the cause of apneas. It only indicates an alarm if no breath is detected when a pre-adjusted time has elapsed since the last detected breath. Therefore, it cannot be used for diagnostic purpose.

16.4.7 Choosing a Time Interval for Peak-Picking

For microstream modules, you can select a time interval for picking the highest CO_2 as the $EtCO_2$ and the lowest as the $FiCO_2$.

To set the time interval:

- 1. Select the CO₂ parameter window to access the [**CO2 Setup**] menu.
- 2. Select [Max Hold]
- 3. Select [Single Breath], [10 s], [20 s] or [30 s]
 - [Single Breath]: EtCO₂ and FiCO₂ are calculated for every breath.
 - [10 s], [20 s] or [30 s]: EtCO₂ and FiCO₂ are calculated using 10, 20 or 30 seconds of data.

16.4.8 Setting the Flow Rate

For the sidestream CO_2 module, you can change the sampling rate of respiratory gas in the patient's airway by setting the flow rate.

To set the flow rate:

- 1. Select the CO₂ parameter window to access the [**CO2 Setup**] menu.
- 2. Select an appropriate setting from [Flow Rate].

WARNING

• Please consider the patient's actual bearing capability and select the appropriate flow rate when setting the flow rate.

16.4.9 Setting up the CO₂ Wave

Select the CO₂ parameter window to access the [**CO2 Setup**] menu, in which you can:

- Select [Wave Type] and then select [Draw] or [Fill]:
 - [**Draw**]: The CO_2 wave is displayed as a curved line.
 - [Fill]: The CO₂ wave is displayed as a filled area.
- Select [Sweep], and then select the appropriate setting. The faster the wave sweeps, the wider the wave is.
- Change the size of the CO₂ waveform by adjusting the wave [**Scale**].

16.5 Setting RR Source

To set RR source:

- 1. Select the CO₂ parameter window to access the [**CO2 Setup**] menu.
- 2. Select [RR Source] and then select a source or [Auto] from the dropdown list.

The [**RR Source**] setting takes effect simultaneously to the Resp, CO₂, and AG modules. For details, please refer to section **9.9 Setting Respiration Rate Source**.

16.6 Barometric Pressure Compensation

Both sidestream and microstream CO_2 modules have the function of automatic barometric pressure compensation (the system automatically measures the barometric pressure which the monitor is exposed to).

16.7 Measurement Limitations

Some adverse effects can influence the CO₂ performance.

CAUTION

- The following factors may influence the accuracy of measurement:
 - Leaks or internal venting of sampled gas
 - Mechanical shock
 - ◆ Cyclic pressure up to 10 kPa (100 cmH₂O)
 - Other sources of interference, if any
- Measurement accuracy may be affected by the breath rate and I/E ratio as follow:
 - etCO₂ is within specification for breath rate \leq 60 bpm and I/E ratio \leq 1:1;
 - etCO₂ is within specification for breath rate \leq 30 bpm and I/E ratio \leq 2:1.
- Measurement accuracy is unspecified for breath rate larger than 60 bpm.

16.8 Leakage test

When the sidestream CO_2 module needs maintenance, the monitor displays the message in the CO_2 waveform window: [Need maintenance. Enter CO2 setup menu.] Then:

- Select [Main Menu]→[Maintenance >>]→[User Maintenance >>]→enter the required password→[Maintain CO2 >>].
- 2. Perform leakage test according to the prompt messages on the menu.

16.9 Troubleshooting the Sidestream CO₂ Sampling System

When the sampling system of the sidestream CO_2 module does not work correctly, check to see if the sampling line is kinked. If the sampling line is not kinked, remove it from the watertrap. If the monitor gives a message indicating the sampling system still does not work correctly, it indicates that the watertrap is blocked and should be replaced with a new one. Otherwise, the sampling line is blocked and should be replaced with a new one.

16.10 Removing Exhaust Gases from the System

WARNING

 Anesthetics: When using the Sidestream or Microstream CO₂ measurement on patients who are receiving or have recently received anesthetics, connect the outlet to a scavenging system to avoid exposing medical staff to anesthetics.

To remove the sample gas to a scavenging system, connect an exhaust tube to the gas outlet connector of the module.

16.11 Zeroing the Sensor

Zero calibration eliminates the effect of baseline drift during CO_2 measurement exerted on the readings and therefore maintains the accuracy of the CO_2 measurements.

For sidestream and microstream CO_2 modules, a zero calibration is carried out automatically when necessary. You can also start a manual zero calibration if necessary.

To manually start a zero calibration:

- Select [Main Menu]→[Maintenance >>]→[User Maintenance >>]→enter the required password→[Maintain CO2 >>].
- 2. Select [Calibrate CO2 >>]→[Start Zero Cal.].

Disconnecting the patient airway is not required when performing a zero calibration.

WARNING

• Do not rely on the readings during zeroing.

16.12 Calibrating the Sensor

For sidestream or microstream CO₂ modules, a calibration should be performed once every year or when the readings go far beyond the range. For details, refer to the chapter **26** User Maintenance.

16.13 Oridion Information

Microstream

Oridion Patents

This posting serves as notice under 35 U.S.C.§287(a) for Covidien patents: http://www.covidien.com/patents.

No Implied License

Possession or purchase of this device does not convey any express or implied license to use the device with unauthorized CO_2 sampling consumables which would, alone, or in combination with this device, fall within the scope of one or more of the patents relating to this device and/or CO_2 sampling consumable.

17.1 Introduction

The anasthetic gas (AG) module measures the patient's anesthetic and respiratory gases by connecting to the airway of intubated patients or collecting the gases with specified accessories. It also incorporates the features of the O₂ module and BIS module. The AG measurement is applicable for adult, pediatric and neonatal patients. AG monitoring is for cPM 12 patient monitors only.

The AG module determines the concentration of certain gases using the infrared (IR) light absorption measurement. The gases that can be measured by the AG module absorb IR light. Each gas has its own absorption characteristic. The gas is transported into a sample cell, and an optical IR filter selects a specific band of IR light to pass through the gas. For multiple gas measurements, there are multiple IR filters. The higher the concentration of gas in a given volume the more IR light is absorbed. This means that higher concentration of IR absorbing gas cause a lower transmission of IR light. The amount of IR light transmitted after it has been passed through an IR absorbing gas is measured. From the amount of IR light measured, the concentration of gas present can be calculated.

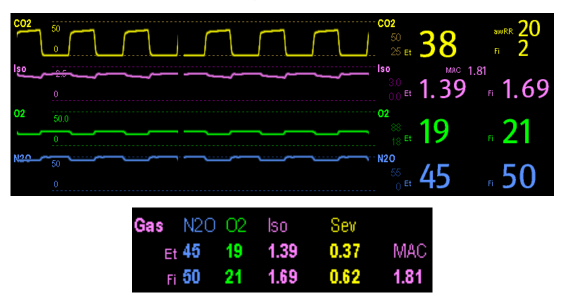
Oxygen does not absorb IR light as other breathing gases and is therefore measured relying on its paramagnetic properties. Inside the O_2 sensor are two nitrogen-filled glass spheres mounted on a strong rare metal taut-band suspension. This assembly is suspended in a symmetrical non-uniform magnetic field. In the presence of paramagnetic oxygen, the glass spheres are pushed further away from the strongest part of the magnetic field. The strength of the torque acting on the suspension is proportional to the oxygen concentration. From the strength of the torque, the concentration of oxygen is calculated.

AG module can identify two anesthetic gases in a mixture automatically and distinguish between them according to their contributions to the MAC value for display as the primary and secondary anesthetic agents.

NOTE

- The AG module is configured with automatic barometric pressure compensation function.
- Perform the measurement in a well-ventilated environment.

17.2 Understanding the AG Display



The AG module can send waves and numerics for all measured anesthetic gases for display on the monitor, including:

- CO₂, O₂, N₂O and AA waves
- awRR: airway respiratory rate
- MAC: minimum alveolar concentration
- End tidal (Et) and fraction of inspired (Fi) numerics for CO₂, O₂, N₂O and AA

Where AA represents Des (desflurane), Iso (isoflurane), Enf (enflurane), Sev (sevoflurane), or Hal (halothane). The AA waveform area displays the primary anesthetic gas's waveform. The O₂ waveform will be displayed only when the O₂ waveform is currently switched on.

WARNING

• To avoid explosion hazard, do not use flammable anesthetic agent such as ether and cyclopropane with this equipment.

17.3 MAC Values

Minimum alveolar concentration (MAC) is the minimum concentration of the agent in the alveoli. It is a basic index to indicate the depth of anesthesia. The standard ISO 21647 defines MAC as this: alveolar concentration of an inhaled anesthetic agent that, in the absence of other anesthetic agents and at equilibrium, prevents 50% of patients from moving in response to a standard surgical stimulus.

Minimum alveolar concentration (MAC) values are listed below:

Agent	Des	lso	Enf	Sev	Hal	N2O
1 MAC	6%*	1.15%	1.7%	2.1%	0.77%	105%**

* The data is taken from a 25-year-old patient.

** indicates 1 MAC nitrous oxide can only be reached in hyperbaric chamber.

NOTE

- The MAC values shown in the table above are those published by the U.S. Food and Drug Administration for a healthy 40-year-old adult male patient.
- In actual applications, the MAC value may be affected by age, weight and other factors.

The formula to calculate the MAC value is as follows:

$$MAC = \sum_{i=0}^{N-1} \frac{EtAgent_i}{AgentVol_{age}i}$$

Where N is the number of all agents (including N_2O) that the AG module can measure, EtAgenti is the concentration of each agent, and AgentVol_{ace}i is the concentration of each agent at 1 MAC with age correction.

The formula for calculating age correction of 1 MAC is:

$$\textit{MAC}_{age} = \textit{MAC}_{40} \times 10^{(-0.00269 \times (age-40))}$$

For example, the Des concentration at 1 MAC of a 60-year old patient is $6\% \times 10^{(-0.00269 \times (60-40))} = 6\% \times 0.88$.

The AG module measures there are 4% of Des, 0.5% of Hal and 50% of N_2O in the patient's end-tidal gas:

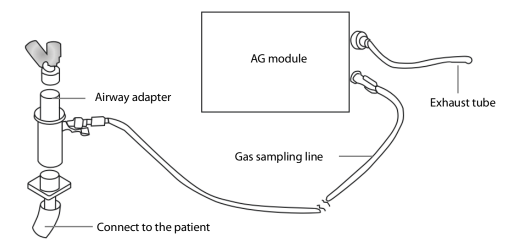
$$MAC = \frac{4.0\%}{6\% \times 0.88} + \frac{0.5\%}{0.77\% \times 0.88} + \frac{50\%}{105\% \times 0.88} = 2.04$$

NOTE

• The formula above is only suitable for patients who are older than one year. If the patient is less than one year, the system uses one year to do age correction.

17.4 Preparing to Measure AG

- 1. Select an appropriate watertrap according to patient category and attach it to the module.
- 2. Connect the gas sample line to the watertrap connector.
- 3. Connect the other end of the gas sampling line to the patient via the airway adapter.
- 4. Connect the gas outlet to a scavenging system using an exhaust tube.



5. Insert the AG module to the monitor and the monitor will prompt [**AG Startup**]. Then the AG module starts to warmup and at the same time the monitor prompts [**AG Warmup**]. After 45 seconds, the AG module enters the iso accuracy mode. After 10 minutes, the module enters the full accuracy mode.

WARNING

- Perform the AG leakage test every time before the AG measurement. For instructions on how to perform the leakage test, see26.7.1AG Leakage Test.
- Make sure that the connections are tight. Any leak in the system can result in erroneous readings due to ambient air mixing with patient gases.
- Do not apply adult watertrap to the neonate patient. Otherwise, patient injury could result.
- Using high-frequency electrosurgical units may increase the risk of skin burn. In this case, do not use antistatic or conductive respiratory tubing.
- Check that the alarm limit settings are appropriate before taking measurement.

CAUTION

- Position the airway adapter so that the part connecting to the gas sample line is pointing upwards. This prevents condensed water from passing into the gas sample line and causing an occlusion.
- The watertrap collects water drops condensed in the sampling line and therefore prevents them from entering the module. To avoid blocking the airway, empty the watertrap container whenever half full. Dispose of accumulated fluids in accordance with the hospital policy or your local regulations.
- The watertrap has a filter preventing bacterium, water and secretions from entering the module. After long-term use, dust or other substances may compromise the performance of the filter or even block the airway. In this case, replace the watertrap. Replacing the watertrap once a month is recommended.

17.5 Changing AG Settings

17.5.1 Setting Gas Unit

For N2O and AA, the unit of the measured gas is fixed to 1%

To set the gas unit:

- 1. Select [Main Menu] \rightarrow [Maintenance >>] \rightarrow [User Maintenance] \rightarrow enter the required password \rightarrow [Unit Setup >>].
- 2. In the Unit Setup menu, you can select [CO2 Unit] or [O2 Unit] and then select [mmHg], [%] or [kPa].

17.5.2 Setting the Apnea Alarm Delay

To set the apnea alarm delay:

- 1. Select the AG parameter area to access the [AG Setup] menu.
- 2. Select [Apnea Delay] and select the appropriate setting.

The monitor alarms if the patient has stopped breathing for longer than the preset apnea time. The [**Apnea Delay**] setting is universally applied to the Resp, CO₂, and AG modules.

WARNING

• The respiration measurement does not recognize the cause of apneas. It only indicates an alarm if no breath is detected when a preset time has elapsed since the last detected breath. Therefore, it cannot be used for diagnostic purposes.

17.5.3 Changing the Sample Flow Rate

In the setup menu for any gas, select [Flow Rate] and then choose either:

- [High]: 200 ml/min for adult and pediatric patients, and 120 ml/min for neonatal patients.
- [Med]: 150 ml/min for adult and pediatric patients, and 90 ml/min for neonatal patients.
- **Low**]: 120 ml/min for adult and pediatric patients, and 70 ml/min for neonatal patients.

17.5.4 Entering the Standby Mode

For the AG module, the default operating mode is measure. When you set the AG module to the Standby mode, the AG gas sample intake pump automatically sets the sample flow rate to zero. When exiting the Standby mode, the AG module continues to work at the previously set sample flow rate with no need to warm up again. After nearly 1 minute, the module enters the full accuracy mode. The AG module Standby mode relates to the Standby mode of the monitor as follows:

- If the monitor enters the Standby mode, the AG module also enters the Standby mode.
- If the monitor exits the Standby mode, the AG module also exits the Standby mode.
- If the AG module enters or exits the Standby mode, it does not affect the monitor.

To enter or exit the Standby mode manually, in the agent's setup menu, select [**Operating Mode**] and then select [**Standby**] or [**Measure**]. You can also set a period of time after which the AG module enters the Standby mode automatically if no breath is detected since the last detected breath. To set the Standby time, in the agent's setup menu, select [**Auto Standby (min**)] and then select the appropriate setting.

17.5.5 Setting up the AG Wave

Select the AG parameter area to access the [AG Setup] menu, in which you can:

- Select [CO2 Wave Type] and then select [Draw] or [Fill]:
 - [**Draw**]: The CO_2 wave is displayed as a curved line.
 - [Fill]: The CO₂ wave is displayed as a filled area.
- Select [Sweep] and then select the appropriate setting. The faster the wave sweeps, the wider the wave is.
- Change the size of the waveform by adjusting the scale.

17.5.6 Setting RR Source

To set RR source:

- 1. Select the AG parameter area to access the [AG Setup] menu.
- 2. Select [RR Source] and then select a source or [Auto] from the dropdown list.

The [**RR Source**] setting is universally applied to the Resp, CO₂, and AG modules. For details, please refer to the section *9.9 Setting Respiration Rate Source*.

17.6 Changing the Anesthetic Agent

When the anesthetic agent used on the patient is changed, the AG module detects the mixed anesthetic gas during the transition of two anesthetic agents. The time required for completing the replacement of anesthetic agent depends on anesthesia type (low flow or high flow) and the characteristics of anesthetic agents (pharmacokinetics).

The AG module can identify two anesthetic agents automatically. When the proportion of the primary and secondary anesthetic agents in the mixture changes, the AG module can distinguish between them according to their contributions to the MAC value. Then the primary and secondary anesthetic agents will be exchanged for display.

CAUTION

• During the transition of two anesthetic agents, the monitor gives no prompt messages and the MAC value displayed may be inaccurate.

17.7 Measurement Limitations

The following factors may influence the measurement accuracy:

- Leaks or internal venting of sampled gas
- Mechanical shock
- Cyclic pressure up to 10 kPa (100 cmH₂O)
- Other sources of interference, if any

17.8 Troubleshooting

17.8.1 When the Gas Inlet is Blocked

If the gas inlet (including watertrap, sampling line and airway adapter) is occluded by condensed water, the message [AG Airway Occluded] appears.

To remove the occlusion:

- Check the airway adapter for an occlusion and replace if necessary.
- Check the sampling line for an occlusion or kinking and replace if necessary.
- Check the watertrap for a build up of water. Empty the watertrap. If the problem persists, replace the watertrap.

17.8.2 When an Internal Occlusion Occurs

Condensed water may enter the module and cause contamination and/or internal occlusions. In this case, the message [AG Airway Occluded] displays.

To remove the occlusion:

- Check for any occlusion in the gas inlet and/or outlet system.
- If the problem persists, internal occlusions may exist. Contact your service personnel.

17.9 Removing Exhaust Gases from the System

WARNING

• Anesthetics: When using the AG measurement on patients who are receiving or have recently received anesthetics, connect the outlet to a scavenging system to avoid exposing medical staff to anesthetics.

To remove the sample gas to a scavenging system, connect an exhaust tube to the gas outlet connector of the module.

FOR YOUR NOTES

During patient monitoring, the freeze feature allows you to freeze the currently displayed waveforms on the screen so that you can have a close examination of the patient's status. Additionally, you can select any frozen waveform for recording.

18.1 Freezing Waveforms

To freeze waveforms, select the [Freeze] hardkey on the monitor's front panel.

■ The system closes the displayed menu (if any), and opens the [Freeze] menu.

Freeze					×
Wave 1	II	Wave 2	I	Wave 3	Pleth
		Scroll		Record	

■ All displayed waveforms are frozen, i.e. the waveforms stop being refreshed.

NOTE

• The freeze feature does not affect the split-screen view of minitrends, OxyCRG and other patients.

18.2 Viewing Frozen Waveforms

To view the frozen waveforms, you can either:

- Select the [Scroll] button and then rotate the knob clockwise or counter-clockwise, or
- Directly select the ◀ or ▶ beside the [Scroll] button using the touchscreen.

The frozen waveforms will scroll left or right accordingly. And meanwhile, at the lower right corner of the last waveform, there is an upward arrow. The freeze time displays below the arrow and the initial frozen time is [**0** s]. With the waveforms scrolling, the freeze time changes at intervals of 1 second. This change will be applied for all waveforms on the screen.

18.3 Unfreezing Waveforms

To unfreeze the frozen waveforms, you can either:

- Select the X button at the upper right corner of the [Freeze] menu,
- Select the [Freeze] hardkey on the monitor's front panel, or
- Perform any other action that causes the screen to be readjusted or opens a menu, such as inserting or removing a module, pressing the [Menu] hardkey, etc.

18.4 Recording Frozen Waveforms

To record the frozen waveforms:

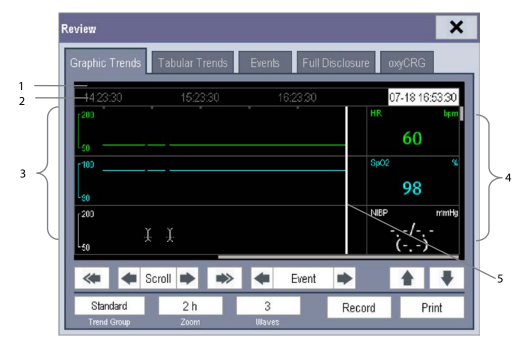
- 1. In the [Freeze] menu, select, in order, [Wave 1], [Wave 2] and [Wave 3] and then select your desired waveforms.
- 2. Select the [**Record**] button. The selected waveforms and all numerics at the frozen time are printed out by the recorder.

19.1 Accessing Respective Review Windows

- 1. Select the [**Review**] QuickKey, or [**Main Menu**]→[**Review** >>].
- 2. Select [Graphic Trends], [Tabular Trends], [Events], or [Full Disclosure] to access their respective review windows.

19.2 Reviewing Graphic Trends

In the [Review] menu, select [Graphic Trends] to access the following window.



1.Event mark area2. Time axis3. Graphic trends area4.Parameter area5. Cursor

i. Fuldificter dicu

Events are marked with colors in the event mark area. Red represents high level alarm event. Yellow represents medium/low level alarm event. Green represents manual event.

In this review window:

- Select [Trend Group] and you can select a trend group for viewing in the popup menu. If [Custom 1] or [Custom 2] is selected, you can further select [Define Trend Group]. Then you can select the parameters for viewing in the popup menu.
- Select [**Zoom**] to set the time length of the review window.
- Select [Waves] to set the number of waves displayed in one page.

- To browse the graphic trends, you can either:
 - ◆ Select ≤ or ▶ beside [Scroll] to move the cursor one step to the left or right to navigate through the graphic trends, or
 - Select select select to move the cursor one page to the left or right to navigate through the graphic trends.

A time indicating your current position displays above the parameter area. Numeric measurement values corresponding to the cursor location change as the cursor is moved. The measurement value that triggered a high level alarm has a red background. The one that triggered a medium/low level alarm has a yellow background.

- Select select
- Select the [**Record**] button to print the currently displayed graphic trends by the recorder.
- Select the [Print] button to set and print the graphic trends report by the printer. For how to set the graphic trends report, refer to chapter 22 Printing.

19.3 Reviewing Tabular Trends

In the [Review] menu, select [Tabular Trends] to access the following window.

eview						×
Graphic Trends	Tabular	Trends	Events	Full Disclos	ure oxy(CRG
08-24	14:10:00	14:10:30	14:11:00	14:11:30	14:12:00	14:12:30
HR	60	60	60	60	60	60
SpO2/SpO2b	98/92	98/92	98/92	98/92	98/92	98/92
∆SpO2	6	6	6	6	6	6
NIBP	 ()	/ ()	 ()	 ()	 ()	l ()
RR	20	20	20	20	20	20
-	Scroll 🗭		🗲 Ev	ent 🕨	1	h 👎
Standard Trend Group	30 s Interv		Record (All) Re	cord	Print

Events are marked with colors in window's top area. Red represents high level alarm event. Yellow represents medium/low level alarm event. Green represents manual event.

In this review window:

- Select [Trend Group] and you can select a trend group for viewing in the popup menu. If [Custom 1] or [Custom 2] is selected, you can further select [Define Trend Group]. Then you can select the parameters for viewing in the popup menu.
- Select [Interval] to change the resolution of the trend data and then select the appropriate setting:
 - [5 s] or [30 s]: select to view up to 4 hours of tabular trends at 5- or 30-second resolution.
 - [1 min], [5 min], [10 min], [15 min], [30 min], [1 h], [2 h] or [3 h]: select to view up to 120 hours of tabular trends at your selected resolution.
 - [NIBP]: select to view the tabular trends when NIBP measurements were acquired.
- To browse the tabular trends, you can either:
 - ◆ Select < or 🖻 beside [Scroll] to drag the scrollbar left or right to navigate through the trend database, or
 - ◆ Select ≪ or 💌 to scroll left or right to navigate through the trend database.

The measurement value that triggered a high level alarm has a red background. The one that triggered a medium/low level alarm has a yellow background.

- Select select
- Select the [Record] button to access the [Record Setup] menu and set the start and end time of the tabular trends you want to record. This feature is not available when reviewing a patient's history. By further selecting [Record], you can print the displayed tabular trends to the recorder.
- Select the [Record (AII)] button to print all of the tabular trends to the recorder. The trends will be printed in reverse chronological order from the current time to the last time a patient was admitted to the patient monitor.
- Select the [Print] button to set and print the tabular trends report with the printer. For how to configure the tabular trends report, refer to chapter 22 Printing.

19.4 Reviewing Events

19.4.1 Marking Events

During monitoring, some events may affect the patient and change the displayed waveforms or numerics displayed on the monitor. To help analyze these waveforms or numeric changes, you can mark these events.

Select [Main Menu] \rightarrow [Mark Event >>]. In the popup menu, you can select the waves to store after triggering a manual event. You can select [Trigger Manual Event] from the [Mark Event] menu or the [Manual Event] QuickKey to trigger a manual event and store it at the same time.

When you are reviewing graphic trends, tabular trends or full-disclosure waveforms, the manual event symbol is displayed at the time the event is triggered.

19.4.2 Reviewing Events

The monitor saves the events in real time. You can review these events.

In the [Review] menu, select [Events] to access the following window.

You can review the parameter alarm events, arrhythmia alarm events and manual events. When an event occurs, the system stores all the measurement numerics at the event triggering time and related waveforms 4 seconds, 8 seconds, or 16 seconds before and after the event triggering time, as per the setting of recording length.

NOTE

- Pausing or switching off alarms will not be recorded as events. The time of these operations will not be recorded in the system log.
- Earlier-recorded events might be overwritten by later ones if the events log reaches capacity.

R	eview					×
	Graphic Trends	Tabular Tren	ds Events	Full Disclosure	oxyCRG	
	Tim	е		Event		
	2016-08-24	14:04:42		** HR Too Low < 61		
	Details		Scroll 🖶 🤅	All	All	-1
L				▼ Event	Level	

In this window:

- Select [**Event**] to view the desired events.
- Selecting [Level] to view the desired events according to the alarm priority .

After selecting the desired event, you can select [**Details**] to access the following window. In this window, the waveform area displays the waveforms related to the event, and the parameter area displays the parameter values happened at the event trigger time.

NOTE

• A total loss of power has no impact on the saved events.



- 1. Waveform area
- 2. Parameter area

In this window:

- You can select or beside the [Event] button to switch between events.
- By selecting the [Beat Annotation] button, you can display or hide beat labels above the first ECG waveform on the screen. The white beat labels indiate how the monitor is classifying beats and may help explain suspected, missed, or false arrhythmia calls.
 - ♦ N = Normal
 - V = Ventricular Ectopic
 - S = Supraventricular Premature
 - P = Paced
 - L = Learning
 - ? = Insufficient information to classify beats
 - ◆ I = Inoperative (for example, Lead Off)
 - M = No beat detected

When beat labels are displayed above the ECG waveform, they can be printed via the recorder or a laser printer. If you select [**Beat Annotation**] in the [**Events**] tab to display or hide beat labels, these beat labels will also be displayed or hidden in the [**Full Disclosure**] tab, and vise versa.

CAUTION

• Ensure that you have selected the optimal leads with the best waveform amplitude and the highest signal-to-noise ratio. Selecting the optimal leads is important for beat detection, beat classification, and V-Fib detection.

- You can set the desired [**Gain**] for ECG waveform.
- You can set the desired [**Sweep**].
- By selecting the [**Record**] button, you can print the displayed alarm events with the recorder.
- Select [**Print**] to print the displayed alarm event to the pre-connected laser printer.
- Select [**Events List**] to view the events list.

19.5 Reviewing Waveforms

In the [Review] menu, select [Full Disclosure] to access the following window.



A. Waveform area

B. Parameter area

In this review window:

- To review full-disclosure waveforms, you need to save waveforms first. Select [Save Waves >>] and then select the parameters whose waveforms you want to view. To save full-disclosure waveforms, your monitor must have a storage card.
- To view the waveforms, you can either:
 - Select sel
 - Select select select is to move the cursor one page left or right to navigate through the waveforms.

A time indicating your current position is displayed at the top of the waveform area. Numeric measurement values corresponding to the cursor location are displayed in the parameter area, and change as the cursor is moved.

- By selecting the [Beat Annotation] button, you can display or hide beat labels above the first ECG waveform on the screen. For more information on the beat annotation function, refer to the section 19.4.2 Reviewing Events.
- You can change the ECG wave gain by selecting [Gain] and then selecting the appropriate setting.
- You can change the waveform sweep speed by selecting [**Sweep**] and then selecting the appropriate setting.
- By selecting the [Record] button, you can print out the first three waveforms and measurement numerics by the recorder.
- By selecting selecting selecting beside the [Event] button, you can position the cursor between events.

20.1 Introduction

The calculation feature is available with your monitor. The calculated values, which are not directly measured, are computed based on the values you provide.

Your can perform the following calculations:

- Dose calculations
- Oxygenation calculations
- Ventilation calculations
- Hemodynamic calculations
- Renal calculations

To perform a calculation, select [**Main Menu**] \rightarrow [**Calc** >>], or the [**Calculations**] QuickKey and then select the calculation you want to perform.

WARNING

• After the calculation is finished, verify the entered values are correct and the calculated values are appropriate. We assume no responsibility for any consequences caused by wrong entries and improper operations.

NOTE

• The calculation feature is independent of other monitoring functions and can be therefore used for patients being monitored by other monitors. Any operation in a calculation window does not affect the patient monitoring by the local monitor.

20.2 Dose Calculations

20.2.1 Performing Calculations

To perform a dose calculation:

- 1. Select [Main Menu]→[Calc >>]→[Dose >>], or select [Calculations] QuickKey→[Dose >>].
- 2. Select [Patient Cat.] and [Drug Name] and then select the appropriate settings. The dose calculation program has a library of commonly used drugs, of which Drug A through Drug E are for those not specified in this library.
 - Drug A, B, C, D, E Isuprel
 - Aminophylline

- Lidocaine

- Dobutamine Dopamine
- Epinephrine
- Heparin
- 3. The system gives a set of default values when the above steps are finished. However, these values cannot be used as the calculated values. The user must enter values following the doctor's instructions, and only then can the calculated values be used
- 4. Enter the patient's weight.
- 5. Enter other values.
- 6. Verify if the calculated values are correct.

20.2.2 Selecting the Proper Drug Unit

Each drug has its fixed unit or unit series. Among a unit series, one unit may change to another automatically depending on the entered value.

The units for each drug are as follows:

- Drug A, B, C, Aminophylline, Dobutamine, Dopamine, Epinephrine, Isuprel, Lidocaine, Nipride and Nitroglycerin use the unit series: g, mg and mcg.
- Drug D, Heparin and Pitocin use the unit series: Unit, KU (kilo units) and MU (million units).
- Drug E uses the unit: mEq (milli-equivalents).

You must select the proper drug name (A, B, C, D or E) according to the units when you define a drug not listed in this library.

NOTE

For neonate patients, [Drip Rate] and [Drop Size] are disabled. •

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- Nltroglycerin
- Pitocin

- Nipride

20.2.3 Titration Table

To open the titration table, select [**Titration Table** >>] in the [**Dose Calculation**] window after the dose calculation is finished.

In the titration table, when you change:

- [Reference]
- [Interval]
- [Dose Type]

The titrated values change accordingly.

You can also:

- Select \$\Rightarrow\$ or \$\Figs\$, or \$\Delta\$ or \$\Delta\$ beside the vertical scrollbar to view more values.
- Select [Record] to print the displayed titrated values by the recorder.

20.2.4 Drug Calculation Formulas

Abbreviation	Unit	Formula
Conc.	g/ml, unit/ml or mEq/ml	Amount / Volume
Dose	Dose/hr, Dose/kg/min	Rate × Conc.
Volume	ml	Rate × Duration
Amount	g, unit, mEq	Rate × Duration
Duration	h	Amount/Dose
Drip Rate	gtt/min	INF Rate × Drop Size / 60

20.3 Oxygenation Calculations

20.3.1 Performing Calculations

To perform an oxygenation calculation:

- 1. Select [Main Menu] \rightarrow [Calc >>] \rightarrow [Oxygenation >>], or select [Calculations] QuickKey \rightarrow [Oxygenation >>].
- 2. Enter values for calculation.
- 3. Select the [**Calculate**] button. The system performs a calculation per the current settings and displays the calculated values.
 - If a calculated value is outside the range, its background will highlight in yellow. You can select [Range] to view its normal range in the unit field.
 - Invalid values are displayed as [---].

In the [Oxygenation Calculation] window, you can:

- Change the pressure unit, Hb unit and oxygen content unit by selecting [Press. Unit], [Hb Unit] and [OxyCont
 Unit] and then selecting the appropriate settings. The changes take effect automatically.
- Trigger a recording by selecting the [Record] button. The displayed oxygenation calculations are printed out by the recorder.
- Review the previously performed calculations by selecting [**Review**].

Abbreviation	Unit	Full spelling	
C.O.	L/min	cardiac output	
FiO ₂	%	percentage fraction of inspired oxygen	
PaO ₂	mmHg	partial pressure of oxygen in the arteries	
PaCO ₂	mmHg	partial pressure of carbon dioxide in the arteries	
SaO ₂	%	arterial oxygen saturation	
PvO ₂	mmHg	partial pressure of oxygen in venous blood	
SvO ₂	%	venous oxygen saturation	
Hb	g/L	hemoglobin	
CaO ₂	ml/L	arterial oxygen content	
CvO ₂	ml/L	venous oxygen content	
VO ₂	ml/min	oxygen consumption	
RQ	None	respiratory quotient	
ATMP	mmHg	atmospheric pressure	
Height	cm	height	
Weight	kg	weight	

20.3.2 Entered Parameters

20.3.3 Calculated Parameters and Formulas

Abbreviation	Unit	Full spelling	Formula
BSA	m ²	body surface area	Wt $^{0.425}$ × Ht $^{0.725}$ × 0.007184
VO ₂ calc	ml/min	oxygen consumption	$C(a-v)O_2 \times C.O.$
C(a-v)O ₂	ml/L	arteriovenous oxygen content difference	$CaO_2 - CvO_2$
O ₂ ER	%	oxygen extraction ratio	100×C(a-v)O ₂ / CaO ₂
DO ₂	ml/min	oxygen transport	$C.O. \times CaO_2$
DAO	mmHa	partial pressure of oxygen in the alveoli	$FiO_2 / 100 \times (ATMP - 47) - PaCO_2 \times [FiO_2 / 100$
PAO ₂	mmHg		$+ (1 - FiO_2 / 100) / RQ]$
AaDO ₂	mmHg	alveolar-arterial oxygen difference	$PAO_2 - PaO_2$
CcO ₂	ml/L	capillary oxygen content	$Hb \times 1.34 + 0.031 \times PAO_2$
		venous admixture	$100 \times [1.34 \times Hb \times (1 - SaO_2 / 100) + 0.031 \times$
Qs/Qt	%		$(PAO_2 - PaO_2)] / [1.34 \times Hb \times (1 - SvO_2 / 100)$
			$+ 0.031 \times (PAO_2 - PvO_2)]$
C.O. calc	L/min	calculated cardiac output	$VO_2/(CaO_2 - CvO_2)$

20.4 Ventilation Calculations

20.4.1 Performing Calculations

To perform a ventilation calculation:

- 1. Select [Main Menu] \rightarrow [Calc >>] \rightarrow [Ventilation >>], or select [Calculations] QuickKey \rightarrow [Ventilation >>].
- 2. Enter values for calculation. If the monitor is connected to an anesthesia machine or a ventilator, the system automatically loads the supported parameter values to the [Ventilation Calculation] window.
- 3. Select the [**Calculate**] button. The system performs a calculation per the current settings and displays the calculated values.
 - If a calculated value is outside the range, its background will highlight in yellow. You can select [**Range**] to view its normal range in the unit field.
 - Invalid values are displayed as [---].

In the [Ventilation Calculation] window, you can:

- Change the pressure unit by selecting [Press. Unit] and then selecting the appropriate setting. Corresponding pressure values shall convert and update automatically.
- Trigger a recording by selecting the [Record] button. The displayed ventilation calculations are printed out by the recorder.
- Review the previously performed calculations by selecting [**Review**].

Abbreviation	Unit	Full spelling	
FiO ₂	%	percentage fraction of inspired oxygen	
RR	rpm	respiration rate	
PeCO ₂	mmHg	partial pressure of mixed expiratory CO ₂	
PaCO ₂	mmHg	partial pressure of carbon dioxide in the arteries	
PaO ₂	mmHg	partial pressure of oxygen in the arteries	
TV	ml	tidal volume	
RQ	None	respiratory quotient	
ATMP	mmHg	atmospheric pressure	

20.4.2 Entered Parameters

20.4.3 Calculated Parameters and Formulas

Abbreviation	Unit	Full spelling	Formula
PAO ₂	mmHa	partial pressure of oxygen in the alveoli	$(ATMP-47) \times FiO_2/100 - PaCO_2 \times$
PAO ₂	mmHg	partial pressure of oxygen in the alveoli	$[FiO_2/100 + (1-FiO_2/100)/RQ]$
AaDO ₂	mmHg	alveolar-arterial oxygen difference	$PAO_2 - PaO_2$
Pa/FiO ₂	mmHg	oxygenation ratio	$100 \times PaO_2 / FiO_2$
a/AO ₂	%	arterial to alveolar oxygen ratio	$100 \times PaO_2 / PAO_2$
MV	L/min	minute volume	(TV × RR) / 1000
Vd	ml	volume of physiological dead space	$TV \times (1 - PeCO_2 / PaCO_2)$
Vd/Vt	%	physiologic dead space in percent of tidal volume	100 × Vd/TV
VA	L/min	alveolar volume	$(TV - Vd) \times RR / 1000$

20.5 Hemodynamic Calculations

20.5.1 Performing Calculations

To perform a hemodynamic calculation:

- 1. Select [Main Menu]→[Calc>>]→[Hemodynamic>>], or select [Calculations] QuickKey→[Hemodynamic>>].
- 2. Enter values for calculation.
 - For a monitored patient, [HR], [Art mean], [PA mean] and [CVP] are automatically taken from the currently measured values. If you have just performed C.O. measurements, [C.O.] is the average of multiple thermodilution measurements. [Height] and [Weight] are the patient's height and weight you have entered. If the monitor does not provide these values, their fields are blank.
 - For an unmonitored patient, confirm the entered values.
- 3. Select the [**Calculate**] button. The system performs a calculation per the current settings and displays the calculated values.
 - If a calculated value is outside the range, its background will highlight in yellow. You can select [**Range**] to view its normal range in the unit field.
 - Invalid values are displayed as [---].

In the [Hemodynamic Calculation] window, you can:

- Trigger a recording by selecting the [Record] button. The currently displayed hemodynamic calculations are printed out by the recorder.
- Review the previously performed calculations by selecting [**Review**].

Unit	Full spelling		
L/min	cardiac output		
bpm	heart rate		
mmHg	pulmonary artery wedge pressure		
mmHg	artery mean pressure		
mmHg	pulmonary artery mean pressure		
mmHg	central venous pressure		
ml	end-diastolic volume		
cm	height		
kg	weight		
	Unit L/min bpm mmHg mmHg mmHg mmHg ml cm		

20.5.2 Entered Parameters

20.5.3 Calculated Parameters and Formulas

Abbreviation	Unit	Full spelling	Formula
BSA	m²	body surface area	Wt ^{0.425} × Ht ^{0.725} × 0.007184
C.I.	L/min/m ²	cardiac index	C.O. / BSA
SV	ml	stroke volume	C.O. / HR × 1000
SI	ml/m ²	stroke index	SV/ BSA

Abbreviation	Unit	Full spelling	Formula
SVR	DS/cm⁵	systemic vascular resistance	79.96 × (AP MAP – CVP) / C.O.
SVRI	DS⋅m²/cm ⁵	systemic vascular resistance index	SVR × BSA
PVR	DS/cm⁵	pulmonary vascular resistance	79.96 × (PAMAP – PAWP) / C.O.
PVRI	DS⋅m²/cm ⁵	pulmonary vascular resistance index	PVR × BSA
LCW	kg∙m	left cardiac work	0.0136 × APMAP × C.O.
LCWI	kg·m/m²	left cardiac work index	LCW / BSA
LVSW	g∙m	left ventricular stroke work	$0.0136 \times APMAP \times SV$
LVSWI	g·m/m²	left ventricular stroke work index	LVSW / BSA
RCW	kg∙m	right cardiac work	0.0136 × PAMAP × C.O.
RCWI	kg·m/m ²	right cardiac work index	RCW / BSA
RVSW	g∙m	right ventricular stroke work	0.0136 × PAMAP × SV
RVSWI	g·m/m²	right ventricular stroke work index	RVSW / BSA
EF	%	ejection fraction	$100 \times SV / EDV$

20.6 Renal Calculations

20.6.1 Performing Calculations

To perform a renal calculation:

- 1. Select [Main Menu] \rightarrow [Calc >>] \rightarrow [Renal >>], or select [Calculations] QuickKey \rightarrow [Renal >>].
- 2. Enter values for calculation.
- 3. Select the [**Calculate**] button. The system performs a calculation per the current settings and displays the calculated values.
 - If a calculated value is outside the range, its background will highlight in yellow. You can select [**Range**] to view its normal range in the unit field.
 - Invalid values are displayed as [---].

In the [Renal Calculation] window, you can:

- Trigger a recording by selecting the [Record] button. The displayed renal calculations are printed by the recorder.
- Review the previously performed calculations by selecting [**Review**].

Abbreviation	Unit	Full spelling
URK	mmol/L	urine pstassium
URNa	mmol/L	urinary sodium
Urine	ml/24h	urine
Posm	mOsm/ kgH ₂ O	plasm osmolality
Uosm	mOsm/ kgH ₂ O	urine osmolality
SerNa	mmol/L	serum sodium
Cr	μmol/L	creatinine

20.6.2 Entered Parameters

Abbreviation	Unit	Full spelling
UCr	μmol/L	urine creatinine
BUN	mmol/L	blood urea nitrogen
Height	cm	height
Weight	kg	weight

20.6.3 Calculated Parameters and Formulas

Abbreviation	Unit	Full spelling	Formula
URNaEx	mmol/24h	urine sodium excretion	Urine × URNa / 1000
URKEx	mmol/24h	urine potassium excretion	Urine × URK / 1000
Na/K	%	sodium potassium ratio	100 × URNa / URK
CNa	ml/24h	clearance of sodium	URNa \times Urine / SerNa
Clcr	ml/min	creatinine clearance rate	Ucr × Urine / Cr / (BSA / 1.73) / 1440
FENa	%	fractional excretion of sodium	$100 \times (URNa \times Cr) / (SerNa \times Ucr)$
Cosm	ml/min	osmolar clearance	Uosm × Urine / Posm / 1440
CH ₂ O	ml/h	free water clearance	Urine × (1—Uosm / Posm) / 24
U/P osm	None	urine to plasma osmolality ratio	Uosm / Posm
BUN/Cr	None*	blood urea nitrogen creatinine ratio	1000 × BUN / Cr
U/Cr	None	urine-serum creatinine ratio	Ucr / Cr

*: BUN/Cr is a ratio under the unit of mol.

20.7 Understanding the Review Window

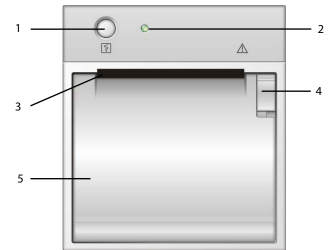
With the review feature, you can review oxygenation, ventilation, hemodynamic and renal calculations. The review window for each calculation is similar. Take the hemodynamic calculations review window for example, you can access it by selecting [**Review**] in the [Hemodynamic Calculation] window.

In this review window:

- You can select ◀, ▶ ◀◀ or ▶▶ to view more values.
- The values that exceed the range are displayed in a yellow background. The [Unit] field displays parameter units. If some parameter values are outside of their normal ranges, you can view their normal range in the [Unit] field by selecting [Range].
- You can review an individual calculation by selecting its corresponding column and then selecting [Original Calc]. You can record the currently displayed calculations or perform another calculation is this window.

21.1 Using a Recorder

The thermal recorder prints patient information, measurement numeric and waveforms, etc.



- 1. Start/Stop key: press to start a recording or stop the current recording.
- 2. Indicator
 - On: when the recorder works correctly.
 - Off: when the monitor is switched off.
 - Flashes: if an error occurred to the recorder, e.g., the recorder runs out of paper.
- 3. Paper outlet
- 4. Latch
- 5. Recorder door

21.2 Overview of Recording Types

By the way recordings are triggered, the recordings can be classified into the following categories:

- Manually-triggered realtime recordings.
- Timed recordings.
- Alarm recordings triggered by an alarm limit violation or an arrhythmia event.
- Manually-triggered, task-related recordings.

NOTE

- For details about alarm recording, refer to the chapter 7 Alarms.
- For details about task-related recordings, refer to respective sections of this manual.

21.3 Starting and Stopping Recordings

To manually start a recording, you can either:

- Press the [**Record**] hardkey on the front of the monitor,
- Press the I hardkey on the front of the recorder module, or
- Select the [**Record**] button from the current menu or window.

Automatic recordings will be triggered in the following conditions:

- Timed recordings will start automatically at preset intervals.
- If both [Alarm] and [Alm Rec] for a measurement are set on, an alarm recording will be triggered automatically as alarms occur.

When a recording is in process, to manually stop the recording, you can either:

- Press the [**Record**] hardkey on the front of the monitor;
- Press the I hardkey on the front of the recorder module; or
- Select [Clear All Tasks] in the [Record Setup] menu.

Recordings stop automatically when:

- A recording is completed.
- The recorder runs out of paper.
- When the recorder has an alarm condition.

When a recording is stopped, the following markers will be added:

- Automatically stopped recording: prints two columns of '*' at the end of the report.
- Manually or abnormally stopped recording: prints one column of '*' at the end of the report.

21.4 Setting up the Recorder

21.4.1 Accessing the Record Setup Menu

By selecting [Main Menu]→[Record Setup >>], you can access the [Record Setup] menu.

21.4.2 Selecting Waveforms for Recording

The recorder can record up to 3 waveforms at a time. You can select, in order, [Waveform 1], [Waveform 2] and [Waveform 3] in the [Record Setup] menu, and then select the waveforms you want. You can also turn off a waveform recording by selecting [Off]. These settings are intended for realtime and scheduled recordings.

21.4.3 Setting the Realtime Recording Length

After starting a realtime recording, the recording time depends on your monitor's settings. In the [Record Setup] menu, select [Length] and then select [8 s] or [Continuous].

- [8 s]: record 4-second waveforms respectively before and after current moment.
- **Continuous**]: record the waveforms from the current moment until stopped manually.

21.4.4 Setting the Interval between Timed Recordings

Timed recordings start automatically at preset intervals. Each recording lasts 8 seconds. To set the interval between timed recordings: in the [**Record Setup**] menu, select [**Interval**] and then select the appropriate setting.

21.4.5 Changing the Recording Speed

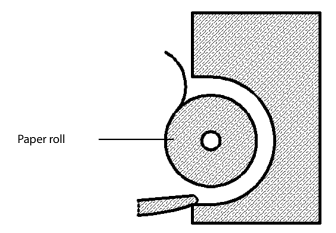
In the [Record Setup] menu, select [Paper Speed] and then select [25 mm/s] or [50 mm/s]. This setting is for all recordings containing waveforms.

21.4.6 Clearing Recording Tasks

In the [Record Setup] menu, select [Clear All Tasks]. All queued recording tasks are cleared and the current recording is stopped.

21.5 Loading Paper

- 1. Press the latch in the upper right corner of the recorder door to open it.
- 2. Insert a new roll into the compartment as shown below.
- 3. Close the recorder door.
- 4. Check if paper is loaded correctly and the paper end is feeding from the top.



CAUTION

- Use only specified thermal paper. Otherwise, it may cause damage to the recorder's printhead, the recorder may be unable to print, or poor print quality may result.
- Never pull the recorder paper with force when a recording is in process. Otherwise, it may cause damage to the recorder.
- Do not leave the recorder door open unless you are reloading paper or resolving an issue.

21.6 Removing Paper Jam

If the recorder works incorrectly or produces unusual sounds, check if there is a paper jam first. If a paper jam is detected, follow this procedure to remove it:

- 1. Open the recorder door.
- 2. Take out the paper and tear off the draped part.
- 3. Reload the paper and close the recorder door.

21.7 Cleaning the Recorder Printhead

If the recorder has been used for a long time, deposits of paper debris may collect on the printhead compromising the print quality and shortening the lifetime of the roller. Follow this procedure to clean the printhead:

- 1. Take measures against the static electricity such as Disposable Wrist Strap for the work.
- 2. Open the recorder door and take out the paper.
- 3. Gently wipe around the printhead using cotton swabs dampened with alcohol.
- 4. After the alcohol has completely dried, reload the paper and close the recorder door.

CAUTION

- Do not use anything that may destroy the thermal element.
- Do not add unnecessary force to the thermal head.

22.1 Printer

The monitor can output patient reports via a connected printer. The monitor supports the following printer:

- HP LaserJet 1505n
- HP LaserJet P2035n
- HP LaserJet P4015n
- HP LaserJet Pro 400 M401n401dn
- HP LaserJet 600 M602
- HP LaserJet Enterprise M605
- HP LaserJet Enterprise M608n

The reports specifications are:

- Paper: A4, Letter
- Resolution: 300 dpi

For more details about the printer, see the document accompanying the printer. With product upgrades, the monitor may support more printers without prior notice will be given. If you have any questions or doubt about your printer, contact our company.

22.2 Connecting a printer

To print the reports or the trend data of a patient, you can connect the local printer and the monitor directly with a network cable.

22.3 Setting Up the Printer

To set the printer's properties, select [Main Menu] \rightarrow [Print Setup >>] \rightarrow [Printer Setup >>]. In the [Printer Setup] menu, you can:

Select a connected printer.

Select [Printer] and then select a connected printer as the monitor's printer.

Search for a printer.

If your selected printer is not in the list or a new printer is added into the network, you can select the [Search Printer] to search for all printers in the network.

Set up the paper.

Select [Paper Size] and and select [A4] or [Letter].

Reports	Contents	Procedures
ECG reports	ECG waveforms and relevant	Select [Main Menu]→[Print Setup >>]→[ECG Reports >>]→
	parameter values	[Print]
OT servert	QT parameter values and waveforms	Select [ECG Setup] \rightarrow [QT Analysis >>] \rightarrow [QT View >>] \rightarrow [Print]
QT report		Note: [QT Analysis] must be set to [On].
Tabular trends	Depends on the selected parameter	Select [Main Menu]→[Print Setup >>]→[Tabular Trends
	group, resolution and time period	Reports >>]→[Print], or select [Main Menu]→[Review >>]→
		[Tabular Trends]→[Print]→[Print]
Graphic trends	Depends on the selected parameter	Select [Main Menu]→[Print Setup >>]→[Graphic Trends
	group, resolution and time period	Reports >>]→[Print], or select [Main Menu]→[Review >>]→
		[Graphic Trends]→[Print]→[Print]
Arrh. alarm	ECG waveforms and relevant	Select [Print] in [Arrh. Events]
review	parameter values	
Parameter alarm	Depends on the selected alarms	Select [Main Menu]→[Review >>]→[Alarms]→[Print]
review		
Realtime waves	Depends on the selected waveforms	Select [Main Menu]→[Print Setup >>]→[Realtime Reports >>]
		→[Print]

22.4 Starting Report Printouts

22.5 Stopping Report Printouts

To stop report printouts, select [Main Menu]→[Print Setup >>]→[Stop All Reports].

22.6 Setting Up Reports

22.6.1 Setting Up ECG Reports

You can print ECG reports only from the full-screen or half-screen. To set up ECG reports, select [Main Menu] \rightarrow [Print Setup >>] \rightarrow [ECG Reports >>].

- [Amplitude]: set the amplitude of the ECG waveforms.
- [Sweep]: set the wave print speed to 25 mm/s or 50 mm/s.
- [Auto Interval]: if [Auto Interval] is set to [On], the system will automatically adjust the space between waveforms to avoid overlapping.
- [Gridlines]: choose whether to show gridlines.

22.6.2 Setting Up Tabular Trends Reports

To set up tabular trends reports, select [Main Menu]→[Print Setup >>]→[Tabular Trends Reports >>].

- [Start time]: you can set a time period whose trend data will be printed out by setting [From] and [Back]. For example, if you set [From] as 2007-4-2 10:00:00 and [Back] as [2 h], the outputted data will be from 2014-4-2 08:00:00 to 2014-4-2 10:00:00. In addition, the [Back] can be set to either:
 - [Auto]: if [ReportFormat] is set to [Time Oriented], the report will be printed by time. If [ReportFormat] is set to [Parameter Oriented], the report will be printed by parameter.
 - [AII]: if you select [AII], all trend data will be printed out. In this case, there is no need to set [From].
- [Interval]: choose the resolution of the tabular trends printed on the report.
- [ReportFormat]: if you select [Time Oriented], the report will be printed by time. If you select [Parameter Oriented], the report will be printed by parameter.
- [Select Parameter >>]: from the pop-up menu, you can:
 - [Currently Displayed Trended Parameters]: print the parameter trend data selected from the [Tabular Trends].
 - [Standard Parameter Group]: select the standard parameter group for printing.
 - [Custom]: define a parameter group for printing from the parameters displayed in the lower part of the menu.

22.6.3 Setting Up Graphic Trends Reports

To set up graphic trends reports, select [Main Menu] \rightarrow [Print Setup >>] \rightarrow [Graphic Trends Reports >>]. Setting up graphic trends reports is similar with tabular trends reports. Refer to section 22.6.2 Setting Up Tabular Trends Reports for details.

22.6.4 Setting Up Realtime Reports

To set up realtime reports, select [Main Menu]→[Print Setup >>]→[Realtime Reports >>].

- [Sweep]: set the wave print speed to 12.5 mm/s, 25 mm/s, 50 mm/s, or Auto.
- [Select Wave >>]: from the pop-up menu, you can:
 - [Current]: print the currently displayed waves.
 - [Select Wave]: select the desired waves for printing.

22.7 End Case Reports

ECG reports, tabular trends reports, graphic trends reports, NIBP review reports and realtime reports can be set as end case reports. When you discharge a patient, the system will automatically print all contents that are set as end case reports.

For example, to set the ECG report as an end case report:

- 1. Select [Main Menu] \rightarrow [Print Setup >>] \rightarrow [ECG Report >>].
- 2. Select [End Case Report]→[Set as End Case Report] and then select [Ok] from the pop-up dialog box.
- 3. Set as described in section 22.6.1 Setting Up ECG Reports.

22.8 Printer Statuses

22.8.1 Printer Out of Paper

When the printer is out of paper, the printer will not print until the paper is replaced. If there are too many print jobs that are not printed, a printer error may occur. In this case, you need to install paper and then re-send the print request. Restart the printer if necessary.

To avoid this, ensure that there is enough paper in the printer before sending a print request.

22.8.2 Printer Status Messages

Printer Status Message	Possible causes and suggested action	
Printer unavailable	The selected printer is not available. Check if the printer is switched on or correctly	
Phinter unavailable	connected or installed with paper.	

23.1 Analog Output

The monitor provides analog output signals to accessory equipment via the multifunctional connector on the rear of the monitor. The analog output signals include analog ECG, IBP, and defibrillator synchronization signals, among which the ECG signals support pace pulses to be enhanced.

NOTE

• The analog output feature is seldom applied in clinical applications. You can contact your service personnel for more details.

23.2 Setting DIAP Baud Rate

The monitor can output measurement numerics and alarm limits of parameter modules at certain baud rate to external devices via the DIAP protocol. The parameter modules are ECG, RR, IBP, NIBP, SpO₂, PR, TEMP, CO₂, CO, and AG.

To set the desired baud rate:

- 1. Select [Main Menu] \rightarrow [Maintenance >>] \rightarrow [User Maintenance >>] \rightarrow enter the required password \rightarrow [Others >>].
- 2. Set [DIAP Baud Rate]. The higher the DIAP baud rate goes, the faster data is sent.

23.3 Nurse Call

The monitor provides a nurse call connector to output nurse call signal when a user-defined alarm occurs. To obtain nurse call signal, use the nurse call cable (*PN: 009-003116-00*) we supply to connect the hospital nurse call system to the nurse call connector of the monitor and then follow this procedure:

- 1. Select [Main Menu] \rightarrow [Maintenance >>] \rightarrow [User Maintenance >>] \rightarrow enter the required password.
- 2. Select [**Others** >>] to access the [**Others**] menu.
- 3. Select [Nurse Call Setup >>] to change the nurse call settings as follows:
- Select [Signal Type] and then select [Pulse] or [Continuous].
 - [Pulse]: the nurse call signal is a pulse signal and each pulse lasts one second. When multiple alarms occur simultaneously, only one pulse signal is output. If an alarm occurs but the previous one is not cleared, a new pulse signal will also be output.
 - [Continuous]: the nurse call signal lasts until the alarm ends, i.e. the duration of a nurse call signal is equal to that of the alarm condition.
- Select [Contact Type] and then select [Normally Open] or [Normally Closed].
 - [Normally Open]: select if your hospital's nurse call relay contact is normally open.
 - [Normally Closed]: select if your hospital's nurse call relay contact is normally closed.

- Select [**Alm Lev**] and set the alarm level for nurse call-triggering alarms.
- Select [Alarm Cat.] and then select the category to which the nurse call-triggering alarms belong.

Alarm conditions are indicated to nurses only when:

- The nurse call system is enabled,
- An alarm that meets your preset requirements occurs, and
- The monitor is not in the alarm paused or silence status.

WARNING

- To obtain the nurse call signal, use the nurse call cable (*PN: 009-003116-00*) we supply. Otherwise the nurse call function will not work and the monitor may be damaged.
- Do not rely exclusively on the nurse call system for alarm notification. Remember that the most reliable alarm notification combines audible and visual alarm indications with the patient's clinical condition.

NOTE

• If no setting is selected from [Alm Lev] or [Alarm Cat.], no nurse call signal will be triggered whatever alarms occur.

23.4 Network Setup

CAUTION

• Disconnecting from the network may result in loss of transmitted data, including parameter waveforms and measurements, alarm events, trends and patient data, or cause function failure. In case of network disconnection, check the patient and solve the network problem as soon as possible.

23.4.1 Setting the Network Service Quality Level

To set the quality of service (QoS):

- Select [Main Menu]→[Maintenance>>]→[User Maintenance>>]→enter the required password.→[Network Setup >>]→[QoS Setup >>].
- 2. Select the desired value for [**Realtime Monitoring**]. This sets the service quality of network connection for important realtime network transactions such as parameter measurements, waveforms, and alarms. The value ranges from 0 to 7. The greater the value, the higher priority the network transaction.
- 3. Select the desired value for [**Others**]. This sets the service quality of network connection for secondary non-realtime network transactions. The value ranges from 0 to7. The greater the value, the higher priority the network transaction.

23.4.2 Setting the Multicast Parameters

Multicast parameters must be configured before use on a network.

To set the multicast parameters:

- 1 Select [Main Menu]→[Maintenance>>]→[User Maintenance>>]→enter the required password.→[Network Setup >>]→[Multicast Setup >>].
- 2. Set [Multicast Addr] and [TTL].
- 3. Select [**Ok**] to save the setting.

23.5 MLDAP

MLDAP refers to Mindray LDAP (Lightweight Directory Access Protocol). It is an independent process which can be installed on application server (Windows). MLDAP provides user authorization and authentication.

The MLDAP server is connected to the hospital's LDAP or ADT server. All monitoring devices are connected to the MLDAP server to perform authorization and authentication for the following operations:

- Changing alarm settings
- Changing arrhythmia settings
- Accessing the [User Maintenance] menu

23.5.1 Setting MLDAP

To access the MLDAP server, you should set your monitor as follows:

- Select [Main Menu]→[Maintenance>>]→[User Maintenance>>]→enter the required password.→[Network Setup >>]→[User Authentication Service Setup >>].
- 2. Set IP address and port of the MLDAP server.

23.5.2 Selecting Password for User Authentication

You can select what password is used when changing alarm settings, arrhythmia settings, and accessing the Maintenance menu. To select the passwords:

- Select [Main Menu]→[Maintenance>>]→[User Maintenance>>]→enter the required password.→[Password Protection >>].
- 2. Select [Alarm Setup] to set password for changing alarm settings.
 - [No Password]: changing alarm settings is not password protected.
 - [Local Password]: changing alarm ON/OFF switch, alarm limit, and alarm priority is password protected. The monitor's local password is required to change alarm settings.
 - [User Password]: changing alarm ON/OFF switch, alarm limit, and alarm priority is password protected. The user name and password saved in the MLDAP server are required to change alarm settings.
- 3. Select [Arrh. Alarm Settings] to set password for changing arrhythmia settings.
 - [No Password]: changing arrhythmia settings is not password protected.
 - [Local Password]: changing arrhythmia ON/OFF switch, alarm priority, and arrhythmia threshold is password protected. The monitor's local password is required to change arrhythmia settings.

- [User Password]: changing arrhythmia ON/OFF switch, alarm priority, and arrhythmia threshold is password protected. The user name and password saved in the MLDAP server are required to change arrhythmia settings.
- 4. Select [User Maintenance] to set password for changing user maintenance settings.
 - [Local Password]: changing user maintenance settings is password protected. The monitor's local password is required to change user maintenance settings.
 - [User Password]: changing user maintenance settings is password protected. The user name and password saved in the MLDAP server are required to change user maintenance settings.

To change the monitor's local password for changing alarm settings and arrhythmia settings, select [Modify Local Alarm Password>>], enter the new password, and select [OK].

To change the monitor's local password for accessing the [User Maintenance] menu, select [Modify Local User Maintenance Password>>], enter the new password, and select [OK].

24.1 Overview

This monitor is designed to operate from rechargeable Lithium-ion battery power when AC power is not available. It is recommended to always install fully charged batteries into the monitor to ensure normal monitoring in case of accidental power failure. If the AC power is interrupted during patient monitoring, the monitor automatically runs from the battery. The cPM 12 patient monitor is capable of storing up to two batteries. The cPM 8 patient monitor is capable of storing one battery.

On-screen battery symbols indicate the battery status as follows:

Indicates full battery capacity.

Indicates that the battery works correctly. The solid portion represents the current charge level of the battery in proportion to its maximum charge level.

Indicates that the battery has a low charge level and needs to be charged.

Indicates that the battery is almost depleted and needs to be charged immediately.

Indicates that no battery is installed.

The capacity of the battery is limited. When the battery is low, the technical alarm area displays [**Low Battery**], the alarm lamp flashes, and monitor produces an alarm sound.

If the battery is depleted, the battery symbol on the screen flashes, the technical alarm area displays [**Battery Depleted**], the alarm lamp flashes, and the monitor produces alarm sound. Connect the equipment to AC mains to run the equipment and charge the battery. Otherwise the equipment will shut down.

24.2 Safety

CAUTION

- Keep the battery out of children's reach.
- Use only specified batteries.
- Keep the batteries in their original package until you are ready to use them.
- Do not expose batteries to liquid.
- High ambient temperature shortens battery run time.
- Extremely high ambient temperature may cause battery overheat protection, and then the monitor will turn off.
- If the battery shows signs of damage or signs of leakage, replace it immediately. Do not use a faulty monitor battery.
- Do not crush, drop or puncture the battery. Mechanical abuse can lead to internal damage and internal short circuits. If a battery has been dropped or banged against a hard surface, whether damage is externally visible or not, remove the battery from use and dispose of it properly.
- Please replace your battery when it reaches the end of its service life. Failure to replace the battery may cause serious damage to your device from battery overheating.

NOTE

• Charge the batteries in the monitor.

24.3 Battery Guidelines

Battery life expectancy depends on how frequent and how long it is used. For a properly maintained and stored lithium ion battery, its life expectancy is about 3 years. For more aggressive use models, life expectancy can be less. We recommend replacing lithium ion batteries after 500 full charge/discharge cycles or every 3 years from first use, whichever occurs first.

To get the most out of the battery, observe the following guidelines:

- The battery performance test must be performed every six months, before monitor repairs, or whenever poor battery performance is suspected.
- Condition the batteries regularly to maintain their useful life. For details, refer to **24.6 Conditioning a Battery**.
- Remove the battery before the monitor is shipped or will not be used for more than two weeks. Leaving the battery in a monitor that is not in regular use will shorten the life of the battery.
- The shelf life of a lithium lon battery is about 6 months when the battery is stored with the battery power being 50% of the total power. In 6 months the battery power must be depleted before the Lithium lon battery is fully charged. Then run the monitor on this fully charged battery. When its battery power becomes 50% of the total power, take out the battery from the monitor and store it.
- Store batteries properly. For details, refer to **24.8 Storing a Battery**.

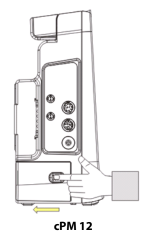
24.4 Installing or Replacing a Battery

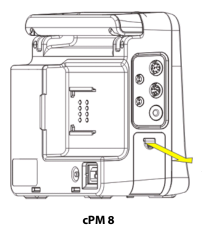
When the cPM 12 patient monitor uses two battery packs, one battery pack can be easily exchanged while the monitor operates from the other. If the cPM 12 patient monitor uses one battery pack, you should insert a new battery pack before the old one depletes.

When the cPM 8 patient monitor is operating on battery power, make sure the monitor is powered off before replacing a battery.

To install or replace a battery:

1. Open the battery door.





Pull here to open the battery door

- 2. Push aside the latch securing the battery and then remove the battery.
- 3. Place the new battery into the slot.
- 4. If necessary, replace the other battery following the steps above.
- 5. Restore the latch to the original position and close the battery door.

24.5 Charging a Battery

To optimize performance, a fully (or nearly fully) discharged battery should be charged as soon as possible. The battery is charged regardless of whether or not the monitor is currently turned on.

To charge the battery:

- 1. Install the battery in the monitor.
- 2. Connect the monitor with the AC power source.

When the battery is charging, the AC power indicator and battery indicator are both on. If the monitor is powered on, the battery status symbol on the monitor screen displays when the charging is complete.

24.6 Conditioning a Battery

The performance of rechargeable batteries may deteriorate over time. If a battery is not conditioned for a prolonged time, its charge indication may not be accurate and you may wrongly evaluate the remaining battery runtime. Keeping the battery continuously fully charged without conditioning will speed up battery aging and shorten its life time. You should condition a battery every three months, before it is used for the first time, or when its run time becomes noticeably shorter.

A battery conditioning cycle is one uninterrupted charge of the battery, followed by an uninterrupted battery discharge and charge.

NOTE

• The battery charge level indicator does not indicate the capacity or operating time of the battery. It only indicates the current battery charge level. The actual battery capacity decreases over time with the use of the battery. For an old battery, its capacity and operating time may not fulfill battery specifications even when the battery charge level indicates the battery is fully charged. Please replace the battery if its operating time is significantly lower than the specified time.

To condition a battery:

- 1. Disconnect the monitor from the patient and stop all monitoring and measuring procedures.
- 2. Insert the battery to be conditioned in the monitor battery slots. For cPM 8, you need to turn it off before performing this step.
- 3. Apply AC power to the monitor and allow the battery to charge uninterruptedly for 10 hours.
- 4. Remove AC power and allow the monitor to run from the battery until it shuts off.
- 5. Apply AC power again to the monitor and allow the battery to charge uninterruptedly for 10 hours.
- 6. This battery is now conditioned and the monitor can be returned to service.

CAUTION

• Do not use the monitor to monitor the patient during battery conditioning.

NOTE

• Do not interrupt battery conditioning.

24.7 Checking Battery Performance

The performance of a rechargeable battery will deteriorate over time. The battery performance test should be performed every six months, before monitor repairs, or whenever the battery performance is diminished.

To check the battery performance:

- 1. Disconnect the monitor from the patient and stop all monitoring and measuring procedures.
- 2. Turn off the monitor.
- 3. Apply AC power to the monitor and allow the battery to charge uninterruptedly for 6 hours.
- 4. Remove AC power and allow the monitor to run from the battery until it shuts off.
- 5. Record the battery operating time.

The battery operating time directly reflects its performance. If the operating time of a battery is noticeably shorter than that stated in the specifications, replace the battery or contact the Mindray service personnel.

NOTE

• Battery operating time depends on the device configuration and operation. For example, high display brightness or measuring NIBP repeatedly will shorten the battery operating time.

24.8 Storing a Battery

When storing batteries, make sure that the battery terminals do not come into contact with metallic objects. If batteries are stored for an extended period of time, they should be placed in a cool place with a partial charge of 40% to 60% capacity.

Stored batteries should be conditioned every 6 months. Refer to section 24.6 Conditioning a Battery for details.

NOTE

- Remove the battery from the equipment if the equipment is not used for a prolonged period of time (for example, several weeks). Otherwise the battery may be over discharged and damaged.
- Storing batteries at high temperature for an extended period of time will significantly shorten their life expectancy.
- Storing batteries in a cool place will slow the aging process. Ideally the batteries should be stored at 15 °C. Do not store the batteries in an environment above 60 °C or lower than -20 °C.

24.9 Recycling a Battery

Remove the old battery from the monitor and recycle it properly. Follow local laws for proper battery disposal.

WARNING

• Do not open batteries, heat above 60 °C, incinerate batteries, or short the battery terminals. They may ignite, explode, leak, or heat up, causing personal injury.

25.1 General Points

In this chapter we only describe cleaning and disinfection of the main unit, modules and certain accessories. For the cleaning and disinfection of other reusable accessories, refer to instructions for use of the corresponding accessories.

Keep your equipment and accessories clean. To avoid damage to the equipment, follow these guidelines:

- Always follow the manufacturer's instructions for each cleaning/disinfecting agent.
- Do not immerse part of the equipment or accessories into liquid.
- Do not pour liquid onto the equipment or accessories.
- Do not allow liquid to enter the case.
- Never use abrasive materials (such as steel wool or silver polish), or erosive cleaners (such as acetone or acetone-based cleaners).

WARNING

• Be sure to turn off the system and disconnect all power cables from the outlets before cleaning the equipment.

CAUTION

- If you spill liquid on the equipment or accessories, contact Mindray or your service personnel.
- Avoid wetting the pins and metal parts of the main unit, modules or accessories during cleaning and disinfection.
- Use only Mindray approved cleaners and disinfectants and methods listed in this chapter to clean or disinfect your equipment. Warranty does not cover damage caused by unapproved substances or methods.
- We make no claims regarding the efficacy of the listed chemicals or methods as a means for controlling infection. For the method to control infection, consult your hospital's Infection Control Officer or Epidemiologist.
- The responsible hospital or institution shall carry out all cleaning and disinfection procedures specified in this chapter.
- Refer to the respective instructions for use of the cleaning agents and disinfectants.
- Do not mix disinfecting solutions (such as bleach and ammonia) as hazardous gases may result.

NOTES

• To clean or disinfect reusable accessories, refer to the instructions delivered with the accessories.

25.2 Cleaning

Your equipment should be cleaned on a regular basis. Before cleaning the equipment, consult your hospital's regulations for cleaning the equipment.

Product Name	Product Type	Ingredients	
Sodium hypochlorite bleach	Liquid	Sodium hypochlorite 10%	
Hydrogen peroxide	Liquid	Hydrogen peroxide 3%	
Isopropanol	Liquid	Isopropanol 70%	
Rely+On [™] Virkon [®]	Powder	Used as 1% solution*	
		Biocidal active:	
		Pentapotassium bis	
		(peroxymonosulphate) bis (sulphate)	
		(500g/kg)	
		Contains dipotassium peroxodisulphate.	
1-Propanol	Liquid	1-Propanol 50%	
Perform [®]	Powder	Used as 1% solution*	
		100g contains 45g Pentapotassium bis	
		(peroxymonosulphate) bis (sulphate).	
		5 % to 15% anionic surfactants;	
		<5% non-ionic surfactants;	
		<5% soap and phosphonates, perfumes	

Note: The "*" symbol indicates a cleaning solution should be prepared according to the concentration directed by the accompanying instructions for use.

To clean your equipment, follow these rules:

- 1. Shut down the monitor and disconnect it from AC power.
- 2. Clean the display screen using a soft, clean cloth dampened with a cleaning agent listed above.
- 3. Clean the exterior surface of the equipment using a soft cloth dampened with a cleaning agent listed above.
- 4. Wipe off all the cleaning solution with a dry cloth after cleaning if necessary.
- 5. Dry your equipment in a ventilated, cool place.

25.3 Disinfecting

Disinfect the monitor as required in your hospital's servicing schedule. Cleaning equipment before disinfecting is recommended.

The following table lists approved disinfectants:

Product Name	Product Type	Ingredients
Sodium hypochlorite bleach	Liquid	Sodium hypochlorite 10%
Isopropanol	Liquid	Isopropanol 70%
Rely+On™Virkon [®]	Powder	Used as 1% solution*
		Biocidal active:
		Pentapotassium bis
		(peroxymonosulphate) bis (sulphate)
		(500g/kg)
		Contains dipotassium peroxodisulphate.
1-Propanol	Liquid	1-Propanol 50%
Perform [®]	Powder	Used as 1% solution*
		100g contains 45g Pentapotassium bis
		(peroxymonosulphate) bis (sulphate).
		5 % to 15% anionic
		surfactants;<5% non-ionic
		surfactants; <5% soap and
		phosphonates, perfumes

Note: The "*" symbol indicates a disinfectant should be prepared according to the concentration directed by the accompanying instructions for use.

CAUTION

• Never use EtO or formaldehyde for disinfection.

25.4 Sterilization

Sterilization is not recommended for this equipment, related products, accessories, or supplies unless otherwise indicated in the Instructions for Use that accompany the products, accessories or supplies.

FOR YOUR NOTES

CAUTION

- Failure on the part of the responsible individual hospital or institution using this equipment to implement a recommended maintenance schedule may cause undue equipment failure and possible health hazards.
- The safety checks or maintenance involving any disassembly of the equipment should be performed by professional servicing personnel. Otherwise, undue equipment failure and possible health hazards could result.
- No modification of this equipment is allowed.
- Do not open the equipment housings. All servicing and future upgrades must be carried out by the personnel trained and authorized by Mindray only.
- If you discover a problem with any of the equipment, contact your service personnel or Mindray.
- The service personnel must be properly qualified and thoroughly familiar with the operation of the equipment.

26.1 General Inspection

A thorough inspection should be performed by qualified service personnel to ensure the reliability before first use, after your monitor has been used for 6 to 12 months, or whenever your monitor is repaired or updated.

Follow these guidelines when inspecting the equipment:

- Make sure that the environment and power supply meet the specifications.
- Inspect the equipment and its accessories for mechanical damage.
- Inspect all power cords for damage, and make sure that their insulation is in good condition.
- Make sure that only specified accessories are applied.
- Inspect if the alarm system functions correctly.
- Make sure that the recorder functions correctly and the recorder paper meets the specifications.
- Make sure that the batteries meet the performance specifications.
- Make sure that the monitor is in good working condition.

In case of any damage or abnormity, do not use the monitor. Contact the hospital's biomedical engineers or your service personnel immediately.

26.2 Maintenance and Testing Schedule

The following maintenance and tests, except for visual inspection, power on test, touchscreen calibration, battery check, ECG calibration, NIBP leakage, CO₂ leakage, and AG leakage, should be carried out by the service personnel only. Ensure the monitor is safety and performance tested by qualified service personnel before initial installation, after repair or upgrade, or during regularly scheduled maintenance. Contact your service personnel if any maintenance is required. Make sure to clean and disinfect the equipment before any test and maintenance.

CAUTION

- Changing the settings in [User Settings >>] and [Factory Maintenance >>] menus may cause data loss.
- Service personnel should acquaint themselves with the test tools and make sure that the test tools and cables are applicable.

Check/Maintenance Item		Recommended Frequency	
Performance Tests			
Visual inspection		When first installed or reinstalled.	
ECG test	Performance test	1. If the user suspects that the measurement is incorrect.	
ECG lest	Verificvation	2. Following any repairs or replacement of relevant module.	
Resp performance test		3. At least once every two years.	
SpO ₂ test			
NIBP test	Pressure check	Note: At least once a year is recommended for NIBP, CO ₂ and AG.	
NIDF test	Leakage test		
Temp test			
IBP test	Performance test		
C.O. test			
CO tosts and	Leakage test		
CO ₂ tests and calibration	Pressure check		
	Calibration		
	Leakage test		
AG test	Performance test		
	Calibration		
Nurse call relay performa	ance test	If the user suspects that the analog output does not work well.	
Analog output performa	nce test		
Electrical Safety Tests			
Electrical safety tests		At least once every two years.	
Other Tests			
Power on test		1. When first installed or reinstalled.	
Power on test		2. Following any maintenance or the replacement of any main unit par	
Touchscreen calibration		1. When the touchscreen appears abnormal.	
		2. After the touchscreen is replaced.	
Recorder check		Following any repair or replacement of the recorder.	
	Functionality test	1. When first installed.	
Battery check		2. Whenever a battery is replaced.	
	Performance test	Once a year or if the battery run time reduced significantly.	

26.3 Checking Monitor and Module Information

To view the information about system start time, selftest, etc., select [Main Menu] \rightarrow [Maintenance >>] \rightarrow [Monitor Information >>]. You can print out the information for the convenience of troubleshooting. The information will not be saved during shut down.

You can also view the information about the monitor configuration and system software version by selecting [Main Menu] \rightarrow [Maintenance >>] \rightarrow [Software Version >>].

26.4 ECG Verification

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

- 1. Select the ECG parameter window or waveform area \rightarrow [**Filter**] \rightarrow [**Diagnostic**].
- Select [Main Menu]→[Maintenance >>]→[User Maintenance >>]→enter the required password→[Module Maintenance >>]→[Calibrate ECG]. A square wave appears on the screen and the message [ECG Calibrating] is displayed.
- 3. Compare the amplitude of the square wave with the wave scale. The difference should be within 5%.
- 4. After the calibration is completed, select [Stop Calibrating ECG]

You can print the square wave and wave scale and then measure the difference between them if necessary. If the difference exceeds 5%, contact your service personnel.

26.5 NIBP Leakage Test

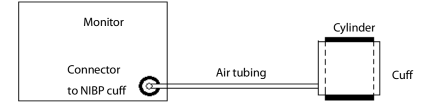
The NIBP leakage test checks the integrity of the system and of the valve. It is required at least once a year or when you doubt the measured NIBP. If the test failed, corresponding prompt messages will be given. If no message is displayed, it means no leakage is detected.

Tools required:

- An adult cuff
- An air tubing
- A correct sized cylinder

Follow this procedure to perform the leakage test:

- 1. Set the patient category to [Adu].
- 2. Connect the cuff to the NIBP connector on the monitor.
- 3. Wrap the cuff around the cylinder as shown below.



- 4. Select [Main Menu]→[Maintenance >>]→[User Maintenance >>]. Enter the required password and then select [Ok].
- 5. Select [Module Maintenance >>]→[NIBP Leakage Test]. The NIBP display shows [Leakage Testing...].

After about 20 seconds, the monitor will automatically deflate. This means the test is completed. If the message [**NIBP Pneumatic Leak**] is displayed, it indicates that the NIBP airway may have leakages. Check the tubing and connections for leakages. If you ensure that the tubing and connections are all correct, perform a leakage test again.

If the problem persists, contact your service personnel.

NOTE

• The leakage test is intended for use to simply determine whether there are leakages in the NIBP airway. It is not the same as that specified in the EN 1060-3 standard.

26.6 CO₂ Leakage Test

For sidestream and microstream CO_2 modules, leakage test is needed every year or when you suspect the measurement. Follow this procedure to perform the test:

- 1. Connect the CO₂ module with the patient module.
- 2. Wait until CO₂ warmup is finished and then use your hand or other objects to completely block the gas inlet of the module or watertrap. The sidestream and microstream CO₂ modules will behave as follows:
 - Sidestream: The alarm message [CO2 FilterLine Err] is displayed on the screen after certain time. Block the gas inlet for another 30 s. If the alarm message does not disappear, it indicates that the module does not leak.
 - Microstream: The alarm message [CO2 Purging] is displayed on the screen after certain time. Block the gas
 inlet for another 30s. If alarm message [CO2 FilterLine Err] is shown, it indicates that the module does not
 leak.

26.7 AG Tests

26.7.1 AG Leakage Test

Follow this procedure to perform the test:

- 1. Plug the AG module into the module rack.
- 2. Wait until the AG module warmup is finished, and then use your hand or other objects to completely block the gas inlet of the AG module. An alarm message [**AG Airway Occluded**] will appear on the screen.
- 3. Block the gas inlet for 60 s. Select [Main menu] \rightarrow [Maintenance >>] \rightarrow [User Maintenance >>] \rightarrow enter the required password \rightarrow [Calibrate AG >>], and check that current flow rate is less than 10ml/min.

If the flow rate is less than 10ml/min and the alarm message [AG Airway Occluded] does not disappear, it indicates that the module does not leak.

If the alarm message does not appear, or the flow rate is greater than or equal to 10ml/min, it indicates that the module leaks. If the problem remains, contact your service personnel for help.

26.8 Calibrating the Touchscreen

- 1. Select [Main Menu] \rightarrow [Maintenance >>] \rightarrow [Cal. Touchscreen].
- 2. 🛨 will, in turn, appear at different positions of the screen. Select each 🛨 as it appears on the screen.
- 3. After the calibration is completed, the message [Screen Calibration Completed!] is displayed. Select [Ok] to confirm the completion of the calibration.

26.9 Entering/Exiting Demo Mode

To enter the Demo mode:

- 1. Select [Main Menu]→[Maintenance >>].
- 2. Select [Demo >>]. Enter the required password and then select [Ok].

To exit the Demo mode:

- 1. Select [Main Menu]→[Maintenance >>].
- 2. Select [Exit Demo] and then select [Ok].
- 3. The monitor exits the Demo mode.

\land WARNING

• The Demo mode is for demonstration purpose only. To avoid the potential risk of the simulated data being mistaken for the monitored patient's data, do not enter the Demo mode while monitoring a patient. Otherwise, improper patient monitoring and delayed treatment could occur.

FOR YOUR NOTES

The accessories listed in this chapter comply with the requirements of IEC 60601-1-2 when in use with the patient monitor. The accessory material that contacts the patients has undertaken the bio-compatibility test and is verified to be in compliance with ISO 10993-1. For details about the accessories, refer to the instructions for use provided with the accessory.

WARNING

- Use accessories specified in this chapter. Using other accessories may cause damage to the patient monitor or not meet the claimed specifications.
- Single-use accessories are not designed to be reused. Reuse may cause a risk of contamination and affect the measurement accuracy.

CAUTION

- Check the accessories and their packages for any sign of damage. Do not use them if any damage is detected.
- Use the accessories before the expiry date if their expiry date is indicated.
- The disposable accessories shall be disposed of according to hospital's regulations.

NOTE

• This manual describes all the accessories that are validated for use. Not all accessories are available in every market.

27.1 ECG Accessories

ECG Electrodes

Model	Description	Patient Category	Part No.
31499224	ECG electrode, Covidien, 10 pieces	Adult	0010-10-12304
2245-50	ECG electrode (3M, package of 50)	Pediatric	9000-10-07469
	Radio Opaque, Pre-wired, 3-lead ECG	Neonate	0681-00-0098-01
/	electrodes, AHA, 18" (45.7 cm), 3 pcs/pack,		
	100 packs/box		
	Radio Translucent, Pre-wired, 3-lead ECG	Neonate	0681-00-0098-02
/	electrodes, AHA, 18" (45.7 cm), 3 pcs/pack,		
	100 packs/box		

12-Pin Separable Trunk Cables

Leadwire	Compatible with	Туре	Patient Category	Part No.
supported				
3-leadwire		Defibrillation-proof		0010-30-42720
3-leadwire		ESU-proof	Infant, neonate	0010-30-42724
3-leadwire		Defibrillation-proof		040-000754-00
3/5-leadwire		Defibrillation-proof	Adult podiatuia	0010-30-42719
3/5-leadwire		ESU-proof	Adult, pediatric	0010-30-42723
3/5/6-leadwire	AHA, IEC			009-005266-00
3/5/6-leadwire		Defibuillation are of	Adult padiatria	009-005267-00
3/5/6-leadwire		Defibrillation-proof	Adult, pediatric	009-005268-00
3/5/6-leadwire				009-005269-00
3/5-leadwire		Defibrillation-proof	Adult podiatric	040-001416-00
3/5-leadwire		ESU-proof	Adult, pediatric	009-003652-00

ECG Leadwires

3-Electro	3-Electrode Leadwires					
Туре	Compatible with	Model	Patient Category	Part No.	Length	Remark
		EL6303A	Adult, pediatric	0010-30-42731	1m	Long
		EL6305A	Infant, neonate	0010-30-42896	1m	Long
		/		0012-00-1514-05	0.6m	24"
Clip	АНА	/		0012-00-1514-06	0.9m	36"
Спр		EY6316A	Adult, pediatric	009-004771-00	0.6m	New telemetry, 24"
		EY6305A		009-004772-00	0.9m	New telemetry, 36"
		EL6301B	Adult, pediatric	0010-30-42734	1m	Long
		EL6311B	Infant, neonate	040-000146-00	1m	Long, disposable
		/		0012-00-1503-05	0.6m	24"
		/		0012-00-1503-06	0.9m	36"
Snan	АНА	/		040-000748-01	0.6m	Disposable, 20 pk
Snap	АПА	EY6316B	Adult, pediatric	009-004765-00	0.6m	New telemetry, 24"
		EY6305B		009-004766-00	0.9m	New telemetry, 36"
		EY6310B		009-004777-00	0.6m	New telemetry, disposable, 24"

5-Electr	5-Electrode Leadwires					
Туре	Compatible with	Model	Patient Category	Part No.	Length	Remark
		/		0012-00-1514-01	0.45m	/
		/		0012-00-1514-02	0.6m	/
		/	-	0012-00-1514-03	0.9m	Long
		EL6501A		0010-30-42727	0.6m	/
Clip	АНА	EL6503A		0010-30-42729	1m to 1.4m	Long
		EY6511A		009-004786-00	0.6m	New telemetry, 24"
		EY6512A	Adult, pediatric	009-004787-00	0.9m	New telemetry, 36"
		/		0012-00-1503-01	0.45m	/
		/		0012-00-1503-02	0.6m	/
		/		0012-00-1503-03	0.9m	Long
	AHA				1.4m for	
		EL6501B		0010-30-42735	RL and LL;	Long
					1m for	Long
					others	
Snap		/		040-000746-01	0.6m	Disposable
		EY6511B		009-004782-00	0.6m	New telemetry, 24"
		51/(5120		009-004783-00		New telemetry,
	АНА	EY6512B	Adult, pediatric		0.9m	36"
]			New
		EY6507B		009-004790-00	0.6m	telemetry ,dispos
						able, 24"

27.2 SpO₂ Accessories

Extension Cable

Model	Part No.	Description	Applicable patient
/	115-020768-00	8-pin, purple connector, 2.1 m, Masimo	All
572A	0010-20-42712	8-pin, 2.5 m, Nellcor	All
4089	040-003381-00	RD to LNC Adapter Cable	All
/	040-003310-00	8-pin Masimo Cable	All

SpO₂ Sensors

Masimo SpO ₂ Module				
Model	PN	Description	Applicable patient	Application site
LNCS DCI	0600-00-0126	Reusable SpO ₂ sensor	Adult	Finger
LNCS DCIP	0600-00-0127	Reusable SpO ₂ sensor	Pediatric	Finger
LNCS NeoPt	0600-00-0156	Disposable SpO ₂ sensor	Neonate	Foot
LNCS Neo	0600-00-0157	Disposable SpO ₂ sensor	Adult (> 40 kg)	Finger
			Neonate (< 3 kg)	Foot
LNCS Inf	0600-00-0158	Disposable SpO ₂ sensor	Infant	Тое
			neonate (3 to 20 kg)	Foot
LNCS Pdtx	0600-00-0122	Disposable SpO ₂ sensor	Pediatric	Finger
LNCS Adtx	0600-00-0121	Disposable SpO ₂ sensor	Adult	Finger
4050	0.40,0000776,00	RD SET DCI, Adult reusable	Adult (> 30 kg)	Finger or toe
4050	040-003376-00	sensor		
4051	040-003377-00	RD SET DCI-P, Pediatric reusable	Pediatric (10 to 50 kg)	Finger or toe
4051	040-003377-00	sensor		
4053	040-003380-00	RD Set TC-I SpO ₂ reusable	Adult (> 30 kg)	Ear
4055		Tip-Clip ear sensor, 3ft		
4000	040-003382-00	RD SET adhesive sensor	Adult (> 30 kg)	Finger or toe
4001	040-003383-00	RD SET PDT adhesive sensor	Pediatric (10 to 50 kg)	Finger or toe
4002	040-003384-00	RD Set Infant adhesive sensor	Infant (3 to 20 kg)	Thumb or great toe
4003		RD Set Neo adhesive sensor	Adult (> 40 kg)	Finger or toe
4003	040-003385-00	RD Set Neo adhesive sensor	Neonate (< 3 kg)	Foot or hand
4004	040-003386-00	RD Set NeoPt adhesive sensor	neonate (< 1 kg)	Foot or hand
4005	040-003387-00	RD Set NeoPt-500 Non-adhesive sensor	neonate (< 1 kg)	Foot or hand

Wavelength emitted by the sensors is between 600 nm and 1000 nm.

The maximum photic output consumption of the sensor is less than 18 mW.

The information about the wavelength range and maximum photic output consumption can be especially useful to clinicians (for example, when photodynamic therapy is performed).

27.3 NIBP Accessories

NIBP hoses

Model	Part No.	Description	Applicable patient
CM1901	6200-30-11560	Reusable NIBP hose	Neonate
CM1903	6200-30-09688	Reusable NIBP hose	Adult, pediatric, infant

Cuffs

Model	Part No.	Description	Applicable patient
CM1500A	001B-30-70692	NIBP cuff, single patient use, size 1, 3.1 to 5.7 cm arm	Neonate
		circumference, 20 pcs/box	
CM1500B	001B-30-70693	NIBP cuff, single patient use, size 2, 4.3 to 8.0 cm arm	Neonate
		circumference, 20 pcs/box	
CM1500C	001B-30-70694	NIBP cuff, single patient use, size 3, 5.8 to 10.9 cm arm	Neonate
		circumference, 20 pcs/box	
CM1500D	001B-30-70695	NIBP cuff, single patient use, size 4, 7.1 to 13.1 cm arm	Neonate
		circumference, 20 pcs/box	
/	115-027563-00	Disposable bladderless cuff, 10 - 19 cm, 10 pcs/box	Child
/	115-027564-00	Disposable bladderless cuff, 18 – 26 cm, 10 pcs/box	Small adult
/	115-027565-00	Disposable bladderless cuff, 24 – 35 cm, 10 pcs/box	Adult
/	115-027566-00	Disposable bladderless cuff, 33 – 47 cm, 10 pcs/box	Large adult
/	115-027567-00	Disposable bladderless cuff, 46 – 66 cm, thigh, 5 pcs/box	Adult
/	115-027568-00	Disposable bladderless long cuff, 24 – 35 cm, 10 pcs/box	Adult
/	115-027569-00	Disposable bladderless long cuff, 33 – 47 cm, 10 pcs/box	Large adult
/	115-027713-00	Reusable bladderless cuff, 10 - 19 cm	Child
/	115-027714-00	Reusable bladderless cuff, 18 – 26 cm	Small adult
/	115-027715-00	Reusable bladderless cuff, 24 – 35 cm	Adult
/	115-027716-00	Reusable bladderless cuff, 33 – 47 cm	Large adult
/	115-027717-00	Reusable bladderless cuff, 46 – 66 cm	Thigh
/	115-027718-00	Reusable bladderless long cuff, 24 – 35 cm	Adult
/	115-027719-00	Reusable bladderless long cuff, 33 – 47 cm	Large adult
/	115-031807-00	Starter kit, reusable bladderless cuff	/

27.4 Temp Accessories

Temp Cable

Туре	Model	Remark	Part No.
Extension cable (reusable)	MR420B	Applicable to sensor MR411 and MR412	0011-30-37391
TEMP adapter cable (2-pin to audio)	MR421	/	0010-30-43056
Temperature adapter cable (2-pin plug)	MR420B	Applicable for all patient categories	040-001235-00

Temp Probes

Туре	Model	Patient Category	Measurement Site	Part No.
	MR401B	Adult	Esophageal/Rectal	0011-30-37392
Reusable	MR403B	Adult	Skin	0011-30-37393
Reusable	MR402B	Dadiatuia infant	Esophageal/Rectal	0011-30-37394
	MR404B	Pediatric, infant	Skin	0011-30-37395
	MR411		Esophageal/Rectal	0011-30-37398
	MR412	 Adult, pediatric, infant 	Skin	0011-30-37397
			Esophageal	0206-03-0112-02
			Stethoscope	
Disposable			Esophageal	0206-03-0118-02
	/	Adult	Stethoscope	
			Esophageal/Rectal	0206-03-0209-02
			Esophageal/Rectal	0206-03-0212-02
			Skin	0206-03-0300-02

27.5 IBP/ICP Accessories

Material	Part No.
IBP adapter cable	0010-30-43055
IBP extended cable with dual-receptacle	040-001029-00
BD IBP Cable, 12 pin	001C-30-70757
Edwards IBP Cable, 12 pin	0010-21-12179
Hospira IBP Cable, 12 pin	001C-30-70759

It is proved through tests that the following accessories are compatible with the patient monitor. Only the accessories proceeded by "*" are available from our company. If you want to purchase other accessories, contact respective manufacturers and make sure if these accessories are approved for sale in local.

Manufacturer	Accessories	
	MX961Z14 Logical Cable, to be used in connection with the Adapter Cable (0010-20-42795)	
	MX960 Reusable Transducer Kit	
Smith Medical	MX261 Logical Clamp For Transducer Bracket	
(Medex)	MX262 Logical Clamp For 2 Transducer Mount Plates	
	MX960E6441 Logical Transducer Mounting Plate	
	(More Logical Clamps are available from Medex. For detailed information, contact Medex.)	
	IBP Reusable Cable (REF: 5203511), to be used in connection with the Adapter Cable (0010-20-42795)	
Braun	Combitrans Monitoring Set (contact Braun for detailed information)	
Braun	Combitrans Attachment Plate Holder (REF:5215800)	
	Combitrans Attachment Plate (contact Braun for detailed information)	
	*Trunk cable (0010-21-43082)	
Managan	SP844 Physiological Pressure Transducer	
Memscap	844-26 Monitoring Line Set	
	84X-49 Mounting Bracket	
	Reusable Blood Pressure Monitor Interface Cable (REF: 650-206)	
	Deltran Disposable Pressure Transducer System	
Utah	(More Deltran sensors are available from Utah. For detailed information, contact Utah.)	
Otan	Pole Mount Unit (ERF: 650-150)	
	Deltran Three Slot Organizer, Attaches to I.V. Pole Mount (REF: 650-100)	
	Deltran Four Slot Organizer, Attaches to I.V. Pole Mount (REF: 650-105)	
	* IBP Truwave Reusable Cable (0010-21-12179)	
	Pressure Monitoring Kit With Truwave Disposable Pressure Transducer.	
Edwards	(More Truwave sensors are available from Edwards. For detailed information, contact Edwards.)	
	DTSC IV Pole Clamp for Model DTH4 Backplate Holder	
	DTH4 Disposable Holder for DPT	

27.6 C.O. Accessories

Model	Material	Part No.
CO7702	12Pin C.O. cable.	0010-30-42743
SP4042	TI Sensor	6000-10-02079
SP5045	TI Sensor Housing	6000-10-02080
MX387	12CC Control Syringe W/1CC Stop W/Rotator	6000-10-02081
93522	In-line inject temperature Probe	0012-00-1520

27.7 CO₂ Accessories

Sidestream CO₂ module

Model	Part No.	Description	Applicable patient
4000	M02A-10-25937	Nasal CO ₂ sample cannula	Adult
4100	M02A-10-25938	Nasal CO ₂ sample cannula	Pediatric
4200	M02B-10-64509	Nasal CO ₂ sample cannula	Neonate
60-15200-00	9200-10-10533	Sampling line, 2.5m, disposable	Adult, pediatric
60-15300-00	9200-10-10555	Sampling line, 2.5m, disposable	Neonate
60-14100-00	9000-10-07486	Airway adapter, disposable, straight	Adult, pediatric
040-001187-00	040-001187-00	Airway adapter, disposable, elbow	Neonate
60-13100-00	125-000005-00	Watertrap, 10pcs/box	Adult, pediatric
60-13200-00	125-000006-00	Watertrap, 10pcs/box	Neonate

27.8 AG Accessories (for cPM 12 only)

Model	Part No.	Description	Applicable patient
60-13100-00	125-000005-00	Watertrap, 10pcs/box	Adult, pediatric
60-13200-00	125-000006-00	Watertrap, 10pcs/box	Neonate
60-14100-00	9000-10-07486	Airway adapter, disposable, straight	/
60-15200-00	9200-10-10533	Sampling line, 2.5m, disposable	Adult, pediatric
60-15300-00	9200-10-10555	Sampling line, 2.5m, disposable	Neonate
60-14200-00	9000-10-07487	Airway adapter, disposable,elbow	Adult, pediatric
/	115-052162-00	AG exhasust gas outlet assembly	Adult, pediatric

27.9 Micellaneous Accessories

Material	Part No.
Lithium battery, LI23S002A	022-000008-00
Power cord (America)	DA8K-10-14452
Grounding cable	1000-21-00122
Nurse call cable (\leq 60W, \leq 2A, \leq 36VDC, \leq 25VAC)	009-003116-00
Analog output cable	009-003117-00
Synchronization cable	009-003118-00
DIAP cable	009-003436-00
Elo ET1929LM, 19-inch black LCD	023-001128-00
USB drive, 4G, USB 2.0 (TRANSCEND)	023-000217-00
USB drive, 4G, USB 2.0 (APACER)	023-000218-00
HP LaserJet Enterprise M608n	023-001566-00
Recorder	TR6F-30-67306
Thermal paper	A30-000001

Material	Part No.
Wall mount bracket for external display	045-000936-00
Bedrail Hook subassembly (cPM 12)	115-012698-00
Bedrail Hook subassembly (cPM 8)	115-012697-00
Transition Plate Kit for rolling stand and wall mount	115-012695-00
cPM 12 rolling stand (Transport)	045-002147-00
Passport12 wall mount (VHM)	045-002148-00
iPM/iMEC rolling stand with quick release mount	045-003428-00
Transport stand for rolling stand (cPM 12)	115-002966-00
Quick Release for wall mount (cPM 8/cPM 12)/Quick release for cPM 12 rolling	045-003424-00
stand	045-005424-00
M series wall mount with Quick Release (cPM 8 and cPM 12)	045-003427-00
VHM wall mount (cPM 8 and cPM 12)	115-006911-00

FOR YOUR NOTES

A.1 Monitor Safety Specifications

A.1.1 Classifications

The monitor is classified, according to IEC60601-1:

Type of protection against electrical shock	Class I, equipment energized from an external and internal electrical	
	power source.	
Degree of protection against electrical shock	Type BF defibrillation proof for CO_2 and AG monitoring.	
Degree of protection against electrical shock	Type CF defibrillation proof for ECG, RESP, TEMP, SpO ₂ , NIBP, IBP and C.O	
Mode of operation	Continuous	
Degree of protection against harmful ingress	IPX1 (Protection against vertically falling water drops)	
of water	IFAT (FIOLECTION against vertically failing water drops)	

A.1.2 Environmental Specifications

CAUTION

• The equipment may not meet the performance specifications if stored or used outside the specified temperature and humidity ranges.

Main unit			
Item	Operating conditions	Storage conditions	
Tomporature (%C)	0 to 40	cPM 8: -30 to 70	
Temperature (°C)		cPM 12: -20 to 60	
Relative humidity (noncondensing)	15% to 95%	10% to 95%	
Barometric (kPa)	57.0 to 107.4	16.0 to 107.4	

Microstream CO ₂ module			
Item	Operating conditions	Storage conditions	
Temperature (°C)	0 to 40	-20 to 60	
Relative humidity (noncondensing)	15% to 95%	10% to 95%	
Barometric (kPa)	57.3 to 105.3	57.3 to 105.3	

Sidestream CO ₂ module		
Item	Operating conditions	Storage conditions
Temperature (°C)	5 to 40	-20 to 60
Relative humidity (noncondensing)	15% to 95%	10% to 95%
Barometric (kPa)	57.3 to 105.3	57.3 to 105.3

AG module		
Item	Operating conditions	Storage conditions
Temperature (°C)	10 to 40	-20 to 60
Relative humidity (noncondensing)	15% to 95%	10% to 95%
Barometric (kPa)	70 to 107.4	70 to 107.4

NOTE

• The environmental specifications of unspecified parameters are the same as those of the main unit of cPM 12.

A.2 Power Supply Specifications

AC power		
Line voltage 100 to 240 VAC (±10%)		100 to 240 VAC (±10%)
Current	ent 1.3 to 0.5 A	
Frequency50/60 Hz (±3Hz)		50/60 Hz (±3Hz)
Battery		
Battery Type		Chargeable Lithium-Ion, 11.1DVC, 4.5 Ah
		\geq 4 hours when powered by one new fully-charged battery (25°C, SpO ₂ , ECG, disconnected from Temp cable, Auto NIBP measurements at intervals of 15 minutes)
Run time	cPM 12	\geq 8 hours when powered by two new fully-charged batteries (25°C, SpO ₂ sensor and ECG cable connected, Temp cable not connected, Auto NIBP measurements at an interval of 15 minutes)
	cPM 8	≥ 6 hours when powered by a new fully-charged battery (25°C, SpO ₂ sensor and ECG cable connected, Temp cable not connected, Auto NIBP measurements at an interval of 15 minutes
Charge time	·	Less than 3 hours to 90%, and less than 4 hours to 100% when the monitor is off. Less than 8 hours to 90%, and less than 12 hours to 100% when the monitor is on.
Shutdown delay at least 20 min (after a low battery alarm first occurs)		at least 20 min (after a low battery alarm first occurs)

A.3 Physical Specifications

Model	Size (Width×Height×Thickness)	Weight	Remark
cPM 12	318mm×274mm×128mm	≤4.5 kg (10 pound)	Standard parameters,
	(12.6 inch $ imes$ 10.8 inch $ imes$ 5.1 inch)	≤4.5 kg (10 poulid)	including touchscreen and
cPM 8 238mm×225mm×128mm		<25 kg(79 pound)	recorder, and battery (2
CFINIO	(9.4 inch $ imes$ 8.9 inch $ imes$ 5.1 inch)	≤3.5 kg (7.8 pound)	batteries for cPM 12)

A.4 Hardware Specifications

A.4.1 Display

Host display				
	Screen Size (diagonal)	Screen type		Resolution
cPM 12	12.1″	color TFT LCD		800×600 pixels
cPM 8	8.4″			
External display				
Screen type Media		Medical-grade TFT LCD		

A.4.2 Recorder

Method	Thermal dot array
Paper width	50 mm±1 mm
Paper speed	25 mm/s or 50 mm/s with accuracy within \pm 5%
Number of waveform channels	Maximum 3

A.4.3 LEDs

Alarm lamp	1 (two color coded: yellow and red)
Power on LED	1 (green)
AC power LED	1 (green)
Battery LED	1 (green)

A.4.4 Audio Indicator

Speaker	Give alarm tones (45 to 85 dB), key tones, QRS tones; support PITCH TONE and
Speaker	multi-level tone modulation; alarm tones comply with IEC60601-1-8.

A.4.5 Monitor Interface Specifications

Power	1 AC power input connector	
Wired network	1 RJ45 connector, 100 Base-TX, IEEE 802.3	
USB	2 connectors, USB 2.0	
Equipotential Grounding Terminal	1	
Multifunctional connector	1	
VGA connector	1	

A.4.6 Outputs

Analog Output			
	Meets the requirements of IEC60601-1 for short-circuit protection and leakage		
Standard	current		
ECG Analog Output			
Dava du vidta	Diagnostic mode: 0.05 to 150 Hz		
Bandwidth (-3dB; reference frequency: 10Hz)	Monitor mode: 0.5 to 40 Hz		
(-Sub, relefence frequency. TOHZ)	Surgical mode: 1 to 20 Hz		
QRS delay	≤ 25 ms (in diagnostic mode, and non-paced)		
Sensitivity	1V/mV ±5%		
	Pace enhancement		
DACE rejection (on bancoment	Signal amplitude: Voh≥2.5V		
PACE rejection/enhancement	Pulse width: 10ms±5%		
	Signal rising and falling time: ≤100µs		
IBP Analog Output (For cPM 12 only)			
Bandwidth (-3dB; reference	DC to 40 Hz		
frequency:1Hz)			
Max transmission delay	30 ms		
Sensitivity	1 V/100 mmHg ±5%		
Nurse Call Signal			
Amplitude	High level: 3.5 to 5 V, providing a maximum of 10 mA output current;		
Ampillude	Low level: < 0.5 V, receiving a maximum of 5 mA input current.		
Rising and falling time	≤1 ms		
Defib Sync Pulse			
Output impedance	≤100 Ω		
Max time delay	35 ms (R-wave peak to leading edge of pulse)		
Amplituda	High level: 3.5 to 5 V, providing a maximum of 10 mA output current;		
Amplitude	Low level: < 0.5 V, receiving a maximum of 5 mA input current.		
Pulse width	100 ms ±10%		
Rising and falling time	≤1 ms		
Alarm output (Network connector)			
Alarm dolay time from the monitor to	The alarm delay time is equal to or smaller than 2 s, measured from the time of		
Alarm delay time from the monitor to the monitor alarm signal generation to the time of the remote equipment			
remote equipment	signal generation.		

A.5 Data Storage

	Trends: 120 hours, at 1 min resolution
Trends	Mid-length trends: 4 hours, at 5 s resolution
	Minitrends: 1 hour, at 1 s resolution
Parameter alarms	100 alarms and manual events and related parameter waveforms. The waveform
	recording length can be 8s, 16s, or 32s.
Arrh. events	100 arrhythmia events and relate waveforms and parameters. The waveform
Arm. events	recording length can be 8s, 16s, or 32s.
NIBP measurements	1000 sets
	48 hours at maximum. The specific storage time depends on the waveforms stored
Full-disclosure waveforms	and the number of stored waveforms.

A.6 Measurement Specifications

The adjustable range of alarm limits is the same as the measurement range for the parameter unless otherwise specified.

ECG			
Standards	Meet standards of EC11, EC13, EN60601-2-27/IEC60601-2-27 and IEC60601-2-25		
Landard	3-lead: I, II, III		
Lead set	5-lead: I, II, III, aVR, aVL, aVF, V		
ECG standard	AHA, IEC		
	1.25 mm/mV (X0.125), 2.5 mm/	mV (X0.25), 5 mm/mV (X0.5), 10 mm/mV (X1), 20	
Display sensitivity	mm/mV (X2), 40 mm/mV (X4)		
	Accuracy: ±5%		
Sween meed	6.25 mm/s, 12.5 mm/s, 25 mm/s	s, 50 mm/s,	
Sweep speed	Accuracy: ±10%		
	Diagnostic mode:	0.05 to 150 Hz	
Bandwidth (-3dB)	Monitor mode:	0.5 to 40 Hz	
	Surgical mode:	1 to 20 Hz	
	Diagnostic mode:	>90 dB	
Common mode rejection ratio	Monitor mode:	>105 dB	
	Surgical mode:	>105 dB	
Notch filter	50/60 Hz		
Differential input impedance	≥5 MΩ		
Input signal range	±8 mV (peak-to-peak value)		
Accuracy of signal reproduction	Based on EC11 to determine system total error and frequency response.		
Electrode offset potential tolerance	±500 mV		
Lead-off detection current	Measuring electrode: <0.1 µA		
	Drive electrode: <1 µA		
Input offset current	Measuring electrode: \leq 0.1 μ A		

A.6.1 ECG

	Drive electrode: ≤1 µA
Baseline recovery time	<5 s (after defibrillation)
Patient leakage current	<10 uA
Calibration signal	1mV (peak-to-peak value)
Calibration signal	Accuracy: ±5%
ESU protection	Cut mode: 300 W
	Coagulate mode: 100 W
	Recovery time: ≤10 s
	In compliance with the requirements in IEC 60601-2-27: Clause 202.6.2.101
	Based on the test method in clause 5.2.9.14 of EC 13, use ECG lead wires which are
ESU noise suppression	in compliance with AAMI. Compared with ECG baseline, the noise of peak to peak
	value ≤2 mV.

Pace Pulse			
	Pace pulses meeting the following conditions are labeled with a PACE marker:		
Pace pulse markers	Amplitude:	±2 to ±700 mV	
race puise markers	Width:	0.1 to 2 ms	
	Rise time:	10 to 100 μs	
	When tested in accordance with the IEC60601-2-27: 201.12.1.101.13, the heart rate		
	meter rejects all pulses meeting the following conditions.		
Pace pulse rejection	Amplitude:	±2 to ±700 mV	
	Width:	0.1 to 2 ms	
	Rise time:	10 to 100 μs	
Pacer pulse detector rejection of fast			
ECG signals	10V/s RTI when measured in accordance with IEC60601-2-27 Section 4.1.4.3.		

HR			
	Neonate:	15 to 350 bpm	
Measurement range	Pediatric:	15 to 350 bpm	
	Adult:	15 to 300 bpm	
Resolution	1 bpm		
Accuracy	±1 bpm or ±1%, whichever i	s greater.	
Sensitivity	200μV (lead ll)		
Minimum ORS detection threshold	Adult and pediatric: 0.16 to 0.48 mV		
Minimum QK3 detection threshold	Neonate: 0.12 to 0.40 mV		
HR averaging method	In compliance with the requirements in IEC60601-2-27: Clause 201.7.9.2.9.101 b) 3), the following method is used: If the last 3 consecutive RR intervals are greater than 1200 ms, the 4 most recent RR intervals are averaged to compute the HR. Otherwise, heart rate is computed by subtracting the maximum and minimum ones from the most recent 12 RR intervals and then averaging them. The HR value displayed on the monitor screen is updated every second.		
Response to irregular rhythm	In compliance with the requirements in IEC60601-2-27: Clause 201.7.9.2.9.101 b) 4), the heart rate after 20 seconds of stabilization is displayed as follows:		

	Ventricular bigeminy (3a):	80±1 bpm	
	Slow alternating ventricular bigeminy (3b): 60±1 bpm		
	Rapid alternating ventricular bigeminy (3c): 120±1 bpm		
	Bidirectional systoles (3d):	90±2 bpm	
	Meets the requirements of IEC60601-2-27:	Clause 201.7.9.2.9.101 b) 5).	
Response time to heart rate change	From 80 to 120 bpm: less than 11 s		
	From 80 to 40 bpm: less than 11 s		
	Meets the requirements of IEC60601-2-27:	Clause 201.7.9.2.9.101 b) 6).	
	Waveform		
	4ah - range: < 11 s		
	4a - range: < 11 s		
Time to alarm for tachycardia	4ad - range: < 11 s		
	Waveform 4bh - range: < 11 s		
	4b - range: < 11 s		
	4bd - range: < 11 s		
	When the test is performed based on IEC60	0601-2-27: Clause 201.12.1.101.17, the	
Tall T-wave rejection capability	maximum T-wave amplitude that can be re	jected is 1.2mV.	
	Asystole, VFib/VTac, Vtac, Vent. Brady, Extre		
Arrhythmia alarms	Bigeminy, Trigeminy, R on T, Run PVCs, PVCs, Tachy, Brady, Missed Beats, Vent.		
	Rhythm, Pacer Not Pacing, Pacer Not Capture, Multif. PVC, Nonsus. Vtac, Pause, Irr.		
	Rhythm, Afib		
QT/QTc Interval Monitoring	1		
	QT: 200 to 800 ms		
Measurement range	QTc: 200 to 800 ms		
	QT-HR: 15 to 150 bpm for adult, 15 to 180 bpm for pediatric and neonate		
Accuracy	QT: ± 30 ms		
Resolution	QT: 4 ms		
	QTc: 1 ms		
HR Alarm limit	Range	Step	
HR High	(low limit + 2) to 300 bpm	HR≪40: 1bpm	
HR Low	15 to (high limit – 2) bpm	HR>40: 5 bpm	
QT/QTc alarm limit	Range (bpm)	Step	
QTc High	200 to 800 ms		
QICHIGH		10	

A.6.2 Resp

Technique	Trans-thoracic impedance
Lead	Options are lead I and II. The default is lead II.
Respiration excitation waveform	<300 μA RMS, 62.8 kHz (±10%)
Baseline impedance range	200 to 2500 Ω (using an ECG cable with 1k Ω resistance)
Bandwidth	0.2 to 2.5 Hz (-3 dB)
Sween mood	6.25 mm/s, 12.5 mm/s or 25 mm/s
Sweep speed	Accuracy: ±10%

Respiration Rate			
	Adult:	0 to 120 rpm	
Measurement range	Pediatric, neonate:	0 to 150 rpm	
Resolution	1 rpm		
Accuracy	7 to 150 rpm:	± 2 rpm or $\pm 2\%$, whichever	ver is greater
Accuracy	0 to 6 rpm:	Not specified.	
Apnea alarm time	10 s, 15 s, 20 s, 25 s, 30 s, 35 s, 40 s		
Alarm limit	Range (rpm)		Step (rpm)
RR High	Adult, pediatric:	(low limit + 2) to 100	
NA HIGH	Neonate:	(low limit + 2) to 150	1
RR Low	0 to (high limit – 2)		

A.6.3 SpO₂

Alarm limit	Range (%)	Step (%)
SpO ₂ High	(low limit + 2) to 100	
	Masimo: Desat to (high limit – 2)	1
SpO ₂ Low	Nellcor: Desat or20 (whichever is greater) to (high limit – 2)	1
Desat	0 to (high limit – 2)	

$Masimo \, \text{SpO}_{\scriptscriptstyle 2} \, \text{Module}$

Standards	Meet standards of ISO9919	
SpO ₂ measurement range	1 to 100%	
Resolution	1%	
	70 to 100%: ±2% (measured without motion in adult/pediatric mode)	
	70 to 100%: \pm 3% (measured without motion in neonate mode)	
Accuracy ¹	70 to 100%: ±3% (measured with motion)	
	1% to 69%: Not specified.	
Refreshing rate	1 s	
SpO ₂ averaging time	2-4 s, 4-6 s, 8 s, 10 s, 12 s, 14 s, 16 s	
Loursefusion conditions	Pulse amplitude: >0.02%	
Low perfusion conditions	Light penetration: >5%	
Low perfusion SpO ₂ accuracy	±2%	
PI measurement range	0.02~20 %	

¹ The Masimo pulse oximeter with sensors have been validated for no motion accuracy in human blood studies on healthy adult volunteers in induced hypoxia studies in the range of 70% to 100% SpO₂ against a laboratory co-oximeter and ECG monitor. This variation equals plus or minus one standard deviation. Plus or minus one standard deviation encompasses 68% of the population. One percent was added to the accuracies for neonatal sensors to account for accuracy variation due to properties of fetal hemoglobin.

The Masimo pulse oximeter with sensors has been validated for motion accuracy in human blood studies on healthy adult volunteers in induced hypoxia studies while performing rubbing and tapping motions at 2 to 4 Hz. At an amplitude of 1 to 2 cm and non-repetitive motion between 1 to 5 Hz. At an amplitude of 2 to 3 cm in induced hypoxia studies in the range of 70% to 100% SpO_2 against a laboratory co-oximeter and ECG monitor. This variation equals plus or minus one standard deviation. Plus or minus one standard deviation encompasses 68% of the population.

² The Masimo pulse oximeter has been validated for low perfusion accuracy in bench top testing against a Biotek Index 2 simulator and Masimo's simulator with signal strengths of greater than 0.02% and a % transmission of greater than 5% for saturations ranging from 70 to 100%. This variation equals plus or minus one standard deviation. Plus or minus one standard deviation encompasses 68% of the population.

Nellcor SpO₂ Module

Standards	Meet standards of ISO9919
Measurement range	0 to 100%
Resolution	1%
	70 to 100%: ±2% (adult/pediatric)
Accuracy	70 to 100%: ±3% (neonate)
0% to 69%: Not specified.	
*: When the SpO $_2$ sensor is applied for neonatal patients as indicated, the specified accuracy range is increased by \pm 1%, to	
compensate for the theoretical effect on oximeter measurements of fetal hemoglobin in neonatal blood.	

Information of the Test Subjects of the Clinical Study Report:

Skin color	Gender	Number	Age (years)	Health
Black	Male	1	28.2±9.19	Healthy
	Female	1		
Yellow	Male	3		
	Female	9		

A.6.4 PR

Alarm limit	Range (bpm)	Step (bpm)
PR High	(low limit +2) to 300	1
PR Low	15 to (high limit-2)	l

PR from Masimo SpO₂

Measurement range	25 to 240 bpm
Resolution	1 bpm
A	±3 bpm (measured without motion)
Accuracy	±5 bpm (measured with motion)
Refreshing rate	1 s
SPO ₂ averaging time	2-4 s, 4-6 s, 8 s, 10 s, 12 s, 14 s, 16 s
	Pulse amplitude: >0.02%
Low perfusion conditions	Light penetration: >5%
Low perfusion PR accuracy	±3 bpm

PR from Nellcor SpO₂

Measurement range	20 to 300 bpm		
Resolution	1 bpm		
Accuracy	20 to 250 bpm: ±3 bpm		
	251 to 300 bpm, not specified		
Refreshing rate	1 s		

PR from NIBP

Measurement range	30 to 300 bpm
Resolution	1 bpm
Accuracy	\pm 3bpm or \pm 3%, whichever is greater

PR from IBP

Measurement range	25 to 350 bpm
Resolution	1 bpm
Accuracy	± 1 bpm or ± 1 %, whichever is greater
Refreshing rate	1 s

A.6.5 NIBP

Standards	Meet standards of EN60601-2-30/IEC60601-2-30, EN1060-1, EN1060-3, SP10 and					
Stariuarus	EN1060-4					
Technique	Oscillometry					
Mode of operation	Manual, Auto and STAT					
Auto mode repetition intervals	1 min, 2 min, 2.5 min, 3 min, 5 min, 10 min, 15 min, 20 min, 30 min, 1 h, 1.5 h, 2 h, 3 h,					
	4 h, 8 h					
STAT mode cycle time	5 min					
Max measurement time	Adult, pediatric: 180 s					
	Neonate: 90 s					
		Adult	Pediatric	Neonate		
Measurement ranges	Systolic:	25 to 290	25 to 240	25 to 140		
(mmHg)	Diastolic:	10 to 250	10 to 200	10 to 115		
	Mean:	15 to 260	15 to 215	15 to 125		
Accuracy	Max mean error: ±5 mmHg					
	Max standard deviation: 8 mmHg					
Resolution	1mmHg					
Initial cuff inflation pressure range (mmHg)	Adult: 80 to 280					
	Pediatric:	atric: 80 to 210				
	Neonate: 60 to 140					
Default initial cuff inflation pressure (mmHg)	Adult:	160				
	Pediatric:	ediatric: 140				
	Neonate:	90				

	Adult:	297±3 mmHg
Software overpressure protection	Pediatric:	297±3 mmHg
	Neonate:	147±3 mmHg
	Adult:	≦330 mmHg
Hardware overpressure protection	Pediatric:	≦330 mmHg
	Neonate:	≦165 mmHg

Alarm limit	Range (mmHg)	Step (mmHg)
	Adult: (low limit+5) to 290	
Sys High	Pediatric: (low limit+5) to 240	
	Neonate: (low limit+5) to 140	
Sys Low	25 to (high limit-5)	
	Adult: (low limit+5) to 260	
Mean High	Pediatric: (low limit+5) to 215	5
	Neonate: (low limit+5) to 125	5
Mean Low	15to (high limit-5)	
	Adult: (low limit+5) to 250	
Dia High	Pediatric: (low limit+5) to 200	
	Neonate: (low limit+5) to 115	
Dia Low	10 to (high limit-5)	

**Measurement accuracy verification: In adult and pediatric modes, the blood pressure measurements measured with this device are in compliance with the American Standard for Non-invasive sphygmomanometers (ISO 81060-2) in terms of mean error and stardard deviation by comparing with intra-arterial or auscultatory measurements (depending on the configuration) in a typical patient population. For auscultatory reference, the 5th Korotkoff sound was used to determine the diastolic pressure.

In neonatal mode, the blood pressure measurements measured with this device are in compliance with the American National Standard for Non-invasive sphygmomanometers (ISO 81060-2) in terms of mean error and stardard deviation by comparing with intra-arterial measurements (depending on the configuration) in a typical patient population.

A.6.6 Temp

Standards	Meet standard of EN12470-4
Technique	Thermal resistance
Measurement range	0 to 50 °C (32 to 122 °F)
Resolution	0.1 °C
Accuracy	±0.1 °C without probe)
Refreshing rate	1 s
Minimum time for accurate	Body surface: <100 s
measurement	Body cavity: <80 s

Alarm limit	Range	Step
T1/T2 High	(low limit +1) to 50°C	
T1/T2 Low	0.1 to (high limit -1) °C	0.1 °C
TD High	0.1 to 50°C	

A.6.7 IBP

Standards	Meet standard of EN60601-2-34/IEC60601-2-34.
Technique	Direct invasive measurement
IBP	
Measurement range	-50 to 300 mmHg
Resolution	1 mmHg
Accuracy	$\pm 2\%$ or ± 1 mmHg, whichever is greater (without sensor)
Refreshing rate	1 s
PPV	
Measurement range	0% to 50%
Pressure transducer	
Excitement voltage	5 VDC, ±2%
Sensitivity	5 μV/V/mmHg
Impedance range	300 to 3000Ω

Alarm limit	Range (mmHg)	Step (mmHg)
Sys High		
Mean High	(low limit + 2) to 300	
Dia High		
Sys Low		
Mean Low	-50 to (high limit – 2)	
Dia Low		

A.6.8 C.O.

Measurement method	Thermodilution method	
	C.O.:	0.1 to 20 L/min
Measurement range	TB:	23 to 43 °C
	TI:	0 to 27 °C
Resolution	C.O.:	0.1 L/min
	TB, TI:	0.1 °C
Accuracy	C.O.:	$\pm 5\%$ or ± 0.1 L /min, whichever is greater
Accuracy	TB, TI:	±0.1 °C (without sensor)
Repeatability	C.O.:	$\pm 2\%$ or ± 0.1 L/min, whichever is greater
Alarm range	ТВ:	23 to 43 ℃

Alarm limit	Range	Step
TB High	(low limit + 1) to 43°C	0.1 °C
TB Low	23 to (high limit - 1) °C	0.1 C

A.6.9 CO₂

Measurement mode	Sidestream, microstream, mainstream	
Technique	Infrared absorption	

Sidestream CO₂ Module

Standard	Meet standard of ISO 21647		
CO ₂ Measurement range	0 to 99 mmHg		
Accuracy*	0 to 40 mmHg: ±2 mmHg		
	41 to 76 mmHg: ±5% of the reading		
	77 to 99 mmHg: ±10% of the reading		
Accuracy drift	Meet the requirement for measurement accuracy within 6 hours		
Resolution	1 mmHg		
Comple flourete	Adult: 70 ml/min, 100 ml/min, 120 ml/min, 150 ml/min		
Sample flowrate	Pediatric, neonate: 70 ml/min, 100 ml/min		
Sample flowrate tolerance	15% or 15 ml/min, whichever is greater.		
Warm-up time	lso accuracy mode: ≤45s		
	Full accuracy mode: ≤10 min		
	Measured with a neonatal watertrap and a 2.5-meter neonatal sampling line, or an		
	adult watertrap and a 2.5-meter adult sampling line:		
Rise time	<400 ms @ 70 ml/min		
Nise time	<330 ms @ 100 ml/min		
	<300 ms @ 120 ml/min		
	<240 ms @ 150 ml/min		
	Measured with a neonatal watertrap and a 2.5-meter neonatal sampling line:		
	<4 s @ 100 ml/min		
	<4.5 s @ 70 ml/min		
Gas sampling delay time	Measured with an adult watertrap and a 2.5-meter adult sampling line:		
	<4.5 s @ 150 ml/min		
	< 5 s @ 120 ml/min		
	<5.5 s @ 100 ml/min		
	<6.5 s @ 70 ml/min		
awRR measurement range	0 to 120 rpm		
awRR measurement precision	±2 rpm		
Apnea time	10 s, 15 s, 20 s, 25 s, 30 s, 35 s, 40 s		

Note: The response time is the sum of the rise time and the delay time.

Effect of interference gases on CO ₂ measurements		
Gas	Concentration (%)	Quantitive effect*
N ₂ O	≤60	
Hal	≤4	
Sev	≤5	±1 mmHg
lso	≤5	
Enf	≤5	
Des	≤15	±2 mmHg

*: means an extra error should be added in case of gas interference when CO_2 measurements are performed between 0-40mmHg.

Alarm limit	Range	Step
EtCO ₂ High	(low limit + 2) to 99 mmHg	
EtCO ₂ Low	1 to (high limit - 2)mmHg	1 mmHg
FiCO ₂ High	1 to 99 mmHg	
awDD High	Adult, pediatric: (low limit + 2) to 100 rpm	
awRR High	Neonate: (low limit + 2) to 150 rpm	
awRR Low	0 to (high limit - 2) rpm	

Microstream CO₂ Module

Standard	Meet standard of ISO 21647			
CO ₂ Measurement range	0 to 99 mmHg			
A	0 to 38 mmHg: ±2 mmHg			
Accuracy*	39 to 99 mmHg:	$\pm 5\%$ of the reading+0	.08% of (the reading-38)	
Accuracy drift	Meet the requirement	for measurement accuracy within	6 hours	
* This accuracy is applied to respirat	ion rate no greater than	80 rpm. For respiration rate greate	r than 80 rpm and EtCO ₂ value	
greater than 18 mmHg, the accurac	y is 4 mmHg or ±12% of	^f the reading, whichever is greater. F	or respiration rate greater than	
60 rpm, the above accuracy can be	achieved by using the C	apnoLine H Set for Infant/Neonatal	. In the presence of interfering	
gases, the accuracy specification de	teriorates by 4% of the a	above accuracy.		
Resolution	1 mmHg			
Sample flow rate	$50^{-7.5}_{+15}$ ml/min			
Initialization time	30 s (typical)			
	Measured with a FilterLine of standard length:			
	2.9 s (typical),			
Response time	4.5 s (Maximum)			
nesponse time	The response time is the sum of the rise time and the delay time			
	Rise time: <190 ms (10% to 90%)			
	Delay time: 2.7 s (typic	cal)		
awRR measurement range	0 to 150 rpm			
	0 to 70 rpm:	±1 rpm		
awRR measurement accuracy	71 to 120 rpm:	±2 rpm		
	121 to 150 rpm:	±3 rpm		
Apnea alarm time	10 s, 15 s, 20 s, 25 s, 30) s, 35 s, 40 s		
Alarm limit	Range Step			
EtCO ₂ High	(low limit + 2) to 99 mmHg			
EtCO ₂ Low	1 to (high limit – 2)mmHg 1 mmHg			
FiCO ₂ High	1 to 99 mmHg			
	Adult, pediatric:	(low limit + 2) to 100 rpm		
awPP High				
awRR High	Neonate:	(low limit + 2) to 150 rpm	1 rpm	

A.6.10 AG (For cPM 12 only)

Standards	Meet standard of ISO 21647			
Technique	Infrared absorption			
	Iso accuracy mode:	≤ 45 s		
Warm-up time	Full accuracy mode:	≤ 10 min		
	Adult, pediatric:	120, 150, 200 ml/min		
Sample flow rate	Neonate:	70, 90, 120 ml/min		
	Accuracy:	±10 ml/min or ±10%, whiche	ver is greater	
	CO ₂ :	0 to 30%		
	O ₂ :	0 to 100%		
	N ₂ O:	0 to 100%		
	Des:	0 to 30%		
Measurement range	Sev:	0 to 30%		
	Enf:	0 to 30%		
	lso:	0 to 30%		
	Hal:	0 to 30%		
	awRR:	2 to 100 rpm		
Resolution	CO ₂ :	1 mmHg		
Resolution	awRR:	1 rpm		
	Gases	Range (% _{REL})	Accuracy (% _{ABS})	
		0 to 1	±0.1	
		1 to 5	±0.2	
	CO2	5 to 7	±0.3	
		7 to 10	±0.5	
		>10	Not specified	
	N ₂ O	0 to 20	±2	
	N ₂ O	20 to 100	±3	
		0 to 25	±1	
	0 ₂	25 to 80	±2	
		80 to 100	±3	
		0 to 1	±0.15	
Full accuracy		1 to 5	±0.2	
Tun accuracy	Des	5 to 10	±0.4	
		10 to 15	±0.6	
		15 to 18	±1	
		>18	Not specified	
		0 to 1	±0.15	
	Sev	1 to 5	±0.2	
		5 to 8	±0.4	
		>8	Not specified	
		0 to 1	±0.15	
	Enf, Iso, Hal	1 to 5	±0.2	
		>5	Not specified	
	awRR	2 to 60 rpm	±1 rpm	
		>60 rpm	Not specified	

Accuracy drift	Meet the requirement	Meet the requirement for measurement accuracy within 6 hours				
Apnea alarm time	10 s, 15 s, 20 s, 25 s, 30	s, 35 s, 40 s				
Refreshing rate	1 s					
	Using the DRYLINE [™] w	vater trap and neor	natal DRYLINE™ sampli	ing line (2.5m):		
		120 ml/min	90 ml/min	70 ml/min		
	CO ₂	≤4S s	≤ 4.5 s	≤5 s		
	N ₂ O	≤4.2 s	≤5 s	≤5.5 s		
	0 ₂	≤4 s	≤5 s	≤6 s		
	Hal, Iso, Sev, Enf	≤4.4 s	≤5.2 s	≤6 s		
Response time	Des	≤4.4 s	≤5 s	≤6 s		
	Using the DRYLINE [™] w	Using the DRYLINE [™] water trap and adult DRYLINE [™] sampling line (2.5m)				
		200 ml/min	150 ml/min	120 ml/min		
	CO ₂	≤4.2 s	≤4.8 s	≤5.5 s		
	N ₂ O	≤4.3 s	≤5 s	≤5.8 s		
	0 ₂	≤4 s	≤4.6 s	≤5.5 s		
	Hal, Iso, Sev, Enf, Des	≤4.5 s	≤5.2 s	≤6 s		
	Primary anesthetic agent					
	In full accuracy mode: 0.15%,					
Anesthetic agent limit	Second anesthetic age	Second anesthetic agent:				
	In full accuracy mode:	0.3% or 5% REL (1	0% in ISO accuracy mo	ode) of primary agent if		
	primary agent is great	primary agent is greater than 10%				

Effect of interference gases on AG measurements					
	Quantitive effect(%ABS) ³⁾				
Concentration(%)	CO ₂	N ₂ O	Agent 1)	02	
/	/	0.1	0	0.2	
/	0.1	/	0.1	0.2	
/	0.1	0.1 ⁵⁾	0.1 ⁴⁾	1.0	
<100%	0.1	0	0	/	
<50%	0.1	0	0	/	
<0.1%	0	0	0	/	
<1%	0.1	0.1	0	/	
<1%	0.1	0.1	0	/	
/	0.1	0	0	/	
/	0.1	0.1	0.1	/	
	Concentration(%) / / / / / / <100%	Concentration(%) Quantitive / / / 0.1 / 0.1 / 0.1 <100%	Quantitive effect(%ABS) ³) Concentration(%) Quantitive effect(%ABS) ³) / CO_2 N_2O / 0.1 0.1 / 0.1 / / 0.1 0.1 / 0.1 0.1 ^{5°} <100%	Quantitive effect(%ABS) ³) Concentration(%) Quantitive effect(%ABS) ³) $(O_2$ N ₂ O Agent ¹) / / 0.1 0 / 0.1 0.1 0.1 / 0.1 0.1 0.1 / 0.1 0.1 ⁵ 0.1 ⁴ 0.1 0 0 0.1 0 0 0.1 0 0 <<100% 0.1 0 0 0 <<10% 0.1 0.1 0 0 <<10% 0.1 0.1 0.1 0 <<11% 0.1 0.1 0 0 <<1% 0.1 0.1 0 0	

1) Agent represents one of Des, Iso, Enf, Sev, and Hal.

2) Multiple agent interference on CO_2 , N_2O and O_2 is typically the same as single agent interference.

3) For CO₂, N₂O and Agents, maximum interference from each gas at concentrations within specified accuracy ranges for each

gas. The total interference of all gases is never larger than $5\%_{\rm REL}$

4) Applicable to type A AG module, representing the interference effect of secondary anesthetic agents on primary anesthetic agent.

5) Measurement interference to type M AG module originates from the applied anesthetic agent that is configured manually.

Alarm limit	Range		Step	
EtCO ₂ High	(low limit + 2) to 99	mmHg		
EtCO ₂ Low	1to (high limit - 2)m	mHg	1 mmHg	
FiCO ₂ High	1 to 99 mmHg			
awRR High	Adult, pediatric: Neonate:	(low limit + 2) to 100 rpm (low limit + 2) to 150 rpm	1 rpm	
awRR Low	0 to (high limit - 2)rp	om		
EtO ₂ High	(low limit + 2) to 100	D %		
EtO ₂ Low	0 to (high limit - 2)%	,)	0.10/	
FiO ₂ High	(low limit + 2) to 100	0 %	0.1%	
FiO ₂ Low	18 to (high limit - 2)	%		
EtN ₂ O High	(low limit + 2) to 100	0 %		
EtN ₂ O Low	0 to (high limit - 2)%	,)	1%	
FiN ₂ O High	(low limit + 2) to 100	(low limit + 2) to 100 %		
FiN ₂ O Low	0 to (high limit - 2)%)		
EtHal/Enf/Iso High	(low limit + 0.2) to 5	.0 %		
EtHal/Enf/Iso Low	0 to (high limit - 0.2)	%		
FiHal/Enf/Iso High	(low limit + 0.2) to 5	.0 %	0.1%	
FiHal/Enf/Iso Low	0 to (high limit - 0.2)	%		
EtSev High	(low limit + 0.2) to 8	.0 %		
EtSev Low	0 to (high limit - 0.2)	%	0.1%	
FiSev High	(low limit + 0.2) to 8	.0 %	0.1%	
FiSev Low	0 to (high limit - 0.2)	%		
EtDes High	(low limit + 0.2) to 1	8.0 %		
EtDes Low	0 to (high limit - 0.2)	%	0.10/	
FiDes High	(low limit + 0.2) to 1	8.0 %	0.1%	
FiDes Low	0 to (high limit - 0.2)	%		

FOR YOUR NOTES

The device meets the requirements of IEC 60601-1-2.

Note

- Using accessories, transducers and cables other than those specified may result in increased electromagnetic emission or decreased electromagnetic immunity of the patient monitoring equipment.
- The device or its components should not be used adjacent to or stacked with other equipment. If adjacent or stacked use is necessary, the device or its components should be observed to verify normal operation in the configuration in which it will be used.
- The device needs special precautions regarding EMC and needs to be installed and put into service according to the EMC information provided below.
- Other devices may interfere with this monitor even though they meet the requirements of CISPR.
- When the inputted signal is below the minimum amplitude provided in technical specifications, erroneous measurements could result.
- Portable and mobile communication equipment may affect the performance of this monitor.

Guidance and Declaration - Electromagnetic Emissions						
The device is intended for use in	The device is intended for use in the electromagnetic environment specified below. The customer or the user of the device					
should assure that it is used in su	ch an environment.					
Emission tests Compliance Electromagnetic environment - guidance						
Radio frequency (RF) emissions	Group 1	The device uses RF energy only for its internal function. Therefore, its				
CISPR 11	RF emissions are very low and are not likely to cause any interference					
		in nearby electronic equipment.				
RF emissions CISPR 11	Class A	The device is suitable for use in all establishments other than				
Harmonic emissions	Class A	domestic and those directly connected to the public				
IEC61000-3-2		low-voltage power supply network that supplies buildings used				
Voltage Fluctuations/Flicker	Complies	for domestic purposes.				
Emissions IEC 61000-3-3						

WARNING

• This equipment/system is intended for use by healthcare professionals only. This equipment/ system may cause radio interference or may disrupt the operation of nearby equipment. It may be necessary to take mitigation measures, such as re-orienting or relocating the [ME EQUIPMENT or ME SYSTEM] or shielding the location.

Guidance and Declaration - Electromagnetic Immunity

The device is intended for use in the electromagnetic environment specified below. The customer or the user of the device should assure that it is used in such an environment.

luc un constant de set	IFC60601 test level	Compliance level	Electromagnetic environment -
Immunity test	IEC60601 test level	Compliance level	guidance
Electrostatic discharge	±6 kV contact	±6 kV contact	Floors should be wood, concrete or
(ESD) IEC 61000-4-2	±8 kV air	±8 kV air	ceramic tile. If floors are covered
			with synthetic material, the relative
			humidity should be at least 30%.
Electrical fast	±2 kV for power supply lines	±2 kV for power supply lines	Mains power quality should be that
transient/burst IEC	±1 kV for input/output lines	±1 kV for input/output lines	of a typical commercial or hospital
61000-4-4			environment.
Surge IEC 61000-4-5	±1 kV line(s) to line(s)	±1 kV line(s) to line(s)	
	±2 kV line(s) to earth	±2 kV line(s) to earth	
Voltage dips, short	<5 % U $_{\rm T}$ (>95 % dip in U $_{\rm T}$) for	<5 % U $_{\rm T}$ (>95 % dip in U $_{\rm T}$) for	Mains power quality should be that
interruptions and	0.5 cycle	0.5 cycle	of a typical commercial or hospital
voltage variations on			environment. If the user of our
power supply input	40 % U _T (60 % dip in U _T) for 5	40 % U $_{\rm T}$ (60 % dip in U $_{\rm T}$) for 5	product requires continued
lines IEC 61000-4-11	cycles	cycles	operation during power mains
			interruptions, it is recommended
	70 % U $_{\rm T}$ (30 % dip in U $_{\rm T}$) for	70 % U _T (30 % dip in U _T) for	that our product be powered from
	25 cycles	25 cycles	an uninterruptible power supply or
			a battery.
	<5 % U _T (>95 % dip in U _T) for	${<}5$ % U_{T} (>95 % dip in U_{T}) for	
	5 s	5 s	
Power frequency	3 A/m	3 A/m	Power frequency magnetic fields
(50/60 HZ) magnetic			should be at levels characteristic of
field IEC 61000-4-8			a typical location in a typical
			commercial or hospital
			environment.

Guidance and Declaration - Electromagnetic Immunity

Incorporate and Compliance Electrometric and and mid-						
assure that it is used in such an environment as described below.						
The device is intended for use in the specified electromagnetic environment. The customer or the user of the device should						

Immunity test	IEC60601 test level	Compliance level	Electromagnetic environment - guidance
Conduced RF	3 Vrms	3Vrms	Portable and mobile RF communications equipment should
IEC61000-4-6	150 kHz to 80 MHz	511115	be used no closer to any part of the system, including cables,
			than the recommended separation distance calculated from
			the equation appropriate for the frequency of the transmitter.
			Recommended separation distances:
			$d = 1.2\sqrt{P}$
Radiated RF	3V/m	3V/m	Recommended separation distances:
IEC61000-4-3	80MHz to 2.5GHz		80 MHz~800 MHz
			$d = 1.2\sqrt{P}$
			800MHz-2.5GHz
			$d = 2.3\sqrt{P}$
			Where, ${\it P}$ is the maximum output power rating of the
			transmitter in watts (W) according to the transmitter
			manufacturer and $m{d}$ is the recommended separation distance
			in meters (m). ^b
			Field strengths from fixed RF transmitters, as determined by
			an electromagnetic site survey ^a , should be less than the
			compliance level in each frequency range ^b .
			Interference may occur in the vicinity of
			equipment marked with the following
			symbol:

Note 1: At 80 MHz to 800 MHz, the separation distance for the higher frequency range applies.

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

^a Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the [ME EQUIPMENT or ME SYSTEM] is used exceeds the applicable RF compliance level above, the [ME EQUIPMENT or ME SYSTEM] should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as re-orienting or relocating the [ME EQUIPMENT or ME SYSTEM].

b Over the frequency range 150 kHz to 80 MHz, field strengths should be less than 3 V/m.

Recommended separation distances between portable and mobile RF communications equipment and the device

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

Rated maximum	Separation distance in meters (m) according to frequency of the transmitter				
output power of	150 kHz - 80 MHz	80 MHz - 800 MHz	800 MHz - 2.5 GHz		
transmitter (W)	$d = 1.2\sqrt{P}$	$d = 1.2\sqrt{P}$	$d = 2.3\sqrt{P}$		
0.01	0.12	0.12	0.23		
0.1	0.38	0.38	0.73		
1	1.20	1.20	2.30		
10	3.80	3.80	7.30		
100	12.00	12.00	23.00		

For transmitters rated at a maximum output power not listed above, the recommended separation distance d in metres (m) can be estimated using the equation applicable to the frequency of the transmitter, where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer.

Note 1: At 80 MHz and 800 MHz, the separation distance for the higher frequency range applies.

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

This chapter lists some of the most important factory default settings for each department in configuration management. You cannot change the factory default configuration itself. However, you can make changes to the settings from the factory default configuration and then save the changed configuration as a user configuration.

C.1 Parameters Configuration

C.1.1 ECG

ECG Setup

	Configurable		
Item Name	In Config Mode	In Monitor Mode	Default
Alm Source	Yes	Yes	HR
Alarm	Yes	Yes	On
Alm Lev	Yes	Yes	Med
			Adu: 120
HR/PR High	Yes	Yes	Ped: 160
			Neo: 200
			Adu: 50
HR/PR Low	Yes	Yes	Ped: 75
			Neo: 100
Gain	Yes	Yes	X1
Sweep	Yes	Yes	25 mm/s
		Yes	General: Monitor
Filter	Vec		OR: Surgery
Filter	Yes		ICU, NICU: Monitor
			CCU: Diagnostic
QRS Volume	Yes	Yes	General, OR: 2
QK3 Volume	les	ies	ICU, NICU, CCU: 1
QRS threhold	No	Yes	0.16 mV
			Auto (if auto lead detection is available);
Lead Set	Yes	Yes	5-lead (if auto lead detection is not
			available)
ECG Display	Yes	Yes	Normal
Notch Filter	Yes	Yes	Weak
Paced	No	Yes	No
Pacer Reject	No	Yes	Off
Smart Lead Off	Yes	Yes	On

QT/QTc Analysis

Item Name	Configurable		Default
item Name	In Config Mode	In Monitor Mode	Delauit
QT Analysis	Yes	Yes	Off
QTc Formula	Yes	Yes	Hodges
Analysis lead	Yes	Yes	All
QTc Alarm	Yes	Yes	Off
∆QTc Alarm	Yes	Yes	Off
QTc Alarm Level	Yes	Yes	Med
∆QTc Alarm Level	Yes	Yes	Med
QTc Record	Yes	Yes	Off
ΔQTc Record	Yes	Yes	Off

Arrh. Analysis

Arrhythmia Threshold Settings				
Item Name	Configurable		Default	
item Name	In Config Mode	In Monitor Mode	Default	
PVCs High	Yes	Yes	10	
Asys. Delay	Yes	Yes	5	
Vtac Rate	Yes	Yes	130	
Vtac PVC	Yes	Yes	6	
Multif. PVC's Window	Yes	Yes	15	
Tashullish	Yes	Yes	Adu: 120	
Tachy High	res	Tes	Ped: 160	
Brady Low	Yes	Yes	Adu: 50	
brady Low	les	Tes	Ped: 75	
Extreme Tachy	Yes	Yes	Adu: 160	
	Tes	Tes	Ped: 180	
Extreme Brady	Yes	Yes	Adu: 35	
	Tes	Tes	Ped: 50	
Vbrd Rate	Yes	Yes	40	
Vbrd PVCs	Yes	Yes	5	
Pause Time	Yes	Yes	2	

Arrhythmia Alarm Settings

Arrnythmia Alarm Settings					
Item Name	Configurable		Default		
item Name	In Config Mode	In Monitor Mode	Alarm Switch	Alarm Level	
Asystole Alarm	Yes	Yes	On	High	
VFib/VTac Alarm	Yes	Yes	On	High	
Vtac Alarm	Yes	Yes	On	High	
Vent. Brady Alarm	Yes	Yes	On	High	
Extreme Tachy Alarm	Yes	Yes	On	High	
Extreme Brady Alarm	Yes	Yes	On	High	

Arrhythmia Alarm Settings				
Item Name	Configurable		Default	
PVCs/min Alarm	Yes	Yes	General, OR, ICU, NICU: Off CCU: On	Med
R on T Alarm	Yes	Yes	General, OR, ICU, NICU: Off CCU: On	Med
Run PVCs Alm Lev Alarm	Yes	Yes	Off	Low
Couplet Alarm	Yes	Yes	Off	Message
Multif. PVC Alarm	Yes	Yes	Off	Med
PVC Alarm	Yes	Yes	Off	Message
Bigeminy Alarm	Yes	Yes	General, OR, ICU, NICU: Off CCU: On	Med
Trigeminy Alarm	Yes	Yes	General, OR, ICU, NICU: Off CCU: On	Med
Tachy Alarm	Yes	Yes	Off	Med
Brady Alarm	Yes	Yes	Off	Med
Pacer Not Pacing Alarm	Yes	Yes	Off	Message
Pacer Not Capture Alarm	Yes	Yes	Off	Message
Missed Beats Alarm	Yes	Yes	Off	Message
Nonsus. Vtac Alarm	Yes	Yes	General, OR, ICU, NICU: Off CCU: On	Med
Vent. Rhythm Alarm	Yes	Yes	Off	Med
Pause Alarm	Yes	Yes	Off	Low
Irr. Rhythm Alarm	Yes	Yes	Off	Message
Afib Alarm	Yes	Yes	Off	Message

C.1.2 RESP

Item Name	Configurable		Default	
item Name	In Config Mode	In Monitor Mode	Derault	
Alarm	Yes	Yes	On	
Alm Lev	Yes	Yes	Med	
RR High	Yes	Yes	Adu, Ped: 30	
кк підп	ies	Tes	Neo: 100	
PP Low	Vec		Adu, Ped: 8	
RR LOW	RR Low Yes Yes	ies	Neo: 30	
Annes Delay	Yes	Yes	Adu, Ped: 20	
Apnea Delay	ies	ies	Neo: 15	
Load	Ver	Vec	Adu, Ped: Auto	
Lead	Yes	Yes	Neo: II	
Gain	Yes	Yes	X2	
Sweep	Yes	Yes	6.25 mm/s	
Detection Mode	Yes	Yes	Auto	
RR Source	No	Yes	Auto	

C.1.3 PR

Item Name	Configurable		— Default	
item Name	In Config Mode	In Monitor Mode		
Alarm	Yes	Yes	On	
Alm Lev	Yes	Yes	Med	
			Adu:	120
HR/PR High	Yes	Yes	Ped:	160
			Neo:	200
			Adu:	50
HR/PR Low	Yes	Yes	Ped:	75
			Neo:	100
PR Source	Yes	Yes	SpO ₂	
Alm Source	Yes	Yes	HR	
Beat Vol	Vec	Ver	General, OR: 2	
	Yes	Yes	ICU, NICU, CCU: 1	

C.1.4 SpO₂

Item Name	Configurable		Default	
item Name	In Config Mode	In Monitor Mode	Delauit	
Alarm	Yes	Yes	On	
Alm Lev	Yes	Yes	SpO _{2:} : Med	
			Desat: High	
SpO ₂ High	Yes	Yes	Adu, Ped: 100	
			Neo:95	
SpO ₂ Low	Yes	Yes	90	
Desat	Yes	Yes	80	
Sensitivity (Masimo)	Yes	Yes	Normal	
Averaging (Masimo)	Yes	Yes	8 s	
Sat-Seconds (Nellcor)	Yes	Yes	0 s	
Sweep	Yes	Yes	25 mm/s	
NIBP Simul	No	Yes	Off	

C.1.5 Temp

Item Name	Configurable		Default
	In Config Mode	In Monitor Mode	Delauit
Alarm	Yes	Yes	On
Alm Lev	Yes	Yes	Med
T1/T2 High (℃)	Yes	Yes	38.0
T1/T2 Low (°C)	Yes	Yes	35.0
TD High (℃)	Yes	Yes	2.0

C.1.6 NIBP

It and Name	Configurable		
Item Name	In Config Mode	In Monitor Mode	— Default
Alarm	Yes	Yes	On
Alm Lev	Yes	Yes	Med
			General: 15 min
			OR: 5 min
Interval	Yes	Yes	ICU: 15 min
			NICU: 30 min
			CCU: 15 min
NIBP End Tone	Yes	Yes	Off
Clock	Yes	Yes	On
			Adu: 160
Initial Pressure (mmHg)	Yes	Yes	Ped: 140
			Neo: 90
			Adu: 80
Cuff Press. (mmHg)	Yes	Yes	Ped: 60
			Neo: 40
Alarm Limit			
			Adu: 160
NIBP-S High (mmHg)	Yes	Yes	Ped: 120
			Neo: 90
			Adu: 90
NIBP-S Low (mmHg)	Yes	Yes	Ped: 70
			Neo: 40
	-		
			Adu: 110
NIBP-M High (mmHg)	Yes	Yes	Ped: 90
			Neo: 70
			Adu: 60
NIBP-M Low (mmHg)	Yes	Yes	Ped: 50
Nibi - Ni Low (mining)	165	165	Neo: 25
			Adu: 90
NIBP-D High (mmHg)	Yes	Yes	Ped: 70
			Neo: 60
			Adu: 50
NIPD D Low (mm La)	Yes	Yes	
NIBP-D Low (mmHg)	ies	165	Ped: 40
			Neo: 20

C.1.7 IBP

	Configurable		
Item Name	In Config Mode	In Monitor Mode	Default
Alarm	Yes	Yes	On
Alm Lev	Yes	Yes	Med
P1 Measure	Yes	Yes	All
P2 Measure	Yes	Yes	All
P3 Measure	Yes	Yes	Mean
P4 Measure	Yes	Yes	Mean
PPV Measurement	Yes	Yes	Off
PPV Source	Yes	Yes	Auto
Sensitivity	Yes	Yes	Med
Sweep	Yes	Yes	25 mm/s
Filter	Yes	Yes	12.5 Hz
Gridlines	Yes	Yes	Off
Art, Ao, UAP, BAP, FAP, I	LV, P1-P2 Arterial Pressu	re Alarm Limits	
			Adu: 160
IBP-S High (mmHg)	Yes	Yes	Ped: 120
			Neo: 90
			Adu: 90
IBP-S Low (mmHg)	Yes	Yes	Ped: 70
			Neo: 55
			Adu: 110
IBP-M High (mmHg)	Yes	Yes	Ped: 90
			Neo: 70
			Adu: 70
IBP-M Low (mmHg)	Yes	Yes	Ped: 50
			Neo: 35
			Adu: 90
IBP-D High (mmHg)	Yes	Yes	Ped: 70
			Neo: 60
			Adu: 50
IBP-D Low (mmHg)	Yes	Yes	Ped: 40
			Neo: 20
PA Alarm Limit			
DA C High (martin)	Vec	Voc	Adu: 35
PA-S High (mmHg)	Yes	Yes	Ped, Neo: 60
PA-S Low (mmHg)	Yes	Yes	Adu: 10
5			Ped, Neo: 24

	Configurable				
Item Name	In Config Mode	In Monitor Mode	Default		
DA M High (mmHg)	Yes	Yes	Adu: 2	20	
PA-M High (mmHg)	res	res	Ped, Neo: 2	26	
PA-M Low (mmHg)	Yes		Adu: 0)	
PA-M LOW (IIIIIIII)	A-M Low (mmHg) Yes Yes	Ped, Neo: 1	2		
PA-D High (mmHg)	Yes	Yes	Adu: 1	6	
	103		Ped, Neo: 4	ł	
PA-D Low (mmHg)	Yes	Yes	Adu: 0)	
	163	fes	Ped, Neo: -	4	
CVP, LAP, RAP, ICP, UVP, P3-P4 V	enous Pressure Alarm Lir	nits			
IBP-M High (mmHg)	Yes	Yes	Adu: 1	0	
	ies	ies	Ped, Neo: 4	ł	
IBP-M Low (mmHg)	Yes	Yes	0		
Art, Ao, BAP, FAP, LV, P1-P2	Arterial Pressure Scale				
Scale (mmHg)	Yes	Yes	0-160		
PA Waveform Scale					
Scale (mmHg)	Yes	Yes	0-30		
CVP, LAP, RAP, ICP, UVP Scal	e				
Scale (mmHg)	Yes	Yes	0-20		
UAP, P3-P4 Venous Pressure	Scale				
Scale (mmHg)	Yes	Yes	0-80		
IBP Overlapping Left Scale					
Scale (mmHg)	Yes	Yes	0-160		
IBP Overlapping Right Scale	IBP Overlapping Right Scale				
Scale (mmHg)	Yes	Yes	0-20		

C.1.8 C.O.

Item Name	Configurable		Defeat	
item Name	In Config Mode	In Monitor Mode	Default	
Alarm	Yes	Yes	On	
Alm Lev	Yes	Yes	Med	
TB High (℃)	Yes	Yes	39.0	
TB Low (°C)	Yes	Yes	36.0	
Comp. Const	Yes	Yes	0.542	
Auto TI	Yes	Yes	Auto	
Manual TI (℃)	Yes	Yes	2.0	
Measuring mode	Yes	Yes	Manual	

C.1.9 CO₂

	Configurable			
Item Name	In Config Mode	In Monitor Mode	Default	
Alarm	Yes	Yes	On	
Alm Lev	Yes	Yes	Med	
Anna Dalau	No		Adu, Ped:	20
Apnea Delay	Yes	Yes	Neo:	15
Operating Mode	Yes	Yes	Measure	
Sweep	Yes	Yes	6.25 mm/s	
Scale (mmHg)	Yes	Yes	50	
RR Source	No	Yes	Auto	
Sidestream CO ₂ Setup				
			Adu,	120 ml/min
Flow Rate	Yes	Yes	Ped:	100 ml/min
			Neo:	70 ml/min
BTPS Compen	Yes	Yes	Off	
N ₂ O Compen	Yes	Yes	0	
			General: 21	
O ₂ Compen	Yes	Yes	OR: 100	
			ICU, NICU, CCU: 21	
Des Compen	Yes	Yes	0	
Microstream CO ₂ Setup				
BTPS Compen	Yes	Yes	Off	
Max Hold	Yes	Yes	20 s	
Auto Standby (min)	Yes	Yes	0	
Alarm Limits				
EtCO ₂ High (mmHg)	Yes	Yes	Adu, Ped:	50
	103	103	Neo:	45
EtCO ₂ Low (mmHg)	Yes	Yes	Adu, Ped:	25
	103	103	Neo:	30
FiCO ₂ High (mmHg)	Yes	Yes	Adu, Ped, Neo:	4
DD High	Vec	Vec	Adu, Ped:	30
RR High	Yes	Yes	Neo:	100
PPLow	Vor	Vor	Adu, Ped:	8
RR Low	Yes	Yes	Neo:	30

C.1.10 AG

	Configurable		
Item Name	In Config Mode	In Monitor Mode	Default
Alarm	Yes	Yes	On
Alm Lev	Yes	Yes	Med
Sweep	Yes	Yes	6.25 mm/s
0.6		N N	OR: On
O ₂ Compen	Yes	Yes	General, ICU, NICU, CCU: Off
Operating Mode	Yes	Yes	Measure
Flow Rate	Yes	Yes	Adu, Ped: 120 ml/min
How Nate	les	163	Neo: 70 ml/min
Auto Standby	Yes	Yes	Off
Apnea Time	Yes	Yes	20 s
RR Source	No	Yes	Auto
CO ₂ Setup			
Wave Type	Yes	Yes	Draw
Scale	Voc	Vec	When unit is mmHg: 50
Scale	Yes	Yes	When unit is % or KPa: 7.0
EtCO Lligh (modella)	Vez	Vec	Adu, Ped: 50
EtCO ₂ High (mmHg)	Yes	Yes	Neo: 45
EtCO Lour (mmHa)	Voc		Adu, Ped: 25
EtCO ₂ Low (mmHg)	Yes	Yes	Neo: 30
$FiCO_2 High (mmHg)$	Yes	Yes	4
DD I lizh	Vec	Vec	Adu, Ped: 30
RR High	Yes	Yes	Neo: 100
RR Low	Yes	Yes	Adu, Ped: 8
RR LOW	les	163	Neo: 30
Gas Setup			
Agent	Yes	Yes	AA
N ₂ O Scale	Yes	Yes	50
O_2 Scale	Yes	Yes	When unit is mmHg: 400
O ₂ Scale	163	163	When unit is % or KPa: 50
AA Scale	Yes	Yes	9.0
Hal/Enf/Iso Scale	Yes	Yes	2.5
Des Scale	Yes	Yes	9.0
Sev Scale	Yes	Yes	4.0
EtO ₂ High	Yes	Yes	88
EtO ₂ Low	Yes	Yes	18
FiO High	Voc	Voc	Adu, Ped: 100
FiO ₂ High	Yes	Yes	Neo: 90
FiO ₂ Low	Yes	Yes	18
EtN ₂ O High	Yes	Yes	55
EtN ₂ O Low	Yes	Yes	0
FiN ₂ O High	Yes	Yes	53

Marrie Marrie	Configurable		Default
Item Name	In Config Mode In Monitor Mode		Detault
FiN ₂ O Low	Yes	Yes	0
EtHal/Enf/Iso High	Yes	Yes	3.0
EtHal/Enf/Iso Low	Yes	Yes	0.0
FiHal/Enf/Iso High	Yes	Yes	2.0
FiHal/Enf/Iso Low	Yes	Yes	0.0
EtSev High	Yes	Yes	6.0
EtSev Low	Yes	Yes	0.0
FiSev High	Yes	Yes	5.0
FiSev Low	Yes	Yes	0.0
EtDes High	Yes	Yes	8.0
EtDes Low	Yes	Yes	0.0
FiDes High	Yes	Yes	6.0
FiDes Low	Yes	Yes	0.0

C.2 Routine Configuration

C.2.1 Alarm

Item Name	Configurable		Default	
nem name	In Config Mode	In Monitor Mode	Delaut	
			General: 2	2
Alm Volume	Yes	Yes	OR:	1
			ICU, NICU, CCU:	2
Reminder Vol	Yes	Yes	Low	
Recording Length	Yes	Yes	16 s	
Arman Dalay	Vec	Vec	Adu, Ped:	20s
Apnea Delay	Yes	Yes	Neo:	15 s
Alarm Delay	Yes	Yes	6 s	
VTac Latching	No	Yes	On	

C.2.2 Screens

Idama Nama		Configurable		Defeat
Item Name		In Config Mode	In Monitor Mode	— Default
Choose Screen		Yes	Yes	Normal Screen
	1			ECG1
	2			ECG2
Colort Mayo Comunition	3	Yes	Yes	SpO ₂ +PR
Select Wave Sequence for Normal Screen	4			Any IBP
for Normal Screen	5			Any IBP
	6			CO ₂
	7			Resp
	Parameter 1			ECG
Select Parameters for Big Numerics Screen	Parameter 2	Vaa	Vec	SpO ₂ +PR
	Parameter 3	Yes	Yes	Resp
	Parameter 4			NIBP

Item Nam	ne	Select QuickKeys		
O.M C		Yes		
0.101	М	No		
General		NIPD Massura - Stop All - Zaro IPD - Davian - Stap day - Scroops - Dationt Satur -		
OR		NIBP Measure \rightarrow Stop All \rightarrow Zero IBP \rightarrow Review \rightarrow Standby \rightarrow Screens \rightarrow Patient Setup \rightarrow		
ICU		Manual Event → Realtime Print → Volume Setup		
NICU		NIBP Measure \rightarrow Stop All \rightarrow oxyCRG \rightarrow Review \rightarrow Standby \rightarrow Screens \rightarrow Patient Setup \rightarrow		
		Manual Event→Realtime Print→Volume Setup		
сси		NIBP Measure→Stop All→Zero IBP→Review→Standby→Screens→Patient Setup→		
		Manual Event→Realtime Print→Volume Setup		
User Defa	ults			

C.2.3 Parameter/Wave Color

Item Name	Configurable		Default
	In Config Mode	In Monitor Mode	Delaut
ECG			Green
NIBP			White
SpO ₂			Cyan
SpO ₂ b	No	Yes	Purple
$\Delta \text{ SpO}_2$			Yellow
PR			Cyan
ТЕМР			White
Art/Ao/UAP/FAP			Red
/BAP/LV/P1~P4 (arterial pressure)			neu
PA			Yellow
CVP/ICP/P1~P4 (venous pressure)			Blue

In the Name	Configurable		Defeat
Item Name	In Config Mode	In Monitor Mode	Default
LAP			Purple
RAP			Orange
UVP			Cyan
CO ₂			Yellow
RESP			Yellow
АА			Yellow
N ₂ O			Blue
O ₂			Green
Hal			Red
Enf			Orange
Iso			Purple
Des			Cyan
Sev]		Yellow
C.O.			White

X represents a waveform label, such as ECG, RESP, CO₂ and so forth. The ECG waveform cannot be set off.

C.2.4 Review

Item Name		Configurable		– Default
		In Config Mode	In Monitor Mode	Delault
				General: 30 min
Tabular Trends	Interval	No	Yes	OR: 5 min
				ICU, NICU, CCU: 30 min
	Trend Group	No	Yes	Standard
	Trend Group	No	Yes	Standard
Graphic Trends	Zoom	No	Yes	90 min
	Waves	No	Yes	2
Minitrend Lengt	h	No	Yes	2 h
	Save Waves	No	Yes	Save ECG1 by default.
Full Disclosure	Gain	No	Yes	x 1
	Sweep	No	Yes	25 mm/s

C.2.5 Event

Item Name	Configurable		Default	
	In Config Mode	In Monitor Mode	Delault	
Waveform 1	No	Yes	Ш	
Waveform 2	N ₋	Yes	General, OR, ICU, CCU: I	
	No		NICU Pleth	
Waveform 3 No	Yes	General, OR, ICU, CCU: Pleth		
		les	NICU Resp	

C.2.6 Record

Item Name		Configurable		Default
		In Config Mode	In Monitor Mode	Derault
Length		No	Yes	8 s
Interval		No	Yes	Off
Paper Speed		No	Yes	25 mm/s
IBP Overlap		No	Yes	Off
Alm Rec	Х	No	Yes	Off

X represents a parameter label.

C.2.7 Print

	Item Name			
		In Config Mode	In Monitor Mode	— Default
Paper Size		No	Yes	Letter
Print On Both Side	5	No	Yes	Off
	Amplitude	No	Yes	10 mm/mV
ECG Reports	Sweep	No	Yes	25 mm/s
	Auto Interval	No	Yes	Off
	Set as End Case Report	No	Yes	Unselected
	Back	No	Yes	Auto
	Spacing	No	Yes	Auto
Tabular Trands	Report Format	No	Yes	Parameter Oriented
Tabular Trends Reports	Currently Displayed Trended Parameters	No	Yes	Selected
	Standard Parameter Group	No	Yes	Unselected
	Custom	No	Yes	Unselected
	Not Print Blank Pages	No	Yes	Selected
	Set as End Case Report	No	Yes	Unselected
Graphic Trends	Back	No	Yes	Auto
Reports	Paginal Tie	No	Yes	Auto
	Set as End Case Report	No	Yes	Unselected
Realtime Reports	Sweep	No	Yes	Auto
	Select Wave	No	Yes	Current

C.2.8 Night Mode

Item Name	Configurable		Default
item Name	In Config Mode	In Monitor Mode	Delaut
Brightness	Yes	Yes	1
Alm Volume	Yes	Yes	2
QRS Volume	Yes	Yes	1
Key Volume	Yes	Yes	0
NIBP End Tone	Yes	Yes	Off

C.2.9 Others

Item Name		Configurable		Default
		In Config Mode	In Monitor Mode	Delauit
Brightness		No	Yes	5
Key Volume		No	Yes	2
View Other Patient	Auto Alarm	No	Yes	On

C.3 User Maintenance Items

	Configurable		
Item Name	In Config Mode	In Monitor Mode	— Default
Changing Pat.Loc.	No	Yes	Protected
Barometric Pressure	No	Yes	760 mmHg
Height Unit	No	Yes	cm
Weight Unit	No	Yes	kg
Press. Unit	No	Yes	mmHg
CVP Unit	No	Yes	cmH ₂ O
CO ₂ Unit	No	Yes	mmHg
O ₂ Unit	No	Yes	%
Temp Unit	No	Yes	°C
Network Type	No	Yes	LAN
Address Type	No	Yes	Manual
Latching Alarms	No	Yes	Lethal
Alarm Pause Time	No	Yes	2 min
Minimum Alarm Volume	No	Yes	General, ICU, NICU, CCU: 2
Minimum Alarm volume			OR: 1
Alarm Sound	No	Yes	ISO
High Alarm Interval (ISO mode)	No	Yes	10
Medium (ISO mode)	No	Yes	20
Low Alarm Interval (ISO mode)	No	Yes	20
Reset Other Bed's Alarms	No	Yes	Off
Alarm Reset By Other Bed	No	Yes	On
Other Bed Disconnection Alm	No	Yes	On
Reminder Interval	No	Yes	3 min
Reminder Tone	No	Yes	On
ECGLeadOff Lev.	No	Yes	Low
SpO ₂ SensorOff Lev.	No	Yes	Low
IBPSensorOff Lev.	No	Yes	Med
Lethal Arrh. OFF	No	Yes	Disable
Extended Arrh.	No	Yes	Enable
Alarm Light on Alarm Reset	No	Yes	On
Alarm Delay	No	Yes	6 s
Vtac Latching Lock	Yes	Yes	Enable

In any Name	Item News			Default
Item Name		In Config Mode	In Monitor Mode	- Default
Wave Line		No	Yes	Med
DIAP Baud Rate		No	Yes	9600
ECG Standard		No	Yes	АНА
Notch Freq.		No	Yes	50 Hz
Data Transfer Method		No	Yes	Off
Transferred Data Lo	Transferred Data Length		Yes	4h
Apply Module Sett	Apply Module Settings		Yes	Off
Parameter Switch		Yes	Yes	Selected
SpO ₂ Tone		No	Yes	Mode 1
	Signal Type	No	Yes	Continuous
Nurse Call Setup	Contact Type	No	Yes	Normally Closed
	Alm Lev	Yes	Yes	High, Med, Low
	Alarm Cat.	Yes	Yes	Phy., Tech.

FOR YOUR NOTES

This chapter lists only the most important physiological and technical alarm messages. Some messages appearing on your monitor may not be included.

In this chapter:

- The "I" column indicates how indications of technical alarms perform after the alarm system is reset: "A" means that some technical alarms are cleared; "B" indicates that some technical alarms are changed to the prompt messages; and "C" indicates that a " √" appears before the alarm message, appears in the alarm symbol area, and the indication of the alarm lamp depends on the alarm light setting. Refer to section **7.8 Resetting Alarms** for details.
- The "Level" field indicates the alarm level: H means high, M means medium and L means low. "*" means the alarm level is user-adjustable.
- XX represents a measurement or parameter label, such as ECG, NIBP, HR, PVCs, RR, SpO₂, PR, etc.

In the "Cause and Solution" column, corresponding solutions are given instructing you to troubleshoot problems. If the problem persists, contact your service personnel.

Measurement	Alarm messages	L	Cause and solution			
	XX Too High	M*	XX value has risen above the high alarm limit or fallen below the low			
ХХ	XX Too Low	M*	alarm limit. Check the patient's condition and check if the patient category and alarm limit settings are correct.			
	ECG Weak Signal	н	The ECG signal is too weak for the monitor to perform ECG analysis. Check the patient's condition and the ECG connections.			
	Asystole	Н				
	VFib/VTac	Н				
	Vtac	Н				
	Vent. Brady	Н				
566	Extreme Tachy	Н				
ECG	Extreme Brady	Н	Arrhythmia has occurred to the patient. Check the patient's			
	R on T	M*	condition and the ECG connections.			
	Run PVCs	M*				
	Couplet	M*				
	PVCs/min	M*				
	Bigeminy	M*				
	Trigeminy	M*				

D.1 Physiological Alarm Messages

Measurement	Alarm messages	L	Cause and solution
	Tachy	M*	
	Brady	M*	
	Missed Beats	M*	
	Afib	M*	
	Irr. Rhythm	M*	
	Vent. Rhythm	M*	
	Multif. PVC	M*	
	Nonsus. Vtac	M*	
	Pause	M*	
	Pacer Not Pacing	M*	The paper appears appermal Check the paper
	Pacer Not Capture	M*	The pacer appears abnormal. Check the pacer.
Resp	Resp Apnea	н	The respiration signal was too weak for the monitor to perform respiration analysis. Check the patient's condition and the Resp connections.
	Resp Artifact	н	The patient's heartbeat has interfered with his respiration. Check the patient's condition and the Resp connections.
6-0	SpO ₂ Desat	н	The SpO_2 value has fallen below the desaturation alarm limit. Check the patient's condition and check if the alarm limit settings are correct.
SpO ₂	No Pulse	н	The pulse signal was too weak for the monitor to perform pulse analysis. Check the patient's condition, SpO_2 sensor and measurement site.
CO ₂	CO ₂ Apnea	н	The patient stops breathing, or the respiration signal was too weak
	AG Apnea	Н	for the monitor to perform respiration analysis. Check the patient's condition and the RM connections.
AG	FiO ₂ Too Low	Н	Check the patient's condition, the ventilated O_2 content and the AG connections.

Measurement	Alarm message	L	I	Cause and solution
	XX SelfTest Err	н	С	
	XX Init Err	н	А	An error occurred to the XX module, or there is a
	XX Init Err N	н	Α	problem with the communications between the
	N is within 1 to 8			module and the monitor. Re-plug the module and restart the monitor, or plug the module into another
	XX Comm Err	Н	Α	monitor.
XX	XX Comm Stop	н	С	
	XX Limit Err	L	с	XX parameter limit is accidentally changed. Contact your service personnel.
	XX Overrange	L	с	The measured XX value is not within the specified range for XX measurement. Contact your service personnel.
	ECG Lead Off	L*	В	
	ECG XX Lead Off	L*	В	The electrode has become detached from the patient or the lead wire has become disconnected from the
	Note: XX represents the leadwires, V, LL, LA, RA, as per AHA standard or C, F, L and R as per IEC standard.			adapter cable. Check the connections of the electrodes and leadwires.
	ECG Noisy	L	A	The ECG signal is noisy. Check for any possible sources of signal noise around the cable and electrode, and check the patient for great motion.
ECG	ECG High Freq. Noise	L	A	High frequency signals are detected on the ECG analysis lead. Check for any possible source of interference around the cable and electrode.
	ECG Low Freq. Noise	L	A	Low frequency signals are detected on the ECG analysis lead. Check for any possible source of interference around the cable and electrode.
	ECG Amplitude Too Small	L	с	The ECG amplitude didn't reach the detected threshold. Check for any possible source of interference around the cable and electrode.
	ECG Config. Err	L	с	ECG configuration is wrongly downloaded. Check the downloaded configuration and re-download the correct configuration.
Resp	Resp Disturbed	L	A	The respiration circuit is disturbed. Restart the monitor.
	Temp Cal. Err	Н	С	A calibration failed. Restart the monitor.
Temp	T1 Sensor Off	L	Α	The Temp sensor has become detached from the
	T2 Sensor Off	L	Α	patient or the module. Check the sensor connections.
6.0	SpO₂ Sensor Off	L*	В	The SpO ₂ sensor has become detached from the
SpO ₂	SpO ₂ Sensor Fault	L	С	patient or the module, or there is a fault with the SpO_2

D.2 Technical Alarm Messages

cPM 12/cPM 8 Operator's Manual

Measurement	Alarm message	L	Т	Cause and solution
	SpO ₂ No Sensor	L	В	sensor, or an unspecified SpO ₂ sensor has been used.
	SpO ₂ Unknown Sensor	L	С	Check the sensor application site and the sensor type,
	SpO ₂ Sensor Incompatible	L	С	and make sure if the sensor is damaged. Reconnect the sensor or use a new sensor.
	SpO_2 Too Much Light	L	с	There is too much light on the SpO_2 sensor. Move the sensor to a place with lower level of ambient light or cover the sensor to minimize the ambient light.
	SpO ₂ Low Signal	L	С	The SpO ₂ signal is too low or too weak. Check the
	SpO ₂ No Pulse	L	с	patient's condition and change the sensor application site. If the error persists, replace the sensor.
	SpO₂ Interference	L	с	The SpO $_2$ signal has been interfered. Check for any possible sources of signal noise around the sensor and check the patient for great motion.
	SpO₂ Comm Abnormal	н	A	An error occurred to the SpO ₂ measurement module, or there is a problem with the communications between the module and the monitor. Restart the monitor. If the error remains, contact your service personnel.
	SpO ₂ Board Fault	L	с	There is a problem with the SpO ₂ measurement board. Do not use the module and contact your service personnel.
	NIBP Loose Cuff	L	А	The NIBP cuff is not properly connected, or there is a
	NIBP Air Leak	L	А	leak in the airway.
	NIBP Pneumatic Leak	L	А	Check the NIBP cuff and pump for leakages.
	NIBP Cuff Type Wrong	L	A	The cuff type applied mismatches the patient category. Verify the patient categor and replace the cuff.
	NIBP Air Pressure Err L A monitor application site requirements and check	An error occurred to the air pressure. Verify that the monitor application site meets the environmental requirements and check if there is any source that affects the air pressure.		
NIBP	NIBP Weak Signal	L	A	The patient's pulse is weak or the cuff is loose. Check the patient's condition and change the cuff application site. If the error persists, replace the cuff.
	NIBP Signal Saturated	L	А	The NIBP signal is saturated due to excess motion or other sources.
	NIBP Overrange	L	А	The measured NIBP value is not within the specified range.
	NIBP-XX Over Upper Limit	L	А	The measured pressure is greater than the specified NIBP measurement upper limit.
	NIBP-XX Over Lower Limit	L	А	The measured pressure is lower than the specified NIBP measurement lower limit.
	XX represents diastolic pressure	e, or systolic pressure.		

Measurement	Alarm message	L	I	Cause and solution
	NIBP Excessive Motion	L	A	Check the patient's condition and reduce thepatient motion.
	NIBP Cuff Overpress.	L	A	The NIBP airway may be occluded. Check the airway and measure again.
	NIBP Equip Err	н	А	An error occurred during NIBP measurement and
	NIBPTimeout	L	А	therefore the monitor cannot perform analysis
	NIBP Measure Failed	L	A	correctly. Check the patient's condition and NIBP connections, or replace the cuff.
	NIBP Illegally Reset	L	A	An illegal reset occurred during NIBP measurement. Check if the airway is occluded.
	XX Sensor Off	M*	А	Check the sensor connection and reconnect the sensor.
IBP	XX Disconnected	н	С	The liquid way is disconneted from the patient, or the three-way valve is open to the air. Check the connection of the liquid way, or check the valve is open to the patient. If the problem remains, contact the Customer Services Dept. for help.
	XX Sensor Fault	М	С	Replace the sensor.
	XX No Pulse	L	А	The catheter may be occluded. Please flush the
	XX represents an IBP label.			catheter.
C.O.	TB Sensor Off	L	А	Check the sensor connection and reconnect the sensor.
	CO ₂ Sensor High Temp	L	С	Check, stop using or replace the sensor.
	CO ₂ Sensor Low Temp	L	С	Check, stop using or replace the sensor.
	CO ₂ Temp Overrange	L	с	The operating temperature of the CO2 module goes beyond the specified range. After it restores within the specified range, the module will restart automatically.
	CO ₂ Airway High Press.	L	С	An error occurred in the airway pressure. Check the
	CO ₂ Airway Low Press.	L	С	patient connection and patient circuit, and then restart the monitor.
	CO ₂ High Barometric Press.	L	С	Check the CO_2 connections, make sure that the
CO ₂	CO ₂ Low Barometric Press.	L	С	monitor application site meets the requirements, and check for special sources that affect the ambient pressure. Restart the monitor.
	CO ₂ FilterLine Occluded	L	С	The airway or watertrap was occluded. Check the airway and remove the occlusion.
	CO ₂ No Watertrap	L	В	Check the watertrap connections.
	CO ₂ Check Adapter	L	A	There is a problem with the airway adapter. Check, clean or replace the adapter.
	CO ₂ FilterLine Err	L	С	Check if there is a leak in the CO_2 sample line or the CO_2 sample line has been occluded.

Measurement	Alarm message	L	Т	Cause and solution
	CO_2 Zero Failed	L	A	Check the CO ₂ connections. After the sensor's temperature becomes stabilized, perform a zero calibration again.
	CO ₂ System Err	L	Α	Re-plug the module or restart the monitor.
	CO_2 Check Cal.	L	С	Perform a calibration.
	CO ₂ Check Airway	L	С	An error occurred to the airway.
	CO ₂ No Filterline	L	Α	Make sure that the filterline is connected.
	CO ₂ No Sensor	L	Α	Make sure that the sensor is connected.
	CO ₂ Main Board Err	Н	С	
	CO ₂ Checking Sensor	L	С	
	CO ₂ Replace Scrubber&Pump	L	С	There is a problem with the CO_2 module. Re-plug the module or restart the monitor.
	CO ₂ 15V Overrange	н	С	module of restart the monitol.
	CO ₂ Hardware Err	н	С	
	AG No Watertrap	L	В	Check the connections of the watertrap and re-connect it.
-	AG Change Watertrap	L	Α	Wait until the change is completed.
	AG Watertrap Type Wrong	L	Α	Make sure that a correct watertrap has been used.
	O ₂ Accuracy Unspecified	L	Α	
	N ₂ O Accuracy Unspecified	L	Α	
	CO ₂ Accuracy Unspecified	L	Α	
	Enf Accuracy Unspecified	L	Α	
AG	Iso Accuracy Unspecified	L	Α	The measured value has exceeded the specified
710	Sev Accuracy Unspecified	L	Α	accuracy range.
	Hal Accuracy Unspecified	L	Α	
	Des Accuracy Unspecified	L	Α	
	awRR Accuracy Unspecified	L	Α	
	AG Hardware Err	н	А	Remove the AG module. Stop using the module and contact your service personnel.
	AG Airway Occluded	L	Α	Check the airway and remove the occlusion.
	AG Zero Failed	L	А	Re-plug the module or restart the monitor, and then perform a zero calibration again.
	12V Too High	н	С	
	12V Too Low	н	С	
	5V Too High	н	С	There is a problem with the system power supply.
Power	5V Too Low	н	С	Restart the monitor.
	3.3V Too High	н	С	
	3.3 Too Low	н	С	

Measurement	Alarm message	L	I	Cause and solution
	Battery Too Low	н	с	Connect the monitor to an AC power source and allow the batteries to charge.
	Different Battery Voltages	м	с	The two batteries have different charge capacity, or the batteries unspecified have been used, or there is a problem with the batteries. Make sure that correct batteries are used and the batteries are not damaged, or replace the batteries.
	Battery Power Overload	н	с	The power consumption of the equipment is too high. Power the monitor using an AC power source.
	RT Clock Not Exist	н	С	Contact your service personnel.
	Recorder Init Err N	L	А	
	N is within 1 to 8.	•	•	Restart the monitor.
	Recorder SelfTest Err	L	А	
	Recorder Comm Err	L	Α	
	Recorder S. Comm Err	L	Α	Stop the recording and restart the monitor.
Recorder	Recorder Unavailable	L	А	
	Recorder Vlt High	L	С	An error occurred to the system power supply. Restart
	Recorder VIt Low	L	С	the monitor.
	Recorder Head Hot	L	с	The recorder has been working for too long time. Stop the recording and resume the recording till the recorder's printhead cools down.
	Rec Paper Wrong Pos.	L	Α	Re-load the recorder paper.
	IP Address Conflict	L	Α	Set a new IP address.
	Other Bed Disconnected	L	А	Check network connection.
	PWR interrupted. Check meas. state	L	A	Power supply failed accidently. Check the measurements when the monitor restarts.
	Restoring Last Config. Failed	L	A	Restart the monitor. If the problem persists, there may be an EEPROM failure. Contact your service personnel.
System	Loading Default Config. Failed.	L	A	Restart the monitor. If the problem persists, there may be an EEPROM failure. Contact your service personnel.
	USB Drive Err	М	A	 Disconnect the USB memory and reconnect it properly. If the problem persists, format the USB memory. If the problem still persists, replace the USB drive.
	Storage Card Err	м	С	Restart the monitor. If the problem persists, format the SD card.

Measurement	Alarm message	L	I	Cause and solution
	Storage Card Space Low	L	В	The storage card has abnormal data. Format the storage card.
	USB Drive Space Low	L	A	Delete unnecessary data from the USB memory, or replace the USB memory.

E.1 Symbols

μΑ	microampere
μV	microvolt
-	Microsecond
μs A	
A	ampere
	ampere hour
bpm	beat per minute
bps	bit per second
°C	centigrade
сс	cubic centimeter
cm	centimeter
dB	decibel
DS	dyne second
٥F	fahrenheit
g	gram
GHz	gigahertz
GTT	gutta
h	hour
Hz	hertz
in	inch
kg	kilogram
kPa	kilopascal
L	litre
lb	pound
m	meter
mAh	milliampere hour
Mb	mega byte
mcg	microgram
mEq	milli-equivalents
mg	milligram
min	minute
ml	milliliter
mm	millimeter
mmHg	millimeters of mercury
cmH2O	centimeters of water
ms	millisecond
mV	millivolt
mW	milliwatt
MΩ	megaohm
nm	nanometer

rpm	breath per minute
S	second
V	volt
VA	volt ampere
Ω	ohm
W	watt
-	minus, negative
%	percent
/	per; divide; or
+	plus
=	equal to
<	less than
>	greater than
≤	less than or equal to
2	greater than or equal to
±	plus or minus
×	multiply

E.2 Abbreviations

AaDO ₂	alveolar-arterial oxygen gradient
AAMI	Association for Advancement of Medical Instrumentation
AC	alternating current
ACI	acceleration index
Adu	adult
AG	anaesthesia gas
АНА	American Heart Association
ANSI	American National Standard Institute
Ao	aortic pressure
Art	arterial
aVF	left foot augmented lead
aVL	left arm augmented lead
aVR	right arm augmented lead
awRR	airway respiratory rate
BAP	brachial arterial pressure
BIS	bispectral index
BP	blood pressure
BPSK	binary phase shift keying
BSA	body surface area
ВТ	blood temperature
BTPS	body temperature and pressure, saturated
C.I.	cardiac index
CCI	Continuous Cardiac Index
C.O.	cardiac output

ссо	Continuous Cardiac Output
CaO ₂	arterial oxygen content
CCU	cardiac (coronary) care unit
CFI	cardiac function index
CIS	Clinical Information System
CISPR	International Special Committee on Radio Interference
CMOS	complementary metal oxide semiconductor
CO ₂	carbon dioxide
COHb	carboxyhemoglobin
СР	cardiopulmonary
CPI	cardiac power index
СРО	Cardiac Power Output
CVP	central venous pressure
DC	direct current
Des	desflurane
Dia	diastolic
DIAP	Datascope Improved ASCII Protocol
DPI	dot per inch
dPmx	left ventricular contractility
DVI	digital video interface
DO ₂	oxygen delivery
DO ₂ I	oxygen delivery index
ECG	electrocardiograph
EDV	end-diastolic volume
EEC	European Economic Community
EEG	electroencephalogram
EMC	electromagnetic compatibility
EMG	electromyography
EMI	electromagnetic interference
Enf	enflurane
ESU	electrosurgical unit
Et	end-tidal
EtCO ₂	end-tidal carbon dioxide
EtN ₂ O	end-tidal nitrous oxide
EtO	ethylene oxide
EtO ₂	end-tidal oxygen
EVLW	extravascular lung water
ELWI	extravascular lung water index
FAP	femoral arterial pressure
FCC	Federal Communication Commission
FDA	Food and Drug Administration
FEV1.0%	first second forced expiratory volume ratio
Fi	fraction of inspired
FiCO ₂	fraction of inspired carbon dioxide
FiN ₂ O	fraction of inspired nitrous oxide
FiO ₂	fraction of inspired oxygen

FPGA	field programmable gate array
FV	flow-volume
GEDV	global end diastolic volume
GEDI	global end diastolic volume index
GEF	global ejection fraction
Hal	halothane
Hct	haematocrit
Hb	hemoglobin
Hb-CO	carbon mono-oxide hemoglobin
HbO ₂	oxyhemoglobin
HIS	hospital information system
HR	heart rate
l:E	
	inspiratory-expiratory ratio
IBP	invasive brood pressure
ICG	impedance cardiography
ICP	intracranial pressure
ICT/B	intracranial catheter tip pressure transducer
ICU	intensive care unit
ID	identification
IEC	International Electrotechnical Commission
IEEE	Institute of Electrical and Electronic Engineers
Ins	inspired minimum
IP	internet protocol
lso	isoflurane
IT	injectate temperature
ITBI	Intrathoracic Blood Volume Index
ITBV	Intrathoracic Blood Volume
LA	left arm
LAP	left atrial pressure
Lat	lateral
LCD	liquid crystal display
LCW	left cardiac work
LCWI	left cardiac work index
LED	light emitting diode
LL	left leg
LVD	low voltage directive
LVDS	low voltage differential signal
LVET	left ventricular ejection time
LVSW	left ventricular stroke work
LVSWI	left ventricular stroke work index
MAC	minimum alveolar concentration
Art mean	mean arterial pressure
MDD	Medical Device Directive
MetHb	methemoglobin
MRI	magnetic resonance imaging
MRN	medical record number

MVe	expiratory minute volume
MVi	inspiratory minute volume
N/A	not applied
N ₂	nitrogen
N ₂ O	nitrous oxide
Neo	neonate
NIBP	noninvasive blood pressure
O ₂	oxygen
O ₂ CI	oxygen consumption index
O ₂ R	oxygen extraction ratio
OR	operating room
OxyCRG	oxygen cardio-respirogram
PA	pulmonary artery
pArt-D	diastolic artery pressure
pArt-M	mean artery pressure
pArt-S	systolic artery pressure
Paw	airway pressure
PAWP	pulmonary artery wedge pressure
PD	photodetector
Ped	pediatric
PEEP	positive end expiratory pressure
PEF	peak expiratory flow
PEP	pre-ejection period
PIF	peak inspiratory flow
PIP	peak inspiratory pressure
Pleth	plethysmogram
Pmean	mean pressure
Pplat	plateau pressure
PPV	Pulse Pressure Variation
PR	pulse rate
PVC	premature ventricular contraction
PVR	pulmonary vascular resistance
PVRI	pulmonary vascular resistance index
PVPI	pulmonary vascular permeability index
pArt	artery pressure
pCVP	central venous pressure
R	right
RA	right arm
RAM	random access memory
RAP	right atrial pressure
Rec	record, recording
Resp	respiration
RHb	reduced hemoglobin
RL	right leg
RM	respiratory mechanics
RR	respiration rate

RSBI	rapid shallow breathing index
SaO ₂	arterial oxygen saturation
SEF	spectral edge frequency
Sev	sevoflurane
SFM	self-maintenance
SI	stroke index
SMR	satellite module rack
SpO ₂	arterial oxygen saturation from pulse oximetry
SQI	signal quality index
SR	suppression ratio
STR	systolic time ratio
SV	stroke volume
SVI	Stroke Volume Index
SVR	systemic vascular resistance
SVRI	systemic vascular resistance index
SVV	stroke volume variation
SvO ₂	mixed venous oxygen saturation
ScvO ₂	central venous oxygen saturation
Sync	synchronization
Sys	systolic pressure
Taxil	axillary temperature
ТВ	Blood Temperature
TD	temperature difference
Temp	temperature
TFC	thoracic fluid content
TFI	thoracic fluid index
TFT	thin-film technology
Toral	oral temperature
ТР	total power
Trect	rectal temperature
TVe	expiratory tidal volume
TVi	inspiratory tidal volume
UAP	umbilical arterial pressure
UPS	uninterruptible power supply
USB	universal serial bus
UVP	umbilical venous pressure
VAC	volts alternating current
VEPT	volume of electrically participating tissue
VI	velocity index
VO ₂	oxygen consumption
VO ₂ I	oxygen consumption index
WLAN	wireless local area network
WPA	Wi-Fi protected access