OMNI

Patient Monitor

USER'S MANUAL

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SAFETY INFORMATION

This section contains important safety information related to general use of the OMNI monitor. Other important safety information appears throughout the manual in sections that relate specifically to the precautionary information. Read all text surrounding all precautionary information.

The OMNI monitor can be powered by an internal battery pack that provides 1 hour of monitoring from fully charged batteries. The batteries are continuously recharged when AC power is connected to the monitor.

A warning message appears on the screen and an audible alarm sounds when the remaining battery power is only enough for 10 minutes of operation. The user should connect the monitor to an external power source to avoid loss of patient monitoring action.

External power sources may be connected, disconnected and reconnected without interrupting the monitoring action.

The integrity of the external protective conductor in the installation or its arrangement is in doubts; equipment shall be operated from its INTERNAL ELECTRICAL POWER SOURCE.

Important! Before use, carefully read this manual, accessory directions for use.

WARNING: The OMNI monitor is defibrillator proof. It may remain attached to the patient during defibrillation or while an electrosurgical unit is in use, but the readings may be inaccurate during use and shortly thereafter.

WARNING: The OMNI monitor is a prescription device and is to be operated by qualified personnel only.

WARNING: Occasionally, electrical signals at the heart do not produce a peripheral pulse. If a patient's beat-to-beat pulse amplitude varies significantly (for example, pulsus alternans, atrial fibrillation, rapid-cycling artificial ventilator), blood pressure and pulse rate readings can be erratic and an alternate measuring method should be used for confirmation.

WARNING: Explosion hazard. Do not use the OMNI monitor in the presence of flammable anesthetics or gases.

WARNING: Do not lift the OMNI monitor by the sensor cable, blood pressure hose, or power cord because the cable, lead, or cord could disconnect from the monitor, causing the monitor to drop on the patient.

WARNING: The OMNI monitor may not operate effectively on patients who are experiencing convulsions or tremors.

WARNING: To prevent electrical hazards to all personnel, these monitors must be properly grounded. The chassis grounding assembly, Universal Switching Power Supply, and the power cord supplied with the equipment provides for this protection. Do not attempt to defeat this protection by modifying the cords or using ungrounded adapters.

WARNING: Enclosure leakage current is less than 100 microamperes (μ A); however, always consider additional leakage current that can be caused by other equipment used on the patient at the same time as these monitors.

WARNING: For pacemaker patients, the OMNI may continue to count pacemaker rate during occurrences of cardiac arrest or some arrhythmias. To reduce the likelihood of this, ensure that the Pacer Detect setting is ON in the ECG menu when monitoring such patients. Do not rely entirely upon the OMNI alarms. Keep pacemaker patients under close surveillance.

WARNING: To ensure that the leakage current protection remains within the specifications, use only the patient cables supplied with, or specifically intended for use with the OMNI II Monitors.

WARNING: Line isolation monitor transients may resemble actual cardiac waveforms and thus inhibit heart rate alarms. Such transients may be minimized by proper electrode and cable placement, as specified in this manual and electrode directions for use.

WARNING: Disconnect the OMNI monitor and sensors during magnetic resonance imaging (MRI) scanning. Use during MRI could cause burns or adversely affect the MRI image or the monitor's accuracy. Also, to avoid burns, remove the sensors from the patient before conducting MRI.

WARNING: ECG cables may be damaged if they are connected to a patient during defibrillation. Cables that have been connected to a patient during defibrillation should be checked for functionality before using again.

CAUTION:

To touchscreen:

- Clean and soft clothes with neutral detergent and with isopropyl alcohol may be used for cleaning.
- Do not use any chemical solvent, acidic or alkali solution.
- The panel is designed with air groove. Insulation and cushioning pads should be designed around the edges of the panel to prevent water and dust.
- Use a plastic stylus (tip R0.8 or over) or finger. Sharp edged or hard articles are prohibited.
- The gathering of dew in the panel may occur with abrupt temperature or humidity changes. A stable environment condition is recommended.
- Keep the surface clean. No adhesives should be applied.
- Avoid high voltage and static charge.

CAUTION:

When connecting the OMNI monitor to any instrument, verify proper operation before clinical use. Both the OMNI monitor and the instrument connected to it must be connected to a grounded outlet. Accessory equipments connected to this patient monitor must be certified according to the respective IEC standards (e.g. IEC 60950 for information technology equipment and IEC 60601-1 for medical electrical equipment). Furthermore all configurations shall comply with the valid version of the system standard IEC 60601-1-1. Any person who connects additional equipment to the signal input or signal output is responsible to ensure the system complies with the requirements of the valid version of the system standard IEC 60601-1-1. If in doubt, contact our company or customer service.

To ensure accurate readings, consider the environmental conditions that are present and the condition of the patient. See the appropriate sections of the manual for specific safety information related to these conditions.

WARNING:

Defibrillation and Electrosurgery: Do not touch the patient, or table, or instruments, during defibrillation.

After defibrillation, the screen display recovers within 10 seconds if the correct electrodes are used and applied in accordance with the manufacturers instructions.

ECG cables can be damaged when connected to a patient during defibrillation. Check cables for functionality before using them again.

According to AAMI specifications the peak of the synchronized defibrillator discharge should be delivered within 60 ms of the peak of the R wave. The signal at the ECG output on the OMNI III patient monitors is delayed by a maximum of 30 ms. Your biomedical engineer should verify that your ECG/Defibrillator combination does not exceed the recommended maximum delay of 60 ms.

When using electrosurgical (ES) equipment, never place ECG electrodes near to the grounding plate of the ES device, as this can cause a lot of interference on the ECG signal.

INTENDED USE ABOUT THIS MANUAL

INTENDED USE

The OMNI monitor is a comprehensive monitoring system with three or six traces compiling, processing, analyzing and displaying data from up to eight different patient parameters. It integrates parameter measuring modules, display and recorder in one device, featuring in compactness, lightweight and portability. Built-in battery facilitates transportation of patient.

The purpose and function of the OMNI monitor is to monitor ECG, heart rate, NIBP (systolic, diastolic, and mean arterial pressures), SpO_2 , respiration, temperature, dual temperature, EtCO₂ and dual IBP for adult, neonate and pediatric patients in all hospital areas and hospital-type facilities. It may be used during hospital transport and in mobile, land-based environments, such as ambulances.

The OMNI monitor offers advanced features such as an intuitive touchscreen with clinical measurements, one-touch commands, crisp and clear display.

WARNING: The OMNI monitor is intended only as an adjunct in patient assessment. It must be used in conjunction with clinical signs and symptoms.

ABOUT THIS MANUAL

This manual explains how to set up and use the OMNI monitor. Important safety information relating to general use of the OMNI monitor appears before this introduction. Other important safety information is located throughout the text where applicable. **Read the entire manual including the** *Safety Information* section before you operate the monitor.

CONTROLS, INDICATORS AND SYMBOLS FRONT PANEL AND LEFT SIDE PANEL REAR PANEL AND RIGHT SIDE PANEL SYMBOLS

FRONT PANEL AND LEFT SIDE PANEL



Figure 1: Front view for main unit

Table 1: Description for controls on front and side panel of OM	NI
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No.	FUNCTION	lcon
1	ALARM INDICATOR	
	In normal mode, no indicator lights.	
	In alarm mode, the alarm indicator flashes.	
2	POWER SWITCH	- •
	This toggle switch turns the secondary power from on to off from the monitor.	$(\cdot)/\dot{(})$
	The monitor will continue to charge the battery as long as the AC cable is	\bigcirc <i>I</i> \bigcirc
2	plugged in, even if the power switch is in the off station.	
3	DC ON	
4	This LED indicates that the monitor is powered by battery.	
4	AC ON	\sim
	This LED indicates that the monitor is powered by AC.	
5	Oxygen saturation sensor port	
6	EtCO ₂ input port (Option)	
7	NIBP port for the connection with the blood pressure cuff hose	
8	AAMI ECG lead socket	
9	IBP port for channel 1 (Option)	
10	IBP port for channel 2 (Option)	
11	Temperature port for channel 1	
12	Temperature port for channel 2	

REAR PANEL AND RIGHT SIDE PANEL



Figure 2: Rear view for main unit

Table 2: Description for con	trols on rear and	side panel of OMNI
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No.	FUNCTION	lcon
1	Equipotentiality Ground	\forall
2	AC Input The AC power connection is where facility line power is connected to this monitor, the AC power fuses must be replaced with the same type and rating fuse.	100-240V ~ 50/60Hz, 150VA
3	Fixed support for EtCO ₂ module (option)	
4	USB port	•
5	Network Interface	
6	RS-232 I/O This digital interface connector provides serial data to most RS-232 devices	RS232
7	Recorder (option)	

SYMBOLS

The following symbols may appear on the packaging, monitor or in user's manual:

Ť	Type BF applied part
┨╋╋	Defibrillation-proof type CF applied part To identify a defibrillation-proof type CF applied part complying with IEC 60601-1. Note 1 - C = Cardial. Note 2 - F = Floating applied part.
(+/←	Rechargeable battery To indicates the positioning of the cells.
SN	Manufacture's serial number
2X T 3DA 250V	Fuse information
\sim	Date of manufacture
	Manufacturer
∎ ⊥	FRAGILE Contents of the transport package are fragile therefore it shall be handled with care.
<u>†</u> †	THIS WAY UP Indicates correct up right position of the transport package.
Ť	KEEP AWAY FROM RAIN Transport package shall be kept away from rain.
× × × × × × × × ×	STACKING LIMIT BY NUMBER Maximum number of identical packages which may be stacked on one another is eight.
Â	General warning, caution, risk of danger Please read the instructions carefully before operating the product.



Figure 3: Display screen partition diagram



Figure 4: Tree diagram for top button



Figure 5: Tree diagram for bottom button

All TFT display screen is divided into five areas:

PARAMETER AREA

This area is used for display monitoring parameters, such as HR, RESP, SPO2, TEMP, NIBP(SYS, DIA, MAP), P1, P2, ETCO2 and so on.

WAVEFORM AREA

This area is used for displaying waveforms and menu setup interface. The user can use menu to distribute the combination of window waveform and NIBP data list.

The waveforms from top to bottom can be selected from ECG I, ECG II, ECG III, ECG AVR, ECG AVF, ECG AVL, ECG V, PLETH, IBP1, IBP2 and ETCO2. Only a relative module is set to be ON, its waveform can be selected.

MESSAGE AREA

The state messages of Time, patient types and correlated information are list here.

TOUCH KEYS AREA

This area lists the main function touch keys. The user can touch and click any of them to enter relevant function.

MENU AREA

Menu Area is in the waveform area, see below:



FACTORY SETUP

Click item in the SYSTEM SETUP→Factory Setup menu to call up input the password menu.

The passward allowes input total 8 characters, for the less than 8 charecaters one, you can add point to full in. The following table is all password you can use:

1."**DEMO....**": Open/close the demonstration mode for the system;

2."**MAKE....**": Switch on/off for "make" item for the SPO2 function;

3."LANGUAGE": Set the language;

4."SCREEN..": Open/close the calibration mode for the touch screen;

5."IP SETUP": Set Ethernet IP address;

6."**FORMAT..**": Format the NAND flash. This function is used for clear all data which for the patient data is too much in order to vacate space in NAND flash, then save patient data again.

RECORDER SETUP

RECORDER LINKING STATUS

Use to display or set the connecting state of recorder.

RECORDER GRID OUTPUT

Set waveforms and parameters printout has a grid background or not.

ALARM RECORD

If this item set to be **ON**, It can record a slip of waveform of 10 seconds (the preceding 4 seconds before the recording till the current 4 seconds) when an alarm is happened.

HOSPITAL NAME

Click this item to input or change the hospital name.

ALARM LEVEL SETUP

It is used to set the alarm priority to be medium or high.

DEFAULT CONFIG SETUP

You can call the default settings by clicking this item.

After return the above confirmation menu, a message of "LOAD DEFAULT CONFIG DATA SUCCESS!" will display in the message highlight area, showing that the system has begun to use the default settings.

SAVE CONFIG SETup

Save current config settings so that system can call up these settings on the next time of open by clicking this item.

After return the above confirmation menu, a message of will display the message of "SAVE DATA SUCCESS!", showing that the system and all monitoring parameter settings have been saved (see each chapter).

LANGUAGE SETUP

Use to select language for the monitor system. The language can be switched only after inputting the correct password of "language".

DEMO display

Use to display demonstration interface. This can be available only after inputting the correct password of "DEMO....".

DISPLAY MODE

It is used to set the display mode to be "6 waveforms", "3 waveforms", "Large font" or "OxyCRG".

ALARM SUSPEND SETUP

There are four options of 1 minute, 2 minutes, 3 minutes and Permanent.

If select "Permanent" item, there is a warning message of "warning: alarm suspend permanently" displaying on the top message area. And alarm indicator will not flash, there are no any alarm messages and alarm sounds.

SCREEN CALIBRATE SETUP

Servicing engineer uses this setup only when input the correct password of "SCREEN..".

Click the button of **Screen Cal** in the above figure, the system enters screen calibration mode. At this time user actions are invalidated except calibration actions.

Four steps need be followed according to information in the message highlight area. The first threes steps are calibration steps that user click the red cross icon accurately to calibrate touch screen.

The fourth step is calibration verification step that user click the red cross icon accurately to verify touch screen calibration result.

After finishing the screen calibration, the system will return to the normal mode when calibration validate is successful, or return to the first step of calibration when calibration is failed.

HOW TO MONITOR

- 1. According to the parameter needed, connect the correlated sensors to the sockets on the left panel;
- 2. Connect with the power supply, press the power switch in the front panel;
- 3. Power indicator is bright, the display screen enter the main screen after 25 seconds;
- 4. Connect corresponding sensors with the patient;
- 5. Set monitoring parameters (see chapters below) ;
- 6. Enter the monitoring state.

CAUTION: If the OMNI is to be stored for a period of 2 months or longer, notify service personnel to remove the battery from the monitor prior to storage. Recharge the battery when the battery has not been recharged for 2 or more months.

CAUTION: Follow local government ordinances and recycle instructions regarding disposal or recycling of device components, including batteries.

ALARM & SOUND

ALARM

When the monitor detects certain conditions that require user attention, the OMNI monitor enters an alarm state. The monitor response is indicated by:

- Visual alarm indicators
- Audible alarm indicators
- Print-on-alarm (if printer installed)
- Identification of out-of-limit vital signs in trend data

WARNING: Each time the monitor is used, check alarm limits to ensure that they are appropriate for the patient being monitored.

WARNING: Do not silence the audible alarm or decrease its volume if patient safety could be compromised.

ALARM PRIORITY

The monitor's visual and audible responses to a detected alarm depend on the priority of the alarm; High, Medium, or Low.

A higher priority alarm will supersede a lower priority alarm.

The three categories of alarms are summarized in the following paragraphs. The text indicates the message shown on the screen.

High Priority:

Indicating that immediate OPERATOR response is required: Asystole (4 seconds have passed with no heart beats from ECG, preceded by detecting valid ECG-derived heart rate data.) Loss of Pulse from SpO2 (and no valid ECG)

Medium Priority:

Indicating that prompt OPERATOR response is required:

High/Low numeric value limits violated (such as High/Low SpO2 limits violated, High/Low Sys./Dia. blood pressure limits violated, High/Low Respiration Rate limits violated, High/Low Temperature limits violated, etc.)

Low Priority:

indicating that OPERATOR awareness is required:

Senor or leads off (such as ECG Leads Off, SpO2 Cable/Sensor Disconnect, Temperature Probe Disconnect, etc.), Low Battery (alarm commences when the OMNI has at least 10 minutes of operating time remaining) and communications errors for modules.

VISUAL ALARM INDICATORS

When an alarm occurs, the OMNI responds with visual alarm indications. The flashing rates for the three categories of alarms are shown. The OMNI uses flashing colors to indicate high and medium priority alarm as following Flashing Rates.

Alarm Category	Flashing Rate
High Priority	Two flashes in 1 second
Medium priority	One flash in 2 seconds
Low priority	Constant (on) (non-flashing)

When a low priority alarm occurs, a non-flashing alarm message appears in the message area. If more than one low priority alarm is present, the alarm

messages "rotate". On the OMNI numeric frame background color will change to a solid yellow for a low priority alarm.

When a medium priority alarm is activated, because a parameter is above or low its alarm limits, the out-of-limit numeric value and the bell icon in the corresponding Numeric Frame flash at the medium priority rate. Only the numeric frame background color will flash yellow for a medium priority alarm in the OMNI.

When the high-priority Asystole alarm occurs, the heart rate numeric value and the corresponding bell icon flash at the high priority rate. Only the numeric frame background color will flash red for a high priority alarm in the OMNI. A nonflashing Asystole message appears in the message area and will override any other messages which may be present (there is no message "rotation" in this instance).

ALARM SUSPEND

If the Alarm Silence switch is depressed, the Alarm Suspend condition is initiated. Pressing the Alarm Silence switch terminates the Alarm Suspend condition.

When Alarm Suspend is initiated, ALL audible alarms and print-on-alarm functions are disabled. Visual alarms and identification of out-of-limits vital signs in trend memory continue to function.

During Alarm Suspend, monitoring continues for all parameters; the numeric values and the top Graphic Frame (typically ECG waveform) continue to operate normally. Trend memory operates normally. The single-function buttons continue to operate normally. No Level 1 or 2 menus may be invoked while in Alarm Suspend State. Access to the Big Numbers screen is denied while in the Alarm Suspend condition.

SOUND

ALARM SOUND

The mild sound of BEEP. There are four items of I, II, III and IM or alarm levels in turn from low to high.

The following encoded auditory alarm signals categorized by alarm condition and priority:

Alarm Category	Encoded Auditory	
High Priority	ССССС	
Medium priority	CCC	
Low priority	eC	
Note1: The characters c,e refer to relative musical pitches and C is one octave c.		
Note2: A high priority alarm signal is generated with the five pulses shown, repeat		
once, for total of 10 pulses.		

HEART-BEAT (PULSE-TONE)

The heart-beat or pulse-tone is a sound of RUB-A-DUB. In the ECG setup menu, there are QRS, PULSE and OFF for choice, when the choice is QRS, the system will sound by heart-beat sound. When the choice is PULSE, the system will sound by pulse-tone sound. When the choice is OFF, the system will close the heart-beat

sound or pulse-tone ..

KEY BEEPS

The key beep sounds come along with clicking function items.

SILENCE

Click this function button to disable all sounds except for the key beeps. A symbol of

displays in the message area, click this button again to restore all sounds except for the key beeps.

ALARM SWITCH

When any alarm switch is set to be $\ensuremath{\mathsf{OFF}}$, the alarm indicator will not light, the relative

alarm parameter will not flash and relative parameter area will appear an icon of

ECG MONITORING

ELECTRODE INSTALLATION SENSOR INSTALLATION ECG SETUP ERROR MESSAGES OF ECG MONITORING MAINTENANCE AND CLEANING

ELECTRODE INSTALLATION

Some points should be paid attention to in ECG monitoring:

- 1. Check the lead and cable, the damaged or ruptured one cannot be used.
- 2. Link up the lead set and cable, and connect the electrode to the lead.
- 3. Choose the suitable skin at which the electrode should be pasted. Use alcohol to clean the skin and remove the skin grease. Paste the electrode on the patient and check that whether they are contact well.
- 4. Follow the methods below to place these 5 –lead electrode, it can be set by the ECG menu.



□ WHITE (RIGHT ARM) ELECTRODE (RA)—is placed near the right shoulder, directly below the clavicle.

□ BLACK (LEFT ARM) ELECTRODE (LA)—is placed near the left shoulder, directly below the clavicle.

□ GREEN (REFERENCE) ELELCTEODE (RL)—is placed on the right hypogastrium.

- □ RED (LEFT LEG) ELELCTEODE (LL)—is placed on the left hypogastrium.
- BROWN(CHEST)ELECTRODE(V or C)-be placed on the chest as illustrated below:

NOTE:

- □ Only the ECG cable presented by our factory can be used.
- □ To ensure patient safety, all leads must be attached to the patient.

For 5-lead set, attach the C-electrode to one of the indicated positions as below:



- \Box V1 is on the 4th intercostal space at the right sterna margin.
- □ V2 is on the 4th intercostal space at the left sterna margin.
- \Box V3 is at the midway between V2 and V4 electrodes.
- □ V4 is on the 5th intercostal space at the left clavicular line.
- □ **V5** is on the left anterior axillary line, horizontal with V4 electrode.
- □ **V6** is on the left middle axillary line, horizontal with V4 electrode.
- □ V3R-V7R is on the right side of the chest in positions corresponding to those on the left.
- \Box VE is over the xyphoid. As for the V-lead position on the back, it should be placed at one of the positions below.
- □ **V7** is on the 5th intercostals space at the left posterior axillary line of back.
- □ **V7R** is on the 5th intercostals space at the right posterior axillary line of back.
- 5. The electrodes must be moved away to check the skin every 24 hours, if the skin is found inflamed of damaged evidently, substituted a new electrode to another position.
- 6. The gain choice of ECG is 0.5, 1.0 and 2.0.
- 7. Make sure no conductive part of electrodes is in contact with the ground and other conductive.

SENSOR INSTALLATION

- 1. Insert the plug of ECG into socket on the left panel of monitor, make sure that the salient of plug must direct to the notch of socket when inserting.
- 2. Connect the electrode lead to the patient's cable.

ECG SETUP





Figure 7: Tree diagram for ECG setup menu

LEAD TYPE

Set the lead type for "5 lead" or "3 lead".

If select 3 lead type, the HR calculate and ECG waveform only can be from one of lead I, lead II and lead III.

If select 5 lead type, the ECG waveforms can be from lead I, II, III, aVR, aVF, AvI and V. But the HR calculates according to lead I, lead II or lead V.

ECG GAIN

The user can freely choose one from items of **X0.25**, **X0.5**, **X1.0**, **X2.0**. Gain adjustment can change the value of ECG waveform and ECG STAFF, and the factory-set is **X 1.0**.

ECG FILTER

Select between ON and OFF, and the factory-set is OFF.

NOTE: On conditions that the interference to ECG waveform is too large or using at operating room, FILTER SWITCH is suggested to set to be **ON**.

HEART SOUND

There are QRS, PULSE and OFF for choice, the factory-set is QRS.

There are QRS, PULSE and OFF for choice, when the choice is QRS, the system will sound by heart-beat sound. When the choice is PULSE, the system will sound by pulse-tone sound. When the choice is OFF, the system will close the heart-beat sound or pulse-tone.

HR CALCULATE SOURCE

Select between lead I, lead II and V, and the factory-set is lead II.

SWEEP SPEED

From 12.5mm/s, 25mm/s and 50mm/s for choice, the factory-set is 25mm/s.

SWEEP COLOR

From white, gray, red, yellow, green, cyan, blue, magenta and default for choice, the factory-set is default.

ST ANALYSIS

It is used to complete Automatic ST-segment analysis function. Refer to the below detailed chapter.

ARRHYTHMIA ANALYSIS

It is used to complete Automatic arrhythmia analysis function. Refer to the below detailed chapter.

ERROR MESSAGES OF ECG MONITORING

Sometimes messages will display above the ECG waveform:

PROMPTS	EXPLANATION
Lead off	ECG leads fall off the skin or the
	monitor
ECG Signal Weak	ECG Signal is Weak

MAINTENANCE AND CLEANING

PATIENT CABLE AND LEAD

After every times of using, the cable must be cleaned and following the methods below: 1. Clear the paste on body and the remainder of electrolyte on the electrode. The paste-eradicator can be used when removing the adhesive tape remainder, but acetone, alcohol, ammonia, chloroform and other strong solvent are not suggested, because they would finally damage the vinyl cable.

2. Use a sponge moistened in mild soap liquid or other suitable detergent solution to clean the cable and then dry them. Do not submerge the cable into the water.

3. Check each cable to see whether they are corroded, damaged or degenerated. Do not Use pressure cooker to disinfect the cable and electrode or heat them to 7°C (167F) and higher temperature. If there is dirt on the material surface, you can use the abluent which will not left remainder to clean and any metal grinding medium like floss is forbidden. The storing temperature should be -20°C till 75°C 68F till 167F). Hang or place them flat so as not to be damaged.

ADDING POINTS

- 1. HR calculating stability has a process, ECG lead switching sometimes affect HR which will become stable after a while. The change of gain and filter may influent the HR calculating stability too. Another factor which affecting HR calculation is the QRS waveform, if T wave is too high, HR will be make mistake too. Arrhythmia sometimes influent HR calculation too.
- 2. Choosing suitable ECG waveform range and complete QRS waveform has important effect in the accuracy of HR calculation.

RESP MONITORING

RESP ELECTRODE INSTALLATION RESP SETUP MAINTENANCE AND CLEANING

RESP ELECTRODE INSTALLATION

The monitor measures respiration from the amount of thoracic impedance between two ECG electrodes. The change of impedance between the two electrodes, (due to the thoracic movement), produces a respiratory waveform on the screen.

For RESP monitoring, it is not necessary for additional electrodes, however, the placing of electrodes is important.

Some patients, due to their clinical condition, expand their chest laterally, causing a negative intrathoracic pressure. In these cases it is better to place the two RESP electrodes laterally in the right axillary and left lateral chest areas at the maximum point of breathing movement to optimize the respiratory waveform.

The sensor of RESP ELECTRODE's installation is same as ECG's.

NOTE:

The RESP monitoring is not recommended to be used on patients who are very active, as this can cause **false alarms**.

The scheme picture for placing the 5 Electrodes for respiratory monitoring is seen as followings:



NOTE:

Place the red and green electrodes diagonally to optimize the respiration waveform. Avoid the liver area and the ventricles of the heart in the line between the RESP electrodes so as to avoid cardiac overlay or artifacts from pulsating blood flow. This is particularly important for neonates.

RESP SETUP



Figure 8: Tree diagram for resp setup menu

The menu can finish settings as below:

ALARM SWITCH

ON and OFF for choice, the factory -set is OFF.

DISPLAY PARAMETER

The **ON** and **OFF** for choice. select **ON** can display RESP rate, select **OFF** would not display the RESP, but this do not influent the actual data of trend.

NOTE: When the patient's thorax or abdomen is subjected too much interference, the RESP monitoring is not accurate, so it is suggested to close the RESP rate display.

SWEEP SPEED

Choose from 12.5mm/s to 25.0mm/s, and the factory-set is 12.5mm/s.

RESP ALARM HIGHER-LIMIT

The RESP alarm upper-limit, the range is from **6** to **120** bpm, and the factory-set is **30** bpm, the single-step adjustable step- length is **1** bpm.

RESP ALARM LOWER-LIMIT

The RESP alarm lower-limit, the range is from **3** to **120** bpm, and the factory-set is **3** bpm, the single-step adjustable step- length is **1** bpm.

MAINTENANCE AND CLEANING

It is the same as the ECG monitoring.

SPO2 MONITORING

SPO2 MONITORING PRINCIPLE SPO2 SENSOR INSTALLATION SPO2 PARAMETER SETUP SPO2 PARAMETER SETUP MEASUREMENT LIMITATIONS SPO2 ERROR MESSAGES

SPO2 MONITORING PRINCIPLE

Arterial oxygen saturation is measured by a method called pulse oximetry. It is a continuous, non-invasive method based on the different absorption spectra of reduced hemoglobin and oxyhemoglobin. It measures how much light, sent from light sources on one side of the sensor, is transmitted through patient tissue (such as a finger or an ear), to a receiver on the other side.

The amount of light transmitted depends on many factors, most of which are constant. However, one of these factors, the blood flow in the arteries, varies with time, because it is pulsating. By measuring the light absorption during a pulsation, it is possible to derive the oxygen saturation of the arterial blood. Detecting the pulsation gives a PLETH waveform and pulse rate signal.

About SpO₂, SaO₂, SjvO₂

SpO₂: It is the arterial blood oxygen saturation lever measuring by oximeter.

□SaO₂: It is the oxygen saturation of arterial blood

SjvO₂: It is the oxygen saturation of the jugular blood.

WARNING: Pulse oximeter can overestimate the SPO2 value in the presence of HB-CO, Met-HB or dye dilution chemicals.

SPO2 SENSOR INSTALLLATION

1. Insert the plug of SPO2 sensor into the **SPO2** socket on the left panel of monitor. Make sure that the salient of plug must direct to the notch of socket when inserting of unplugging, otherwise the measurement will not be reliable and the sensor connector will be damaged.

2. Wear the finger-probe on the finger; make sure that the finger tip is the same direction as the finger direction indicated on the probe.

SPO2 PARAMETER SETUP



Figure 9: Tree diagram for SPO2 setup menu

ALARM SWITCH

ON and OFF for choice, the factory -set is OFF.

ALARM RECORD

ON and OFF for choice, the factory -set is OFF.

If it is the **ON** choice, the recorder can automatically print the current wave and each parameter value when occurring parameter alarm.

WAVEFORM SPEED

Choose from **12.5mm/s** to **25.0mm/s**, and the factory-set is **12.5mm/s**.

SPO2 ALARM UPPER-LIMIT

The SPO2 alarm upper-limit, the range is from **70** to **100**%, and the factory-set is **100%**, the single-step adjustable step- length is **1** %.

SPO2 ALARM LOWER-LIMIT

The SPO2 alarm lower-limit, the range is from **70** to **150**%, and the factory-set is **85%**, the single-step adjustable step- length is **1**%.

MEASUREMENT LIMITATIONS

- 1. The measurement is decided by the pulsating characteristic of blood stream in artery or arterial blood vessel. The arterial blood stream may decreased to the level which cannot be measured in conditions below:
 - Shock
 - Hypothermia
 - □ Vasoactive medicines are applied
 - Anemia

- 2. The measurement are also decided by the condition how the oxyhemoglobin and reduced-hemoglobin absorb the light of special wave-length. If there are other material can absorb the same wave-length light, they can cause the measurement false or lower than the actual value of SPO2, for example:
 - Carboxyhemoglobin
 - Methemoglobin
 - □ Methylene blue
 - Carmine indigo
- 3. The strong light in the environment also can influent measurement. Some suitable light-tight material to cover the sensor which can improve the measure quality.

WARNING:

- □ Prolonged and continuous monitoring may increase jeopardy of unexpected change of dermal condition such as abnormal sensitivity, rubescence, vesicle, repressive putrescence, and so on. It is especially important for neonate and patient of poor perfusion or immature dermogram to check the sensor placement by light collimation and proper attaching strictly according to changes of the skin .Check regularly the sensor placement and move it when the skin deteriorates. More frequent examinations may be required for different patients.
- □ Setting the SpO2 upper alarm limit to 100% is equivalent to switching off the alarm on upper limit. High oxygen levels may predispose a premature infant to retrolental fibroplasias. Therefore, the upper alarm limit for oxygen saturation must be carefully selected in accordance with commonly accepted clinical practices.

SPO2 ERROR MESSAGES

PLETH Waveform may display messages as below:

PROMPTS	EXPLAINATION	
Search Too Long	Search-time of SPO2 is too long	
Searching For Pulse	On searching for pulse signal	
Sensor Off	Sensor falls off or the finger fails to insert into the finger-probe	
SpO ₂ Com Error	SPO2 board has communication error with the mainboard	

NIBP MONITORING

SUMMARY ON NIBP MONITORING NIBP CUFF FITTING NIBP MONITORING INITIALIZATION NIBP MONITORING SETUP MEASUREMENT LIMITATIONS NIBP ERROR MESSAGES MAINTAINENCE AND CLEANING

SUMMARY ON NIBP MONITORING

The Non-invasive Blood Pressure (NIBP) module measures the blood pressure using the oscillometric method.

It is applicable for **adult**, **pediatric**, and **neonatal** usage.

There are three modes of measurement available: **manual**, **automatic** and **continuous**. Each mode displays the diastolic, systolic and mean blood pressure.

WARNING:

- □ You must not perform NIBP measurements on patients with sickle-cell disease or under any condition which the skin is damaged or expected to be damaged.
- □ For a thrombasthenia patient, it is important to determine whether measurement of the blood pressure shall be done automatically. The determination should be based on the clinical evaluation.
- Before starting a measurement, verify that you have selected a setting appropriate for your patient(adult, pediatric or neonate.) Ensure that the correct setting is selected when performing measurements on neonate, because the higher adult BP level is not suitable for neonate, it may be dangerous for the neonate to use an over pressure level.
- Do not apply the cuff to a limb that has an intravenous infusion or catheter in place. This could cause tissue damage around the catheter when infusion is slowed or blocked during cuff inflation.

NIBP CUFF FITTING

1. Whether choosing the suitable cuff which match the arm of patient influent much on the accuracy of NIBP measurement. The cuff width recommend by **AMERICA HEART SOCIETY** is the 40% of upper arm circumference or the 2% of the upper arm length.

2. Apply the blood pressure cuff to the patient's arm:

□ Make sure that the cuff is completely deflated.

 \Box Apply the appropriate size cuff to the patient, and make sure that the symbol " ϕ " is over the appropriate artery. Ensure that the cuff is not wrapped too tightly around the limb. Excessive tightness may cause discoloration and eventual isocheimal of the extremities.

3. Make sure that the cuff has not been twisted..

4. Insert the air pipe into the **NIBP** socket on the left panel of monitor. Ensure that the spring block on the left side of socket has been pressed

when inserting or unplugging the pipe, otherwise measurement process will be irregular and the sensor connector will be damaged.

WARNING:

- □ The width of the cuff should be either 40% of the limb circumference (50% for neonates) or 2/3 of the upper arm length. The inflatable part of the cuff should be long enough to encircle 50-80% of the limb. The wrong size of cuff can cause erroneous readings. If the cuff size is in question, then use a larger cuff.
- \Box Make sure that the cuff edge falls within the range of $\langle \rangle$. If does not, change a more suitable cuff.
- □ Connect the cuff to the air hose. The limb chosen for taking the measurement should be placed at the same level as the patient's heart. If this is not possible you should apply the following corrections to the measured values:

If the cuff is placed higher than the heart level, add 0.9mmHg (0.12kPa) for each inch of difference. If it is placed lower than the heart level, deduct 0.9mmHg (0.12kPa) for each inch of difference.

NIBP MONITORING INITIALIZATION

After opening the host machine, check the information indicating area before NIBP monitoring, if there is a note of NIBP MODULE SELF-CHECK OK, it shows that the NIBP module operate well, then begin NIBP monitoring, and the NIBP monitoring before this information is invalid; if there is NIBP MODULE SELF-CHECK ERROR, it shows that the NIBP module cannot be proceeded, click the button of **START/STOP** to give another time of self-checking or machine-opening, if it is also this information, contact with servicing engineer.

NIBP SETUP



Figure 10: Tree diagram for NIBP setup menu

ALARM SWITCH

ON and OFF for choice, the factory -set is OFF

PRESSURE UNIT

mmHg or kPa, the factory -set is mmHg.

PATIENT TYPE ADULT TYPE:

It can apply to the adult mode. In the initiated measurement, inflate the cuff to 180mmHg (24kPa), if the NIBP value cannot be measured, then inflate the cuff to higher than the form value by 50mmHg (6.7kPa), the maximum value cannot exceed 280mmHg (37.3kPa), and the enduring pressure range is 50-280mmHg.

PEDIATRIC/NEONATE TYPE

It can apply to the **PEDDIATRIC or NEONATE** mode. In the initiated measurement, inflate the cuff to 60mmHg (8kPa), if the NIBP value cannot be measured, then inflate the cuff to higher than the form value by 30mmHg (4kPa), the maximum value cannot exceed 150mmHg (20kPa), and the enduring pressure range is 50-150mmHg.

If this setup is before the NIBP module initiation, information indicating area will give a message of PATIENT TYPE SET ERROR.

The factory –set is **ADULT TYPE**.

Inflating range showing above has been realized on NIBP, NIBP use this inflation range to make sure the safety of patient.

INFLATION TYPE

MANUAL MODE、AUTOMATICAL MODE and STAT MODE

MANUAL MODE:

Click the button of **START/STOP** to begin inflation, the information indicating area display "MANUAL MEASURING... " which shows that it is on measurement just the moment.

If the NIBP value been measured, NIBP parameter area will display it and the information indicating area will give a note of "MANUAL MEASURING END!", then the measurement process finished.

If the NIBP value cannot be measured, NIBP parameter area will display error messages and automatically begin three times of measurement again, if the value cannot be measured also, the information indicating area will give a note of "RETRY OVER!" and never measure again.

During the measurement, click the button of **START/STOP** again will stop the NIBP measurement process and the information indicating area will give a note of STOP MANUAL MEASURING.

AUTOMATICAL MODE:

NIBP parameter area will display the countdown of "Auto measuring..." (TIME INTERVAL), so long as reaching the zero point, machine will automatically precede inflating measurement again and again until the mode be changed.

If the NIBP value been measured, NIBP parameter area will display it and the information indicating area will give a note of "AUTO MEASURING END! ", which show measurement process is finished and automatically begins another measurement until the mode is changed.

If the NIBP value cannot be measured, NIBP parameter area will display error messages and the first measurement automatically begin three times of measurement again, if the value cannot be measured.

also, the information indicating area will give a note of "RETRY OVER!" and automatically go on the next measurement until the mode is changed.

If click the button of **START/STOP** during any period of countdown, it is immediately begin inflation measurement.

During the measurement, click the button of **START/STOP** again will stop this period of NIBP measurement process and the information indicating area will give a note of "STOP AUTO MEASURING", but the automatic measurement period is continuous.

WARNING:

Prolonged non-invasive blood pressure measurements in Auto mode may be associated with purport, isocheimal and neuropathy in the limb wearing the cuff. When monitoring a patient, examine the extremities of the limb frequently for normal color, warmth and sensitivity. If any abnormality is observed, stop the blood pressure measurements.

STAT MODE:

Click the button of **START/STOP** to begin inflation, the information indicating area display "STAT MEASURING..." which shows that it is on measurement just the moment; If the NIBP value been measured, NIBP parameter area will display it and the information indicating area well give a note of "STAT MEASURING END".

If the NIBP value cannot be measured, NIBP parameter area will display error messages and automatically begin three times of measurement again, if the value cannot be measured also, the information indicating area will give a note of "RETRY OVER!", and then continue another time of measurement which lasts 5 minutes and then stop.

During the measurement, if click the button of **START/STOP** again, the information indicating area will give a note of "STOP STAT MEASURING" to stop the NIBP measurement and exit from this mode.

NOTE:

The value having been measured will display on the NIBP parameter area for 240 minutes unless a new measurement begin during this period. On the appropriate trend graph and trend table, the parameter will exist for correlated time length.

TIME INTERVAL

2/3/4/5/10/20/30/40/50/60/120/180/240 minutes for choice. This setting is used supported by **automatic** inflation mode, and the factory-set is **2** minutes.

FACTORY SETUP

Servicing engineer uses this function only.

MEASUREMENT LIMITATIONS

To different patient conditions, the oscillometric measurement has certain limitations. The measurement is in search of regular arterial pressure. In those circumstances when the patient's condition makes it difficult to detect, the measurement becomes unreliable and measuring time increases. The user should be aware that the following conditions could interfere with the measurement, making the measurement unreliable or longer to derive. In some cases, the patient's condition will make a measurement impossible.

PATIENT MOVEMENT

Measurements will be unreliable or may not be possible if the patient is moving, shivering or having convulsions. These motions may interfere with the detection of the arterial pressure pulses. In addition, the measurement time will be prolonged.
CARDIAC ARRHYTHMIA`S

Measurements will be unreliable and may not be possible if the patient's cardiac arrhythmia has caused an irregular heartbeat. The measuring time thus will be prolonged.

HEART-LUNG MACHINE

Measurements will not be possible if the patient is connected to a heart-lung machine.

PRESSURE CHANGES

Measurements will be unreliable and may not be possible if the patient's blood pressure is changing rapidly over the period of time during which the arterial pressure pulses are being analyzed to obtain the measurement.

SEVERE SHOCK

If the patient is in severe shock or hypothermia, measurements will be unreliable since reduced blood flow to the peripheries will cause reduced pulsation of the arteries.

HEART RATE EXTREMES

Measurements cannot be made at a heart rate of less than 40 bpm and greater than 240 bpm.

NIBP ERROR MESSAGES

Message indicating area may display messages like below:

Table 5: NIBP error messages
Patient moving!
Pressure < 10 mmHg!
Pressure < 1.3 kPa!
Pressure > 325 mmHg!
Pressure > 43.3 kPa!
Serial overtime!
Reset error!
Zero reset error!
Serial error
NIBP renew selfcheck
NIBP selfcheck
NIBP selfcheck error!
NIBP inter error !
Patient type error!
Setup patient
NIBP selfcheck ok!

MAINTAINENCE AND CLEANING

NOTE: Do not squeeze the rubber tube on the cuff.

REUSABLE BLOOD PRESSURE CUFF

The cuff can be sterilized by means of conventional autoclaving, gas or radiation sterilization in hot air ovens or disinfected by immersion in decontamination solutions, but remember to remove the rubber bag if you use this method. The cuff should not be dry-cleaned. The cuff can also be machine-washed or hand-washed; the latter method may prolong the service life of the cuff. Before washing, remove the latex rubber bag, and for machine-washing, close the Velcro fastening. Allow the cuff to dry thoroughly after washing, and then reinsert the rubber bag.

To replace the rubber bag in the cuff, first place the bag on top of the cuff so that the rubber tubes line up with the large opening on the long side of the cuff. Now roll the bag lengthwise and insert it into the opening on the long side of the cuff. Hold the tubes and the cuff and shake the complete cuff until the bag is in position. Thread the rubber tubes from inside the cuff, and out through the small hole under the internal flap.

TEMP MONITORING

THEORY OF OPERATION TEMP SENSOR INSTALLATION TEMP PARAMETER SETUP TEMP ERROR MESSAGES MAINTAINENCE AND CLEANING

THEORY OF OPERATION

The monitor provides one or two isolated temperature measurement channels (T1 and T2). When inserting one temperature sensor to any port of T1 or T2, the relative channel's test temperature value will display. If the second temperature channel is installed, the temperature difference between the two channels is an available option. Temperature difference is displayed as " Δ T" (delta temperature).

The monitor utilizes a temperature probe with a thermistor to give continuous electronic temperature readings of either core body temperature via rectal/esophageal probe or skin temperature via an external sensor.

TEMP SENSOR INSTALLATION

1. Insert the plug of **T1 or/and T2** sensor into the sensor socket on the left panel of monitor.

2. Put the probe on the patient according to the explanation of probe usage (lacuna and body).

WARNING:

Inspect the probe for wear or splitting after every disinfection/sterilization process is complete. If wearing or splitting of the probe is found upon visual inspection, a new probe should be used.

TEMP PARAMETER SETUP



Figure 11: Tree diagram for Temp setup menu

ALARM SWITCH

ON and OFF for choice, the factory-set is OFF.

TEMP UNIT

FAHRENHEIT or CELSIUS for choice, the factory-set is CELSIUS.

T ALARM UPPER-LIMIT

The T1 or T2 alarm upper-limit, the range is from **10** to **50°C** and the factory-set is **38.0°C** the single-step adjustable step- length is **0.1°C**

T ALARM LOWER-LIMIT

The T1 or T2 alarm lower-limit, the range is from **10** to **50°C** and the factory-set is **36°C** the single-step adjustable step- length is **0.1°C**

TEMP ERROR MESSAGES

TEMP SENSSOR OFF: the TEMP probe falls off the monitor.

MAINTAINENCE AND CLEANING

REUSABLE TEMP PROBES

1. The TEMP probe should not be heated above $100^{\circ}C(212^{\circ}F)$. It should only be subjected briefly t temperatures between $80^{\circ}C(176^{\circ}F)$ and $100^{\circ}C(212^{\circ}F)$.

2. The probe must not be sterilized in steam.

3. To clean the probe with alcohol detergent solution.

4. To clean the probe, hold the tip with one hand and with the other hand rubbing the probe down in the direction of the connector using a moist lint-free cloth.

ETCO2 MONITORING(OPTION)

THEORY OF OPERATION WARNINGS ABBREVIATIONS AND TERMINOLOGY ZEROING THE CO2 MODULE PATIENT AND TUBING PREPARATION ETCO2 SETUP ADVANCED SETUP CALIBRATION STATUS/ERROR MESSAGES

THEORY OF OPERATION

Carbon dioxide monitoring is used to monitor continuous carbon dioxide and report the End Tidal carbon dioxide ($EtCO_2$), inspired CO_2 and respiratory rate values of the intubated and non-intubated adult, pediatric, infant and neonatal patient.

Carbon dioxide monitoring system is a sidestream sampling system with a 50 ml/minute low sampling rate that is used to measure the CO_2 of non-intubated and intubated neonate, infant, pediatric and adult patients using specially designed sampling cannula and on-airway adapter kits. These kits incorporate a filter and the sample cell that provides maximum filtration of fluids and contaminants and protects the system from aspiration of these fluids.

In carbon dioxide monitoring system, infrared light is generated by the sensor and beamed through the sample cell to a detector on the opposite side. CO_2 from the patient that is aspirated into the sample cell absorbs some of this infrared energy. The monitor determines CO_2 concentration in the breathing gases by measuring the amount of light absorbed by these gases. EtCO₂ is displayed as a numerical value in millimeters of mercury (mmHg), percent (%), or kilopascals (kPa). In addition, a CO_2 waveform (capnogram) may be displayed which is a valuable clinical tool that can be used to assess patient airway integrity and proper endotracheal tube (ETT) placement. Respiration rate is calculated by measuring the time interval between detected breaths.

WARNING

- DO NOT position the sensor cables or tubing in any manner that may cause entanglement or strangulation. Support the carbon dioxide monitoring system airway adapter to prevent stress on the ET tube.
- Reuse, disassembly, cleaning, disinfecting or sterilizing the single patient use cannula kits and on-airway adapters may compromise functionality and system performance leading to a user or patient hazard. Performance is not guaranteed if an item labeled as single patient use is reused.
- Inspect the sidestream on-airway adapters and sidestream sampling kits for damage prior to use. DO NOT use the sidestream on- airway adapters and sidestream sampling kits if they appear to be damaged or broken.
- Replace the sidestream on-airway adapters and sidestream sampling kits if excessive secretions are observed.
- Monitor the CO₂ check waveform (Capnogram). If you see changes or abnormal appearance the patient and the sampling line. Replace line if needed.
- Do not apply excessive tension to any cable.
- DO NOT use device on patients that can not tolerate the withdrawal of 50 ml/min +/-10 ml/min from the airway or patients that can not tolerate the added dead space to

the airway.

- Do not connect the exhaust tube to the ventilator circuit.
- DO NOT stick appendage into sample receptacle.
- Always insert sample cell before inserting the on-airway adapter into the ventilated circuit.
- Always remove the on-airway adapter from the ventilated circuit before removing the sample cell.
- Nitrous oxide, elevated levels of oxygen, helium, Xenon, halogenated hydrocarbons, and barometric pressure can influence the CO₂ measurement.

ABBREVIATIONS AND TERMINOLOGY

EtCO ₂	End tidal carbon dioxide
AWRR	Air-way respiration rate
BARO	Barometric Pressure

ZEROING THE CO2 MODULE

The sample cell zero allows the CO_2 Module to adjust to the optical characteristics of the sample cell only when requested.

Whenever the type of adapter being used with the CO_2 Module is changed. For optimal accuracy, a CO_2 Module zero should also be performed whenever the CO_2 Module is connected to the patient monitor.

Before performing a CO_2 Module zero, the CO_2 Module should be removed from the patient monitor and the airway adapter type to be used in the circuit should be inserted into the CO_2 Module. Care should be taken ensure that the airway adapter is clear of any residual CO_2 gas. The maximum elapsed time for a CO_2 Module zero is 30 seconds. The typical time for a zero is 15 - 20 seconds.

Several CO_2 Module conditions may also request that a zero be performed. These requests stem from changes in the airway adapter that may indicate that the sensor is not in optimal measuring condition. When this occurs, the airway adapter should be checked to ensure optical occlusions such as mucus have not obscured the adapter window. If occlusions are found, the airway adapter should be cleaned or replaced.

NOTE:

- System does not allow adapter zero for 20 seconds after the last breath is detected.
- System does not allow adapter zero if temperature is not stable.
- An adapter zero cannot be performed if a sample cell is not connected to the module.

PATIENT AND TUBING PREPARATION

1. MODULE MOUNTING

a. Put the CO_2 module into the bracket of the rear panel of the monitor.

b. Check that monitor is switched off, Insert the plug of CO_2 sensor into the corresponding sensor socket marked with **EtCO₂** icon on the left panel of monitor.

WARNING: Don't hot plug $EtCO_2$ module, that is make sure that the OMNI is powered off before Insert the connector of CO_2 sensor into $EtCO_2$ socket. Otherwise the CO_2 module may be damaged by power supply from $EtCO_2$ socket of OMNI.

2. CONNECTING THE SAMPLE KIT

a. The sample cell of the sampling kit must be inserted into the sample cell receptacle of the CO_2 Module as shown in following figure. A "click" will be heard when the sample cell is properly inserted.



Figure 12: Connecting diagram for sample kit

b. Connect the CO₂ tubing to Nasal And Nasal/Oral Sidestream Kits.

c. Inserting the sample cell into the receptacle automatically starts the sampling pump. Removal of the sample cell turns the sample pump off.

d. To remove the sampling kit sample cell from the sample cell receptacle, press down on the locking tab and pull the sample cell from the sample cell receptacle.

DIRECTIONS FOR USE OF SINGLE PATIENT USE NASAL AND NASAL/ORAL SIDESTREAM KITS

CAUTION: The Nasal and Nasal/Oral Cannula kits are intended for single patient use. Do NOT reuse or sterilize the cannula kit as system performance will be compromised.

1. Verify that the cannula kit is clean, dry and undamaged. Replace the cannula kit if necessary.

2. Insert the sample cell into the sample cell receptacle as shown in above figure on Connecting the Sample Kit section. A "click" will be heard when properly inserted.

- 3. Perform a sample cell zero if prompted by the host system.
- 4. Place the nasal cannula kits onto the patient as shown in following figure.



Figure 13: Placing method for the nasal cannula kits

- 5. Some patients are prone to mouth breathing. The Oral/Nasal sampling cannula should be used on these patients, as most, if not all of the CO_2 is exhaled through the mouth. If a standard nasal CO_2 sampling cannula is used with these patients, the $EtCO_2$ number and capnogram will be substantially lower than actual.
- 6. When using the Nasal or Oral/Nasal CO₂ sampling kits with oxygen delivery, place the cannula on the patient as shown in Figure 14 and then attach the oxygen supply tubing to the oxygen delivery system and set the prescribed oxygen flow.
- 7. If the oral/nasal cannula is used, the oral sampling tip may need to be trimmed to adequately fit the patient (see following figure). Place the cannula onto the patient as shown in above figure. Observe the length of the oral cannula tip. It should extend down past the teeth and be positioned in the mouth opening. Remove the cannula

from the patient if the tip needs to be trimmed.



Figure 14: Placing method for oral sampling tip

CAUTION: Do NOT cut the oral cannula tip when the cannula is on the patient.

CAUTION: Remove the sampling kit sample cell from the CO_2 Module Inlet Port when not is use.

ETCO2 SETUP

Click the EtCO₂ Parameter Area to pop up the menu of EtCO₂ Setup, see graph below:



Figure 15: Tree diagram for ETCO2 setup menu ALARM SWITCH

ON and OFF for choice, the factory-set is OFF.

ETCO2 ALARM HIGH

The range is from **20** to **100 mmHg**, and the factory-set is **20mmHg**.

ETCO2 ALARM LOW

The range is from 10 to 95 mmHg, and the factory-set is 40mmHg.

AWRR ALARM HIGH

The range is from 10 to 150 mmHg, and the factory-set is 30mmHg.

AWRR ALARM LOW

The range is from **5** to **100 mmHg**, and the factory-set is **20mmHg**. The single-step adjustable length of alarm limit above is **5mmHg**.

ASPHYXIA DELAY

This setting is used to set the no breaths detected time-out. This time-out is the time period in seconds following the last detected breath at which the CO_2 module will signal no breaths detected.

The setting range is from **10** to **60 seconds**, and the factory-set is **20 seconds**.

ETCO2 UNIT

mmHg, kPa or percent (%), the factory –set is mmHg.

ETCO2 PERIOD

This setting is used to set the calculation period of the $EtCO_2$ value. The end-tidal CO_2 value is the highest peak CO_2 value of all end of expirations (end of breaths) over the selected time period. If less than two breaths exist in the selected time period, the value will be the maximum $EtCO_2$ value for the last two breathes.

This setting has 1 breath, 10 seconds and 20 seconds for choice, the factory –set is **20 seconds**.

SWEEP SPEED

From 12.5mm/s and 25mm/s for choice, the factory-set is 25mm/s.

ZERO SETUP

Pick up "**ZERO SETUP**" item to call up the zero setup menu:

Zero steps refer to "Zeroing the CO₂ Module" section detailed.

In above menu, complete the zero procedure by clicking the button "staring zeroing". During zeroing, a message of " $EtCO_2$ Zero Started" will be display on the message area.

NOTE: During the CO₂ module warmup period after the monitor is powered on, the monitor will perform an automatic zero calibration The maximum elapsed time for a CO₂ Module zero is 30 seconds. The typical time for a zero is 15 - 20 seconds.

ZERO GAS TYPE

When performing a zero on room air, this setting should be set to room air (the default). Only change to nitrogen (N_2) when performing a zero on 100% N_2 gas; this is provided for use in a laboratory environment.

ADVANCED SETUP

SET GAS COMPENSATIONS

The measurement of CO_2 is affected by temperature, pressure, and gas compensations. The barometric pressure as well as the presence of O_2 , N_2O , helium, and anesthetic agents in the gas mixture needs to be compensated for by the CO_2 module in order to achieve its stated accuracy. The instrument settings for these parameters should be set when initially connecting to the CO_2 module and whenever there is a change in the conditions at the patient airway.

In the CO_2 module, the temperature of the gas in the airway also effects the CO_2 measurement. It is necessary to adjust the instrument setting for the gas temperature to achieve the maximum accuracy for the CO_2 module.

BAROMETRIC PRESSURE

This setting is used to set current Barometric Pressure. The setting range is from **400** to **850 mmHg**. The factory –set is **760 mmHg**.

GAS TEMPERATURE

This setting is used to set temperature of the gas mixture. This setting is useful when bench testing using static gasses where the temperature is often room temperature or below.

The setting range is from **0** to 50° C. The factory –set is 35° C.

OXYGEN COMPENSATION

The setting range is from 0 to 100%. The factory –set is 16%.

BALANCE GAS

There are room air, N₂O and Helium items to choose.

ANESTHETIC AGENT

Use this setting to correct for the compensation of the gas mixture administered to the patient. Anesthetic agent is ignored when the balance gas is set to helium. The setting range is from **0.0** to **20.0%**. The factory –set is **0.0%**.

NOTE: At 700mmHg of pressure, the correct CO_2 value is 35.0 mmHg.

WAVEFORM SCALE

Use this setting to adjust the amplitude measurement (size) of the displayed \mbox{EtCO}_2 waveform scale manually.

There are two items to choose: $0 \sim 75$ mmHg, $0 \sim 150$ mmHg.

FILL IN WAVEFORM

Use this setting to fill in the bottom portion of the waveform on any channel of the display; the "fill in" can be canceled by choosing NO item.

CALIBRATION

No routine user calibration required.

Safety lock-outs:

- System does not allow sample cell zero for 20 seconds after the last breath is detected.
- System does not allow sample cell zero if temperature is not stable.

• An adapter zero cannot be performed if a sample cell is not connected to the module.

STATUS/ERROR MESSAGES

	Table 6: Staus/error messages for ETCO2
Messages	Descriptions
Sensor Off	The CO ₂ sensor is not connected
Sensor Warm Up	One of the following conditions exist:
	Sensor under temperature
	Temperature not stable
	Source Current unstable
Sensor Over Temp	Make sure sensor is not exposed to extreme heat (heat lamp, etc.). If
	error persists, return sensor to factory for servicing.
Sensor error	Check that the sensor is properly plugged in. Reinsert or reset the sensor
	if necessary. If error persists, return sensor to factory for servicing.
Sensor Zeroing	A zero is currently in progress.
Zero Required	To clear, check airway adapter and clean if necessary. If this does not
	correct the error, perform an adapter zero. If you must adapter zero more
	than once, a possible hardware error may exist.
Check Sampling Line	To clear, clean if sampling line mucus or moisture is seen. If the
	sampling line is clean, perform a zero.
CO ₂ Out of Range	The value being calculated is greater than the upper CO2 limit (150
	mmHg, 20.0 kPa, or 19.7 %). The maximum value output is the upper
	CO ₂ .
Check Airway	To clear, clean airway adapter if mucus or moisture is seen. If the
Adapter	adapter is clean, perform a zero.
Pump Life Exceed	The manufacturer stated pump life has been exceeded. Service may be
	required if Pneumatic System Error is present and can no longer be
	cleared.
Sensor Setup	The CO ₂ sensor is setting process.
EtCO ₂ Zero Error:	The CO_2 sensor is not ready for a EtCO ₂ Zero
Sensor Not Ready.	
EtCO ₂ Zero Error:	Breaths have been detected by the CO ₂ module within the last 20
Breath Detected.	seconds while a CO ₂ module zero was attempted.

MAINTENANCE AND CLEANING

SCHEDULE

The CO₂ Module flow rate accuracy should be verified by direct measurement using a calibrated flow meter every 12 months.

CLEANING

Cleaning the CO2 Module case, Cable and connector:

1. Use a cloth dampened with isopropyl alcohol 70%, a 10% aqueous solution of sodium hypochlorite (bleach), a 2% gluteraldehyde solution, ammonia, mild soap or disinfectant spray cleaner such as Steris Coverage® Spray HB.

2. Wipe down with a clean water-dampened cloth to rinse and dry before use. Make certain that the sensor windows are clean and dry before reuse.

NOTE: Do not immerse or sterilize the CO_2 Module.

Cleaning the Sidestream On-Airway Adapters and Sidestream Sampling Kits:

Sidestream on-airway adapters and sidestream sampling kits are single patient use. Treat in accordance with hospital protocols for handling single patient use devices.

IBP MONITORING(OPTION)

THEORY OF OPERATION INTRDUCTION WARNING PREPARATION FOR MONITORING INSTRUCTIONS FOR USE OF TRANSDUCER MONITORING KIT IBP SETUP SET TRANSDUCER ZERO PROMPT MESSAGE MAINTAINENCE AND CLEANING

THEORY OF OPERATION

There are two ways of measuring blood pressure: Direct (Invasive Pressure or IP) and Indirect (Non-invasive Blood Pressure or NIBP) method. The indirect method uses simple equipment but provides limited physiological information. The direct or invasive method (IP) provides accurate pressure measurements in regions of the cardiovascular system that are inaccessible to the indirect method.

To measure blood pressure by the invasive method, a catheter is inserted in a blood vessel and taken to the point of interest. The catheter has a transducer that provides electrical signals, which are then processed and analyzed by the monitor. Measurement of blood pressure by the invasive method gives the systolic (maximum), diastolic (minimum) and mean pressure.

The invasive pressure range is from -30 to 300 mmHg, allowing the operator to use the monitor for measuring arterial pressure, pulmonary artery pressure and central venous pressure.

INTRDUCTION

When an invasive pressure is selected to be displayed on a waveform channel, the monitor will default to the label P1 or P2, which indicates a general "Invasive Pressure". In addition, the monitor allows the selection of a pressure channel label that more clearly identifies a measurement. The choices for invasive arterial pressures are:

- ART Arterial Blood Pressure
- PA Pulmonary Artery Pressure
- CVP Central Venous Pressure
- RAP Right Arterial Pressure
- LAP Left Arterial Pressure
- ICP Intracranial Pressure

WARNING:

- For invasive pressure monitoring, routinely inspect the catheter and/or pressure line for leaks after zeroing. Always follow the pressure transducer/catheter manufacturer's use recommendations.
- Always zero the pressure transducer(s) prior to patient use.
- Non-physiological pulsatile invasive pressure waveforms (e.g., such as found during intra-aortic balloon pump use) can lead to inaccurate blood pressure readings. If questionable value is observed, re-check patient's pressures by alternate means before administering medication or therapy.
- The operator should avoid contact with the conductive parts of the appurtenance when being connected or applied.
- Disposable IBP transducer or domes should not be reused.
- Use only the pressure transducer designated by our company.
- Verify transducer cables fault detection before beginning of monitoring phase. Unplug the transducer of the channels from the socket, the screen will display the error message and audible alarm is activated, the other channel is the same.
- If any kind of liquid, other than solution to be infused in pressure line or transducer, is splashed on the equipment or its accessories, or may enter the transducer or inside the monitor, contact the hospital service center immediately.
- If there are air bubbles in the pressure line or the transducer, you should flush the system with solution to be infused.
- Calibrate the instrument either whenever a new transducer is used, or as frequently as indicated by your hospital procedures policy.

PREPARATION FOR MONITORING

Preparing for invasive pressure monitoring requires the following steps:

- Installation of transducer cable
- Kit set up
- Purging air from the Lines
- Zeroing, leveling and calibration
- Connecting monitoring system to patient
- Set IP channel and label
- Rescale the IP waveform
- Set the alarm limits
- Select printer option

INSTRUCTIONS FOR USE OF TRANSDUCER MONITORING KIT

INSTALLATION OF TRANSDUCER CABLE

- 1. Insert the plug of IBP transducer cable into the corresponding sensor socket on the left panel of monitor and check that monitor is switched on.
- 2. Prepare the pressure tubing and transducer by flushing through the system with normal saline solution. Ensure the system is free of air bubbles.
- 3. Connect that patient catheter to the pressure line; making sure that there is no air present in the catheter of pressure line.
- 4. Position the transducer so that it is the level with the patient's heart, approximately midaxillary line.
- 5. Check if you have selected the correct lable.

6. Zero the transducer.

KIT SET UP



Figure 16: Diagram for ETCO2 transducer setup

This section detail refers to relate to content detailed of instructions for use for disposable transducer monitoring kit.

PATIENT MOUNT



Figure 17: Patient mount for ETCO2 transducer

This section detail refers to relate to content detailed of instructions for use for disposable transducer monitoring kit.

IBP SETUP

Click the P1 Parameter Area to pop up the menu of IBP Setup, see graph below:



Figure 18: Tree diagram for IBP setup menu

ALARM SWITCH

Click the **alarm switch** item to pop up the IBP1 or IBP2 alarm setup menu.

The alarm setup range for high or low is from 0 to 300mmHg for ART label. The factory-set for high limit is 150 mmHg. The factory-set for low limit is 75 mmHg.

The alarm setup range for high or low is from -10 to 120mmHg for PA label. The factory-set for SYS high limit is 90 mmHg. The factory-set for SYS low limit is 60 mmHg. The factory-set for DIA high limit is 75 mmHg. The factory-set for DIA low limit is 45 mmHg.

The alarm setup range for high or low is from -10 to 40mmHg for CVP, RAP, LAP and ICP label. The factory-set for MAP high limit is 30 mmHg. The factory-set for MAP low limit is 10 mmHg.

ALARM SETUP

ON and OFF for choice, the factory-set is OFF.

SET LABEL

ART, PA, CVP, RAP, LAP and ICP are selectable.

SWEEP SPEED

From 12.5mm/s, 25mm/s and 50mm/s for choice, the factory-set is 25mm/s.

NOTE: The pressure unit is displayed in accord with setup of NIBP menu.

SET TRANSDUCER ZERO

After the system has been primed and mounted, zero the transducer. The following procedure should be completed periodically.

1. Turn zero reference stopcock "off" to the patient and remove yellow nonvented cap from the side port that opens the zero reference stopcock to air.

NOTE: The air-fluid interface of the zero reference stopcock should be at or near the right atrial (mid-axillary) level.

2. Click the button of "zero set" in IBP setup menu to call up the zero menu:

Upon connection of an invasive pressure transducer, the monitor will seek a steady pressure for zeroing. A sequence of on-screen status messages will be displayed.

- a. As soon as the power switch is turned on, "SENSOR OFF!" will be displayed on the screen in the message highlight area.
- b. When an invasive pressure transducer is inserted into the IP receptacle on the left side panel of the monitor, the initial waveform may be visible immediately based upon the most recently selected scale. The waveform scale numbers are not shown until transducer is zeroed. If the pressure transducer or interconnect cable is defective, the on-screen message "SENSOR OFF, UNABLE TO ZERO!" will remain on the screen. In this case, try another transducer or another cable.

NOTE:

It is the responsibility of the user to ensure that a zero procedure has recently been done on the transducer, otherwise there will be on recent, valid zero value for the instrument to use, which may result in inaccurate measurement results.

- Turn off patient 3-way stopcock before you start the zero procedure.
- The transducer must be vented to atmospheric pressure before the zero procedure.

3. Turn zero reference stopcock "off" to the side port. Replace nonvented yellow cap. **WAVEFORM SCALE**

Click the button of "waveform scale" to call up this menu.

The waveform and corresponding scale values will be displayed in the IBP waveform area. These scales can be set according to the table given below:

HI: IBP value of High Limit scale;

LO: IBP value of Low Limit scale.

Labels	High	Low
ART	50-300	0-100
PA	20-150	-10-50
CVP	0-150	-10-150
RAP	0-150	-10-150
LAP	0-150	-10-150
ICP	0-150	-10-150

Table 7: Limits for waveform scale for IBP

PROMPT MESSAGE

	npt message for IDF measure			
Messages	Descriptions			
OVERANGE, ZERO FAIL!	Make sure that the stopcock is vented to atmosphere. If the problem persists, contact service representative if necessary.			
TIMED OUT, ZERO FAIL!	Make sure that monitor is not in DEMO mode. Contact service representative if necessary.			
SENSOR OFF, UNABLE TO ZERO!	Make sure that channel 1 or channel 2 transducer is not off, and then proceed zeroing.			
ZERO IN PROCESS!	A zero is currently in progress.			
ZERO OK!	The zero procedure is completed.			

Table 8: Prompt message for IBP measure

MAINTAINENCE AND CLEANING

Make sure that the device is switched off and disconnected from the power cable before cleaning the monitor or the transducer.

The disposable transducers or caps is a single use kit, must not be re-sterilized or re-used.

OXYCRG DISPLAY

The OxyCRG diagram is used to quickly evaluate the patient status based on SpO2, HR and RESP.

The OxyCRG diagram will display the following three graphs simultaneity :

- 1. Heart rate (HR) data trend;
- 2. SPO2 data trend
- 3. Respiration rate or Constringent the respiration waveform

The information displayed for the above three parameters is 1 minute, 2 minutes, 4 minutes or 8 minutes newly.

Select "OxyCRG" item in the System Setup menu \rightarrow Display Mode to change to the OxyCRG display mode as following picture:



Figure 19: OxyCRG display

Note:

1. The OxyCRG dynamic view display appears in the below area of the screen and overlaps on the realtime waveforms area. At this time all monitoring is processing in and the alarm will still active.

2. The OxyCRG display speed is too slow to have effect on displaying when pressing down the waveform freeze button.

3. The number for the realtime waveforms display which is along with the OxyCRG diagram at the same time is determined by the Max. waveforms display number in the system. This detailed is seen as following table:

"OxyCRG" display is OFF	"OxyCRG" display is ON
6 hannels realtime waveforms	"OxyCRG" diagram + 3channels realtime waveforms
8channels realtime waveforms	"OxyCRG" diagram + 4channels realtime waveforms

DRUG DOSE CALCULATIONS

DESCRIPTIONS

The drug dose calculations function is used to calculate the Drug-perfusion ratio, dosage, amount or volume. The titration table for each drug can be displayed or printed.

Normally there are up to 24 sorts of drugs which can be calculated as following: Aminophylline, Amrinone,Bretylium, Dobutamine, Dopamine, Epinephrine, Heparin, Hydrocortisone, Inocor, Insulin, Isuprel, Levophed, Lidocain, Morphine, Neostigmine, Neosynephrine, Nipride, Nitroglycerine, Pronestyl and Versed.

OPERATING INSTRUCTIONS

1. Enter Tools \rightarrow Drug Calculation >> on the bottom menu bar.



Figure 20: Drug calculation menu

- 2. Choose patient type and drug name. Fifteen drugs below, including user defined drugs Drug A, B, C, D, E:
 - Drug A, B, C, D, E
 - Oxytocin
 - Nitroglycerin
 - Nipride
 - Lidocaine
 - Isoproterenol
 - Heparin
 - Adrenaline
 - Dopamine
 - Dobutamine
 - Aminophylline

Default values will be given, These are not the useful results. Doctor will provide known parameters' value.

3. Input the patient's weight(kg or lb) and the known parameters' value. Unit:

Drug A, B, C: Nitroglycerin, Nipride, Lidocaine, Isoproterenol, Adrenaline, Dopamine, Dobutamine and Aminophylline use units g, mg, mcg. Drug D: Oxytocin and Heparin use units Unit, KU, MU. Drug E: use unit mEq. Choose user defined drugs by the drug's unit. Titration Table:

Click Titration Table >> to open the titration table.

Titration Table - Oxytocin								
Dose	6.00	uni t/h	Rate	60.00	ml/h			
Amount	50.00	Uni t	Volume	500.00	ml			
Weight	154.3	1b	Drip rate	20.00	gtt/min			
Dose	Rate	Drip rate	Dose	Rate	Drip rate			
0.00	0.00	0.00	7.00	70.00	23.33			
1.00	10.00	3.33	8.00	80.00	26.67			
2.00	20.00	6.67	9.00	90.00	30.00			
3.00	30.00	10.00	10.00	100.00	33, 33			
4.00	40.00	13.33	11.00	110.00	36.67			
5.00	50.00	16.67	12.00	120.00	40.00			
6.00	60.00	20.00	13.00	130.00	43.33			
				_	_			
PageUp	PageDown	Interval	1	Benchmark	Dose			
)ose Type	Dose∕h		Record]	Exit			
lip to wat	tch for more	e data						

Figure 21: Titration table-oxytocin diagram

Interval, Becnchmark and Dose Type can be changed. Click PageUp or PageDown to choose the required table. Click Record to print currently displayed table.

PATIENT INFORMATION ADMINISTRATION

PATIENT BASIC INFORMATION SETUP TOOLS SETP NIBP DATA LIST OBSERVATION TREND GRAPH ANALYSIS TREND ANALISIS TRANSFERRING TRENDS VIA RS-232

PATIENT BASIC INFORMATION SETUP

Click the patient information area as following:



Figure 22: Patient info display area diagram

Pop up the menu of Patient Information Setup (see graph below). The menu can set up the following patient record:

ID

Set the ID number of the patient.

NAME

Move the cursor box to the correlated position and press it, turn the ROTARY KNOB after choosing can input the English alphabet or Chinese phonetic letters, the range is: uppercase, A-Z, point(.)and blank character.

Patient name support the display method of English and phonetic letter, and do not support the Chinese character input. The user can input 9 characters at most.

SEX

Set the patient gender, the default setting is MALE.

BLOOD

Set the blood type of the patient. It can be: N/A(unknown type), A, B, O, AB, RH+, RH- and so on, the default setting is N/A.

AGE

Set the age of the patient, the default setting is 20.

NOTE:

□ Once the user choose the method of **YES** to exit from the Patient Information Setup, all information of patient will be refreshed and the trend data will be renovated.

TOOLS SETUP

There are four types of events that you can define. The user can freely define the implication of each type by select **Tools**→ **Mark event**.

MARK EVENT

To mark the event: click and select one from A , B, C and D. There is a **V** mark signal for the one selected.

Choose the button of **YES** to exit, and the event marked is become effective immediately upon the exit, or else it will not become effective.

IMPORTANCE OF EVENT MARKING:

It can classify the circumstances which influence the parameter monitoring on patient, for example, medicine taking, injection and other treatment, These events, displaying on NIBP list, trend graph or table, are very important to the parameter analysis.

NIBP LIST OBSERVATION

							,		
NO.	Time	NIBP	HR	Sp02	PR	T1/T2	RR	ST	ET
1	03/11 14:27:36	120/80 100	60	99	60	37.1/37.4	20	-0.01	N
2	03/11 14:27:11	120/80 100	60	99	60	37.1/37.4	20	-0.01	N
3	03/11 14:26:46	120/80 100	60	99	60	37.1/37.4	20	-0.01	N
4	03/11 14:26:21	120/80 100	60	99	60	37.1/37.4	20	-0.01	N
5	03/11 14:25:55	120/80 100	60	99	60	37.1/37.4	20	-0.01	N
6	03/11 14:25:30	120/80 100	60	99	60	37.1/37.4	20	-0.01	N
7	03/11 14:25:05	120/80 100	60	99	60	37.1/37.4	20	-0.01	N
8	03/11 14:24:40	120/80 100	60	99	60	37.1/37.4	20	-0.01	N
9	03/11 14:24:14	120/80 100	60	99	60	37.1/37.4	20	-0.01	N
	PageDown	PageUp			Rec	ord	Clos	e	

Click the button of **NIBP LIST** to pop up the menu of NIBP LIST, see chart below:

Table 9: NIBP list

NIBP LIST MOVING

Click the choice item to the correlated button position as indicated on chart to complete relevant operation, including in page down, page up, and record.

The NIBP list can save 256 group of data.

NOTE:

Only after the NIBP value has been measured can it be added to the NIBP Data List. NIBP list can save 256 groups of data at all, if exceed, the new data will kick the most former data out of the list and be added to the list automatically.

TREND OBSERVATION

PARAMETER FOR TREND OBSERVATION

The monitoring system will save and trace the trend of parameters below: HEART RATE (HR)、OXYGEN SATURATION (SPO₂) NONTRAUMATIC BLOOD PRESSURE (NIBP-SYSTOLIC BLOOD PRESSURE (SYS, DIASTATIC BLOOD PRESSURE (DIA) TEMPERATURE (TEMP) PULSE (PULSE) RESPIRATION RATE (RESP) ETCO2 IBP1 IBP2 ST EVENT

TREND OBSERVATION ADMITTANCE

Click the function button of **TREND** button to pop up the graph below:



Figure 23: Trend management menu

Data-recording Status Bar:

It is used to show the current data-recording length. For example, the user set a trend of 15 minutes, if the color of bar right moment is red, it means that the data-recording time is shorter than 15 minutes, i.e. the data-recording length is smaller than the time-length choosing by user; if the color is light-blue, it means that the data-recording is equal to the choosing time-length; if the bar presents the light-blue and green alternately, it means that the data-recording length is larger than the setup time length, and the light=blue part is the proportion of data-recording length covered by the time-length, and the green part is the proportion of data-recording length covered by time-length which has not been chosen.

TREND TIME CHOOSING

Trend time is the time length before current time.

There are eleven items for trend time choosing: 15 minutes, 30 minutes, 60 minutes, 90 minutes, 3hours, 6 hours, 12 hours, 18 hours, 24 hours, 36 hours, 48 hours. For instance, if 30 minutes is chose as the reference trend time, then we can recall the trend change of 30 minutes before current time.

TREND TIME INTERVAL

Trend Time Interval means how often the system stores a trend data. Different trend reference time has its correlated trend time interval, the relation between them are show below:

30 minutes: 6 seconds

60 minutes:	12 seconds
90 minutes:	18 seconds
3 hours:	36 seconds
6 hours:	72 seconds
12 hours:	144 seconds
18 hours:	216 seconds
24hours:	288 seconds
36 hours:	432 seconds
48 hours:	576 seconds
72 hours:	864 seconds

TREND GRAPH ANALYSIS

If want to choose the Trend Graph Analysis on trend management menu, click the item of **Trend Graph** to pop up the trend graph menu like the graph below:



Figure 24: Trend graph menu

Each page display the trend chart of one parameter, the user can change it by clicking the item of **PARAMETER**.

TREND TABLE ANALYSIS

TREND TABLE ADMITTANCE

Click the item of **Trend Table** on the trend management menu, the trend table menu will display in the waveform area on the screen. Sixteen groups of parameters are listed each one page. These data will list by following the order of from new to former and the time is displaying at the 24-hour system. The parameter name is display on the top of chart and the invalid data will not display.

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NO.	TIME	NIBP	Sp02	HR	PR	RR	T1	T2	ST	EtC02	IBP1	IBP2	ET
1	2/11 15:11:06	120/80	99	60	60	20	37.1	37.4	- 0.01	38	126/86	32/18	N
2	2/11 15:11:12	120/80	99	60	60	20	37.1	37.4	- 0.01	38	126/86	32/18	N
3	2/11 15:11:18	120/80	99	60	60	20	37.1	37.4	- 0.01	38	126/86	32/18	N
4	2/11 15:11:24	120/80	99	60	60	20	37.1	37.4	- 0.01	38	126/86	32/18	N
5	2/11 15:11:30	120/80	99	60	60	20	37.1	37.4	- 0.01	38	126/86	32/18	N
6	2/11 15:11:36	120/80	99	60	60	20	37.1	37.4	- 0.01	38	126/86	32/18	N
7	2/11 15:11:42	120/80	99	60	60	20	37.1	37.4	- 0.01	38	126/86	32/18	N
8	2/11 15:11:48	120/80	99	60	60	20	37.1	37.4	- 0.01	38	126/86	32/18	N
9	2/11 15:11:54	120/80	99	60	60	20	37.1	37.4	- 0.01	38	126/86	32/18	N
10	2/11 15:12:00	120/80	99	60	60	20	37.1	37.4	- 0.01	38	126/86	32/18	N
11	2/11 15:12:06	120/80	99	60	60	20	37.1	37.4	- 0.01	38	126/86	32/18	N
12	2/11 15:12:12	120/80	99	60	60	20	37.1	37.4	- 0.01	38	126/86	32/18	N
13	2/11 15:12:18	120/80	99	60	60	20	37.1	37.4	- 0.01	38	126/86	32/18	N
14	2/11 15:12:24	120/80	99	60	60	20	37.1	37.4	- 0.01	38	126/86	32/18	N
15	2/11 15:12:30	120/80	99	60	60	20	37.1	37.4	- 0.01	38	126/86	32/18	N
16	2/11 15:12:36	120/80	99	60	60	20	37.1	37.4	- 0.01	38	126/86	32/18	N
16	2/11 15:12:36 PageUP	120/80	99 Pa	60 ageDo	60 wn	20	37.1	37.4 Record	- 0.01	38	126/86	32/18	

Table 10: Trend table

TREND TABLE MOVING

Click the choice item to the correlated item position as indicated on chart to complete relevant operation, including in page down, page up and record.

NOTE: The trend table can save total 43200 groups of data.

ALARM EVENTS RECALL

If click the item of **Alarm Events** on the trend management menu, the Alarm Event Review window will display in the waveform area on the screen. In this window you can select the alarm parameter (10 parameters), alarm waveform (12 waveforms) and alarm times (8 times).

- 1. Alarm waveform select menu It is used to select alarm waveform.
- 2. Alarm times select menu There are eight times.

3. The last waveforms select menu

Click the item of **Last Waveform** to pop up the last waveform review window like the graph below:



Figure 25: Last waveform review

When there are waveforms display for demonstration or realtime measurement, the system only save data for the last 16 seconds and display two selectable waveforms, the

happened time for the late waveforms will display on the title bar in the window as below. Recent Wave Review -- Default 1/ 1/2010 08:20:20



Figure 26: Recent waveform review

ST SEGMENT MONITORING

OPERATON SEQUENCE

Click the button of in the ECG setup \rightarrow **ST Analysis** menu to pop up the ST setup menu.

ST ANALYSIS SWITCH

The default value is **OFF**, only the choice of **ON** can operate the ST Segment Monitoring. The ST Segment result been measured on the lead appointed by the user is the digital form when appearing on the ST Parameter Area. Meanwhile, the **TREND GRAPH** or **TREND TABLE** can be opened by the button of **TREND** to see the tendency displaying on the graph or table.

ST ALARM SWITCH

The default value is **OFF**. The alarm is triggered when the ST measurement value exceeds the alarm limits. If the ST Alarm is **ON**, the ST value blinks, the alarm sounds and the alarm indicator flashes, and the information column will give the note that **ST HIGHER** or **ST LOWER**.

ST ALARM LIMIT

Set the ST alarm upper limit and lower limit separately. The default upper limit is+0.30, The default lower limit is -0.30.

The Most High	gh Limit	the Most Low Limit	Single Adjustment
ST 2	2.00mV	-2.00mV	0.02 mV

ST ADJUST

After choosing, the menu below will appear (the graph is the obtained ST Segment module):



Figure 27: ST adjust setup menu

ISO (Base Point)

Set the baseline point, its adjustable range is $-508ms^{-4}ms$, the default value is -80ms, it shows that the reference point is the position 80ms before the peak of R-wave locates.

ST (Measurement Point)

Set the measuring point, its adjustable range is +508ms \sim +8ms, the default value is +108ms, it shows that the reference point is the position 108ms after the peak of R-wave locates.

These two points can be adjusted by clicking the button of << or >> . The value and the indicating line will change simultaneously. The ST measurement for each beat complex is the vertical difference between the two measurement points.

NOTE: The ST measurement point should be adjusted if the patient's HR or ECG morphology changes significantly.

ATTENTION POINTS

1. ST Segment Measurement Value:

Unit: mV

Implication: The positive number means the ST SEGMENT is elevating, and the negative number means depressing. Measurement Range: -2.0mV \sim +2.0mV

2. Indicating Information Explanation:

ST SEGMENT TOO HIGH: means that the ST value is above the upper limit of alarm. ST SEGMENT TOO LOW: means that the ST value is below the lower limit of alarm.

3. Other Points:

Abnormal QRS complex is not considered in ST segment analysis. If the ST Segment Calibration cannot be entered, it means that no ST module for use. It only appears when the ECG signal isn't existed.

DATA RECALL

Data Recall Storage Data Recall Displays Recall Operation Descriptions

DATA RECALL STORAGE

Data Recall in graphical or tabular format can be displayed on the screen or transferred to on the computer for analysis via RS232 interface, and printed if a printer is installed.

The data recall for all parameters is the average of a 6-second sample of the data. Seventy two (72) hours of recall data is stored in a nonvolatile memory, and remain in storage when the monitor is in Standby.

A new record of data recall is started each time the monitor is turned on. A data recall record is defined as the data from one power on event to the Standby power event. A date/time annotation is included at the start of each new record (up to eight patients') and the record can be correlated with the patient. Once the recall memory has stored 72 hours of data, the oldest recall data will be overwritten by new data.

RECALL DATA DISPLAYS

The data recall are displayed in graphical or tabular format. The recall information in graphical format for a selected parameter is shown as a line connecting each of the points representing the stored 6-second average.

The information stored for each recall episode can include:

- Numeric vital signs for all the measurements monitored
- Waveforms for up to 12 measurements of alarm events for your choice

You can navigate through the recall database to view events retrospectively, and you can document recalls on a recording or report marked with the patient name, patient ID, the data and time.

RECALL OPERATION DESCRIPTIONS

Input ID and name of a patient for recall

After you power on the monitor, there is a window pop out on the screen to remind you input the patient's ID as following:



Figure 28: Prompt window for input patient ID

The above window will be automatically closed in count down 10 seconds.

Note: If you want recall data for a patient at any time, you must input his/her ID number, otherwise you couldn't recall this patient's history data. Of course, you could input this patient's ID later by click the patient setup icon.

If click **YES** item, enter the Patient Setup menu.

Recall Operation

1. Click the **Recall** icon on the bottom of the function keys to open the recall function for up to 8 patients.

2. Select the patient's ID for recall: Complete step 1, select patient's ID.

3. Complete step 1 and step 2, enter the following **Trend Management** window with Patient ID as figure 24.

Note: This trend management-default window is for a patient which has no ID number.

ARR ANALYSIS

Arrhythmia analysis can monitor 13 kinds of arrhythmias:

ASY ---- Asystole

FIB ---- Fibrillation

VTA ---- Ventricular tachycardia

ROT ---- R ON T

RUN ---- Ventricular Run

TPT ---- Ventricular Triplet

CPT ---- Ventricular Couplet

VPB ---- Ventricular prematare beat

BGM ---- Bigeminy

TGM ---- Trigeminy

TAC ---- Tachycardia

BRD ---- Bradycardia

MIS ---- Miss beat

The monitoring system supports the self relearn function to accommodate itself to new conditions such as different patients. The user can edit the arrhythmia type. For each type system saves 8 items arrhythmia and totally saves 104 items.

ECG PACE

Set the ECG Pace on or off. The factory-set is OFF,

If ECG Pace is set to be **on**, the arrhythmia analysis is off to avoid invalid arrhythmia analysis.

OPERATON SEQUENCE

Click the button of ECG setup \rightarrow **ARR Analysis** menu to pop up the ARR setup menu.

ARR ANALYSIS

Set arrhythmia analysis to be ON or OFF. The factory-set is ON,

ARR SOURCE

Select between **lead I, lead II** and **V**, and the factory-set is **lead II**. The user can switch the ECG lead if the current lead's signal is weak.

PVC MONITOR

Set PVC monitor to be **ON** or **OFF**. The factory-set is **ON**, If the premature ventricular contraction times exceed the PVC ALARM COUNT, the system will alarm.

PVC ALARM COUNT

Its set range is from 1 to 10. The factory-set is 10.

ARR RELEARN

Self relearn to accommodate itself to new conditions. Such as different patients, cardiograph change a lot.

ARR ALARM SETUP

Set all kinds of arrhythmia alarm to be **ON** or **OFF**. The all factory-sets are **ON**.

ARR REVIEW

Click the button of **Select** to choose a arrhythmia item.

Click the button of **Wave** to review the selected arrhythmia item includes items of HR, ST, PR, SpO2,NIBP,Temp,Resp and PVC.



Figure 29: ARR detail info

Click the button of **Rename** to rename a selected ARR item. Click the button of **Delete** to delete a selected ARR item. Click the button of **Sort** to sort the arrhythmia items by **time** or **type**. Choose the type of **Sort**. The factory-set is by **Time**.

RECORDING (OPTION)

NOTE: This is thermal recorder which must use the thermal record paper (the specification is 48mm on width). The waveforms can be printed by 2 or 3 traces.

REAL-TIME WAVEFORM RECORDING

1. CONTENTS:

(1) From the preceding 8 seconds before the recording, it can record a burst of two waveforms, Gain and Paper Advance Speed.

(2) Record the parameter report simultaneously which include:

Patient Name, Hospital name, Recording Time, LEAD, HR, RESP, SPO2, T1, T2, ETCO2, IBP1, IBP2, ST Segment, the latest NIBP's Blood Pressures of SYS and DIA (mmHg), see graph below:



Figure 30: Prinoutt sample for 2 traces

2. RECORDER SETUP

3. OPERATION SEQUENCE

Click the function button of **RECORD** on the screen \rightarrow The statement of START RECORDING appear on the bottom of screen, which shows that the print process is going on; If want to terminate print during the printing process, just press the button of RECORD again , the recorder stop immediately as the statement of BREAK RECORDING will appears on the bottom of screen.

NOTE: Each time of pressing the button of RECORD will either carry out REAL-TIME RECORDING or terminate CURRENT RECORDING TASK.

TREND TABLE RECORDING

1. CONTENTS

Record 11 groups of parameter report, include seven items of DATE, TIME, HR, ST SECTION, RESP, SPO2, T1/T2, SYS/DIA, ETCO2, IBP1, and IBP2, see table below:

Date	2008/03/03	2008/03/03	2008/03/03	2008/03/03	2008/03/03	2008/03/03	2008/03/03	2008/03/03
Time	10:30:17	10:30:14	10:30:11	10:30:08	10:30:05	10:30:02	10:29:59	10:29:56
NIBP	/	120 / 80	120 / 80	120 / 80	120 / 80	120 / 80	120 / 80	120 / 80
SP02		and the second sec	100 mm - 100					
HR		59	59	59	59	59	59	59
PR						·		
RESP		20	20	20	20	20	20	20
T1/T2	/	37.1/37.2	37.1/37.2	37.1/37.2	37.1/37.2	37.1/37.2	37.1/37.2	37.1/37.2
ST		+0.00	+0.00	+0.00	+0.00	+0.00	+0.00	+0,00

Table 11: Printout sample for trend table

2. RECORDING SEQUENCE

Click the function button of **TREND** on the screen \rightarrow TREND MANAGEMENT menu appears on screen \rightarrow click the button of TREND TABLE \rightarrow XX TIME TREND TABLE menu appears on screen \rightarrow click the button of RECORD \rightarrow START RECORDING.....If want to stop record, click the button of record again or choose the soft button of RECORD

again \rightarrow BREAK RECORDING.

ALARM RECORD

1. CONTENTS

- (1) It can record a burst of waveform of 10 seconds (the preceding 4 seconds before the recording till the current 4 second), after the alarm, includes Gain and Paper Advance Speed.
- (2) One paper of alarm report, includes Alarm Begin Time, Alarm Lead, HR, RESP, SPO2, T1/T2, ST SECTION, Alarm Type, the latest NIBP's SYS and DIA(mmHg) value, Alarm Types are explained as below:

HR ALM	Heart Rate Alarm
SYS ALM	Systolic Pressure Alarm
DIA ALM	Diastolic Pressure Alarm
ST ALM	ST Segment Alarm
SPO2 ALM	SPO2 Alarm
TEMP ALM	Temperature Alarm
RESP ALM	Respiration Alarm



Figure 31: Printout sample for alarm record

2. RECORDING SEQUENCE

Click the button of **TREND** on the screen— \rightarrow TREND MANAGEMENT menu appears on screen— \rightarrow click the item of ALARM EVENT— \rightarrow ALARM EVENT menu appears on screen— \rightarrow click button of RECORD— \rightarrow START RECORDING, If want to stop printing, click the button of RECORD again or click the soft button of RECORD again— \rightarrow BREAK RECORDING.

If there is several parameters alarm simultaneously, click and choose them on ALARM menu, parameter alarms indicated by red sigh can be record simultaneously. The same parameter alarm can only record one sequence from the alarm state into the normal state; it can only be refreshed until the next state of alarm.

[NOTE] : "------"means invalid parameter.

RECORDING EXPLANTION

1. INSERTING PAPER

Press the button of the catch on the recorder, open the catch and take the old paper roll out and insert a new one into the paper cassette. Pay attention that the paper is turning swiftly. Pull a small length of paper out of the catch from the lower end of the roll (If it is the upper end, the paper reel installed conversely), close the catch, and make sure that the paper is just in the groove, or else paper advance will not be orderly.

2. ATTENTIONS

(1) The time of continuous print cannot exceed 2 minutes.

(2) Do not press the button of RECORD when there is no paper, or the recorder head will be damaged.

(3) Only thermal record paper can be used.

(4) If there is too much dust, use a sponge lightly moistened with alcohol to clean the correlated parts.

3. RECORDING INDICATING MESSAGES

(1) START RECORDING

Recording process is going on

(2) BREAK RECORDING: The button of RECORD been pressed again during the process of printing, so it can press the button once again to re-start it.

(3) DOOR OPEN

Recorder's door is opened.

(4) DOOR CLOSE

Recorder's door has been closed.

(5) PAPER OK:

Showing that record paper has been installed well

(6) NO PAPER

Record paper has been use up.

(7) RECORDER READY

Showing that recorder has been connected well.

(8) RECORDER NOT READY

Showing that recorder hasn't been connected well.

GRID OUTPUT

Some recording paper without grid, in order to observe the waveform easily, you can set the grid form. The set method is as below:

Click the function button of **SETUP** \rightarrow the menu of SYSTEM SETUP \rightarrow recorder setup \rightarrow RECORDER SETUP MENU \rightarrow RECORDER GRID is set to ON (default value is OFF, then the waveform being recorded is in the grid form
BATTERY OPERATION

OMNI type patient monitor is equipped with a rechargeable battery. The monitor through the AC INPUT socket to recharge the battery until it is full. A symbol is displayed in the upper right quarter of the screen to indicate the status of recharging, in which the color part represents the electric energy of the battery.

A new, fully charged battery will provide about 1 hour of monitoring time under the following conditions: no audible alarms, no analog or serial output devices attached, and no backlight.

When operating on battery, the monitor will prompt alarm and shut off automatically when the electric energy is low. When the electric energy is lower than 25% of total power capacity, the alarm will be active, at the same time the message of "Battery Power Low" will display in the message area in the top of screen. The battery signal will change red, Connect the monitor to AC power at this moment can recharge the battery while operating. If keep operating on the battery, the monitor will shut off automatically upon exhaustion of the battery.

NOTE: Whenever the monitor is connected to AC power, the battery is being charged. Therefore, it is recommended that the monitor remain connected to AC power when not in use. This will make available a fully charged battery for use at any time.

NOTE: As the battery is used and recharged over a period of time, the amount of time between the onset of the low battery alarm and the instrument shut-off may become shorter.

If the backlight is turned off during a low battery condition, it cannot be turned back on. It is recommended that the internal battery is replaced by qualified service personnel every 24 months.

CAUTION: If the OMNI is to be stored for a period of 2 months or longer, notify service personnel to remove the battery from the monitor prior to storage. Recharge the battery when it has not been charged for 2 or more months.

WARNING: The batter need to be charged before use. Please note the inserting direction, insert the battery rightly according to the instruction on the label attaching on the battery. Otherwise it will damage the inner structure of the battery housing.

DISPOSAL OF DEVICE COMPONENTS

Follow local governing ordinances and recycling instructions regarding disposal or recycling of device components, including batteries.

PERIODIC SAFETY CHECKS

It is recommended that the following checks be performed every 24 months.

- Inspect the equipment for mechanical and functional damage.
- Inspect the safety relevant labels for legibility.

CLEANING

WARNING: Do not spray, pour, or spill liquid on OMNII, its accessories, connectors, switches, or openings in the chassis. Do not immerse the OMNI II or its accessories in liquid or clean with caustic or abrasive cleaners.

To clean the OMNI, dampen a cloth with a commercial, nonabrasive cleaner and wipe the exterior surfaces lightly. Do not allow any liquids to come in contact with the power connector or switches. Do not allow any liquids to penetrate connectors or openings in the instrument. For cables, sensors, and cuffs, follow the cleaning instructions in the directions for use that accompany these accessories.

SPECIFICATIONS

SAFETY		
Meet the requirement of EN60601 series, CE marking according to MDD93/42/EEC Equipment not suitable for use in the presence of a flammable anaesthetic mixture with air or with oxygen or nitrous oxide		
Type of Protection:	Class I (on AC power), Internally powered (on battery power)	
Degree of Protection: Sterilization or Disinfection methods:	Type BF, defibrillation-proof CF - Applied part 70% isopropyl alcohol solution or a nonstaining disinfectant.	
Operation Mode: Protection Against Ingress of Liquid's:	Continuous Operation IPX0	
APPLICATION Neonatal, pediatric and adult patients		
Physical Dimensions & Weight Base Unit:	310×270×145mm	
Weight:	4.7kgs	
PEFORMANCE SPECIFICATIONS		
Display:	10.1″ color TFT	
Resolution:	1024×860 3 or 6 wayoforms	
Waveforms	ECG(I, II, III, aVR, aVL, aVF, V1-V6), PLETH, RESP, IBP1, IBP2, ETCO2	
Indicator:	Alarm indicator Power indicator	
	QRS beep and alarm sound	
Trend time:	From 30 minutes to 72 hours	
ECG		
Input:	5 lead or 3 lead ECG cable and standard AAM	
Input:	5 lead or 3 lead ECG cable and standard AAMI line for connection	
Input: Lead Choice:	5 lead or 3 lead ECG cable and standard AAMI line for connection I, II, III, aVR, aVF, aVL, V	
Input: Lead Choice: Gain Choice:	5 lead or 3 lead ECG cable and standard AAMI line for connection I, II, III, aVR, aVF, aVL, V x0.5, x1.0, x2.0	
Input: Lead Choice: Gain Choice: CMRR (common mode	5 lead or 3 lead ECG cable and standard AAMI line for connection I, II, III, aVR, aVF, aVL, V ×0.5, ×1.0, ×2.0 >100 dB at 50 Hz or 60 Hz	
Input: Lead Choice: Gain Choice: CMRR (common mode rejection ratio):	5 lead or 3 lead ECG cable and standard AAMI line for connection I, II, III, aVR, aVF, aVL, V ×0.5, ×1.0, ×2.0 >100 dB at 50 Hz or 60 Hz	
Input: Lead Choice: Gain Choice: CMRR (common mode rejection ratio): Frequency Characteristic:	5 lead or 3 lead ECG cable and standard AAMI line for connection I, II, III, aVR, aVF, aVL, V ×0.5, ×1.0, ×2.0 >100 dB at 50 Hz or 60 Hz 0.67~40 Hz (+3dB attenuation) Z channels	
Input: Lead Choice: Gain Choice: CMRR (common mode rejection ratio): Frequency Characteristic: ECG Waveforms: Sweep Speed:	5 lead or 3 lead ECG cable and standard AAMI line for connection I, II, III, aVR, aVF, aVL, V x0.5, x1.0, x2.0 >100 dB at 50 Hz or 60 Hz 0.67~40 Hz (+3dB attenuation) 7 channels 12 5, 25 and 50 mm/s	
Input: Lead Choice: Gain Choice: CMRR (common mode rejection ratio): Frequency Characteristic: ECG Waveforms: Sweep Speed: HR Display Range:	5 lead or 3 lead ECG cable and standard AAMI line for connection I, II, III, aVR, aVF, aVL, V $\times 0.5$, $\times 1.0$, $\times 2.0$ >100 dB at 50 Hz or 60 Hz 0.67~40 Hz (+3dB attenuation) 7 channels 12.5, 25 and 50 mm/s 30~300bpm	
Input: Lead Choice: Gain Choice: CMRR (common mode rejection ratio): Frequency Characteristic: ECG Waveforms: Sweep Speed: HR Display Range: Accuracy:	 5 lead or 3 lead ECG cable and standard AAMI line for connection I, II, III, aVR, aVF, aVL, V ×0.5, ×1.0, ×2.0 >100 dB at 50 Hz or 60 Hz 0.67~40 Hz (+3dB attenuation) 7 channels 12.5, 25 and 50 mm/s 30~300bpm +1bpm or +1% whichever is greater 	
Input: Lead Choice: Gain Choice: CMRR (common mode rejection ratio): Frequency Characteristic: ECG Waveforms: Sweep Speed: HR Display Range: Accuracy: Alarm Limit Range	 5 lead or 3 lead ECG cable and standard AAMI line for connection I, II, III, aVR, aVF, aVL, V x0.5, x1.0, x2.0 >100 dB at 50 Hz or 60 Hz 0.67~40 Hz (+3dB attenuation) 7 channels 12.5, 25 and 50 mm/s 30~300bpm ±1bpm or ±1%, whichever is greater Upper limit:80~400bpm Lower limit: 20~150bpm 	
Input: Lead Choice: Gain Choice: CMRR (common mode rejection ratio): Frequency Characteristic: ECG Waveforms: Sweep Speed: HR Display Range: Accuracy: Alarm Limit Range	5 lead or 3 lead ECG cable and standard AAMI line for connection I, II, III, aVR, aVF, aVL, V $\times 0.5$, $\times 1.0$, $\times 2.0$ >100 dB at 50 Hz or 60 Hz $0.67 \sim 40$ Hz (+3dB attenuation) 7 channels 12.5, 25 and 50 mm/s $30 \sim 300$ bpm ± 1 bpm or ± 1 %, whichever is greater Upper limit: $80 \sim 400$ bpm Lower limit: $20 \sim 150$ bpm ± 5 mV AC	
Input: Lead Choice: Gain Choice: CMRR (common mode rejection ratio): Frequency Characteristic: ECG Waveforms: Sweep Speed: HR Display Range: Accuracy: Alarm Limit Range Input Dynamic Range: Defibrillator Discharge:	5 lead or 3 lead ECG cable and standard AAMI line for connection I, II, III, aVR, aVF, aVL, V $\times 0.5$, $\times 1.0$, $\times 2.0$ >100 dB at 50 Hz or 60 Hz $0.67 \sim 40$ Hz (+3dB attenuation) 7 channels 12.5, 25 and 50 mm/s $30 \sim 300$ bpm ± 1 bpm or ± 1 %, whichever is greater Upper limit: $80 \sim 400$ bpm Lower limit: $20 \sim 150$ bpm ± 5 mV AC	
Input: Lead Choice: Gain Choice: CMRR (common mode rejection ratio): Frequency Characteristic: ECG Waveforms: Sweep Speed: HR Display Range: Accuracy: Alarm Limit Range Input Dynamic Range: Defibrillator Discharge: Differential Input Impedance:	5 lead or 3 lead ECG cable and standard AAMI line for connection I, II, III, aVR, aVF, aVL, V $\times 0.5$, $\times 1.0$, $\times 2.0$ >100 dB at 50 Hz or 60 Hz $0.67 \sim 40$ Hz (+3dB attenuation) 7 channels 12.5, 25 and 50 mm/s $30 \sim 300$ bpm ± 1 bpm or ± 1 %, whichever is greater Upper limit: $80 \sim 400$ bpm Lower limit: $20 \sim 150$ bpm ± 5 mV AC <5 sec >5 M O	
Input: Lead Choice: Gain Choice: CMRR (common mode rejection ratio): Frequency Characteristic: ECG Waveforms: Sweep Speed: HR Display Range: Accuracy: Alarm Limit Range Input Dynamic Range: Defibrillator Discharge: Differential Input Impedance: Bandwidth:	5 lead or 3 lead ECG cable and standard AAMI line for connection I, II, III, aVR, aVF, aVL, V $\times 0.5$, $\times 1.0$, $\times 2.0$ >100 dB at 50 Hz or 60 Hz $0.67 \sim 40$ Hz (+3dB attenuation) 7 channels 12.5, 25 and 50 mm/s $30 \sim 300$ bpm ± 1 bpm or ± 1 %, whichever is greater Upper limit: $80 \sim 400$ bpm Lower limit: $20 \sim 150$ bpm ± 5 mV AC <5 sec >5 M Ω 0 5 to 30 Hz (filter mode) 0.5 to 40 Hz (non filter	
Input: Lead Choice: Gain Choice: CMRR (common mode rejection ratio): Frequency Characteristic: ECG Waveforms: Sweep Speed: HR Display Range: Accuracy: Alarm Limit Range Input Dynamic Range: Defibrillator Discharge: Differential Input Impedance: Bandwidth:	5 lead or 3 lead ECG cable and standard AAMI line for connection I, II, III, aVR, aVF, aVL, V $\times 0.5$, $\times 1.0$, $\times 2.0$ >100 dB at 50 Hz or 60 Hz $0.67 \sim 40$ Hz (+3dB attenuation) 7 channels 12.5, 25 and 50 mm/s $30 \sim 300$ bpm ± 1 bpm or ± 1 %, whichever is greater Upper limit: $80 \sim 400$ bpm Lower limit: $20 \sim 150$ bpm ± 5 mV AC <5 sec >5 M Ω 0.5 to 30 Hz (filter mode), 0.5 to 40 Hz (non filter mode)	
Input: Lead Choice: Gain Choice: CMRR (common mode rejection ratio): Frequency Characteristic: ECG Waveforms: Sweep Speed: HR Display Range: Accuracy: Alarm Limit Range Input Dynamic Range: Defibrillator Discharge: Differential Input Impedance: Bandwidth: Electrode Offset Potential Tolerance:	5 lead or 3 lead ECG cable and standard AAMI line for connection I, II, III, aVR, aVF, aVL, V $\times 0.5$, $\times 1.0$, $\times 2.0$ >100 dB at 50 Hz or 60 Hz $0.67 \sim 40$ Hz (+3dB attenuation) 7 channels 12.5, 25 and 50 mm/s $30 \sim 300$ bpm ± 1 bpm or ± 1 %, whichever is greater Upper limit: $80 \sim 400$ bpm Lower limit: $20 \sim 150$ bpm ± 5 mV AC <5 sec >5 M Ω 0.5 to 30 Hz (filter mode), 0.5 to 40 Hz (non filter mode) ± 300 mV	
Input: Lead Choice: Gain Choice: CMRR (common mode rejection ratio): Frequency Characteristic: ECG Waveforms: Sweep Speed: HR Display Range: Accuracy: Alarm Limit Range Input Dynamic Range: Defibrillator Discharge: Differential Input Impedance: Bandwidth: Electrode Offset Potential Tolerance: Input Signal Range:	5 lead or 3 lead ECG cable and standard AAMI line for connection I, II, III, aVR, aVF, aVL, V $\times 0.5$, $\times 1.0$, $\times 2.0$ >100 dB at 50 Hz or 60 Hz $0.67 \sim 40$ Hz (+3dB attenuation) 7 channels 12.5, 25 and 50 mm/s $30 \sim 300$ bpm ± 1 bpm or ± 1 %, whichever is greater Upper limit: $80 \sim 400$ bpm Lower limit: $20 \sim 150$ bpm ± 5 mV AC <5 sec >5 M Ω 0.5 to 30 Hz (filter mode), 0.5 to 40 Hz (non filter mode) ± 300 mV ± 5 mV	

Meets the performance standards of ANS standardization voltage (section 4.2.2.9), in the ECG display, along with the ECG s references particular sections of ANSI/AA	I/AAMI EC13:2002. Instead a fixed, 1 cm reference bar ize setting expressed in mV/ MI EC13:2002.	of a 1 mV is always present /cm. The following
Respiration, leads-off sensing	Sinusoidal signal, 260 µ A	. 40.5 kHz
waveform.	Noise Suppression: RL	drive gain 44 dB
Noise suppression 4 1 2 1(b)	max max voltage 1.8 Vrm	
Tall T wave rejection $4.1.2.1(s)$	Exceede ANSI/AAM EC 1	2 Sect. 4 = 2 = 1 (a)
	Exceeds ANSI/AAIVII EC I.	5 Sect. 4.1.2.1(c),
	Recommended less than	1.2 mV I-Wave
	amplitude	
Heart Rate Averaging Method	Normally, heart rate is com	puted by averaging
4.1.2.1(d)	the 12 most recent RR inte	rvals. For runs of
	PVCs, up to 8 RR intervals	are averaged to
	compute the HR. If each of	4 consecutive RR
	intervals is greater than 12	00 ms (that is. rate
	less than 50 bpm) then the	e 4 most recent RR
	intervals are averaged to c	ompute the HR
	Thereafter computing by in	creasing 1 RR interval
	till recover 12 PP intervale	
	The freeb rote for displayin	
	The fresh rate for displayin	g HR value is 1
	second.	
Response to irregular	a) Ventricular bigeminy - 80) BPM
rhythm. 4.1.2.1(e)	b) Slow alternating ventricu	ılar bigeminy - 60
	BPM.	
	c) Rapid alternating vent	ricular bigeminy -
	120 BPM	
	d) Bi-directional systoles - 9	90 BPM
Heart rate meter response	HR change from 80 to 120	bom:
time $4 1 2 1(f)$	Average: 10 seconds	.
	HP change from 80 to 40 h	nm:
	Average: 10 accords	ipm.
	Average. To seconds	Average Time to Alerm
Heart rate meter response	0.5 mV	
time. 4.1.2.1(g)	2 mV	10 sec
	Waveform 4(b), Amplitude	10 sec
	1 mV	Average Time to Alarm
	2 mV	10 sec
	4 mV	10 sec
		10 sec
Pacemaker pulse rejection.	With the exceptions noted	below, the monitor
4.1.4.1, 4.1.4.3	will reject all pacemake	er pulses having
	amplitudes of ± 2 to ± 70	00 mV and pulse
	widths from 0.1 to 2.0 ms.	·
RESPIRATION		
Measure Method:	RA-LL impedance	
Range.	$0 \sim 120 \text{ rpm}$	
Acouracy		
Alorm Linner lower Limit		
Alarm Opper-lower Limit	Upper limit: $6 \sim 120$ rpm,	
	Lower limit: $3\sim$ 120 rpm	
Sweep Speed:	12.5 and 25mm/s	
NIBP		
Measuring Technology:	Automatic oscillating meas	urement
Cuff Inflating:	<30s (0~300 mmHa, sta	indard adult cuff)
Measuring Period [.]	AVF<40s	
Mode	Manual Auto STAT	
Measuring Interval in ALITO Mode	$2 \min_{\alpha \neq 1} A hro$	
	∠ IIIII/~4 IIIS	
Pulse Rate Range:	30 bpm \sim 250 bpm	

Measuring Range:	Adult/Pediatric ModeSYS $40 \sim 250 \text{ (mmHg)}$ DIA $15 \sim 200 \text{ (mmHg)}$ Neonatal ModeSYSSYS $40 \sim 135 \text{ (mmHg)}$
Resolution:	DIA 15~100 (mmHg) 1mmHg
Pressure Accuracy:	Maximum Mean error: ±5mmHg
Overpressure Protection:	Adult Mode 280(mmHg) Neonatal Mode 150 (mmHg)
Alarm Limit:	SYS 50~240 mmHg DIA 15~180 mmHg
TEMPERATURE	
Range:	25∼50 (°C)
Accuracy:	+ 0.2℃ (25.0~34.9℃)
-	_ + 0.1℃(35.0~39.9℃)
	$+0.2^{\circ}$ (40.0~44.9°C)
	$+ 0.3^{\circ}$ (45.0~50.0°C)
Display Resolution:	<u>1</u> 0.000 (+0.000000) 0.1℃
Alarm Upper-lower Limit:	Upper limit $0 \sim 50^{\circ}$
Channel	Lower minit 0^{-50}
Alarm Limit:	
	10~50 (())
	Anti motion SpO
SpO ₂ % Bange:	Anti-motion SpO ₂ $0 \sim 100\%$
	$0^{\prime} \sim 100\%$
SpO_2 Accuracy.	$\pm 2\%$ (70~100%, non-motion)
	<u>+</u> 3% (70~100%, motion)
Pulse Rate Range:	30-250 bpm
Pulse Rate Accuracy:	±2 bpm(non-motion)
Alarm Upper-lower Limit:	± 3 bpm (motion)
	Lower limit $70 \sim 100\%$
SnO_{2} Probe	Red light LED wavelength 660nm+5nm
000211000	Infrared light LED wavelength 940nm+10nm
IBP(OPTION)	
Measurement Range:	-10~300mmHg
Channel:	2 channels
Pressure transducer:	Sensitivity, 5µV/V/mmHg
Impedance range:	300~3000Ω
Transducer sites:	ART, PA,CVP, RAP, LAP, ICP
Resolution:	1mmHg
Accuracy:	±1mmHg or ±2%, whichever is greater
Alarm range:	-10~300mmHg
EtCO ₂ (OPTION)	
Mode of Sampling:	Sidestream or Mainstream
Principle of Operation:	Non-dispersive infrared (NDIR) single beam
00	optics, dual wavelength, no moving parts.
CO_2 measurement Range:	U to 150 mmHg (U to 19.7%, U to 20 kPa)
CO_2 Calculation Method:	DIFS (Body lemperature Pressure Saturated) 0.1 mmHz(0.60 mmHz)
	o. mm⊓g(o-osmm⊓g),

CO ₂ Accuracy: Sampling rate: Respiration Rate: Respiration Rate accuracy: Response Time:	0.25mmHg(70-150mmHg) $0 \sim 40 \text{ mmHg} \pm 2 \text{ mmHg}$ $41 \sim 70 \text{ mmHg} \pm 5\%$ of reading $71 \sim 100 \text{ mmHg} \pm 8\%$ of reading $101 \sim 150 \text{ mmHg} \pm 10\%$ of reading Above 80 breath per minute $\pm 12\%$ of reading 100Hz $2 \sim 150 \text{ bpm}$ $\pm 1 \text{ breath}$ <3 seconds - includes transport time and rise
	time
Inspired CO ₂ measurement Range:	$3\sim$ 50 mmHg
NETWORKING	
Wired Networking Wireless Networking	Industry standard: IEEE 802.3 wired network Connected bedside number: Up to 16 bedside monitors RJ45 interface or RS232 serial port Up to 100m indoors Frequency Range: 2.412~2.484 GHz Industry standard 802.11b/g wireless Supports TCP/IP and UDP/IP Protocols
POWER	
Source: AC Power: Battery:	External AC power and internal battery 100~240VAC, 50/60Hz, 150VA Rechargeable Sealed Lead Type: FB 1223 12v-2.3Ah Operating time under the normal condition: 1 hour
	Operating time after the first alarm of low battery: 10 minutes Manufacturer: Pilot Battery Co.,Ltd.
Charge Time:	Operating time after the first alarm of low battery: 10 minutes Manufacturer: Pilot Battery Co.,Ltd. 4 hours
Charge Time: Operating Time:	Operating time after the first alarm of low battery: 10 minutes Manufacturer: Pilot Battery Co.,Ltd. 4 hours 1 hour
Charge Time: Operating Time: ENVIRONMENTAL SPECIFICATIONS	Operating time after the first alarm of low battery: 10 minutes Manufacturer: Pilot Battery Co.,Ltd. 4 hours 1 hour Operating: 5~40 °C
Charge Time: Operating Time: ENVIRONMENTAL SPECIFICATIONS Temperature:	Operating time after the first alarm of low battery: 10 minutes Manufacturer: Pilot Battery Co.,Ltd. 4 hours 1 hour Operating: 5~40 °C Starsage: 10x 45 °C
Charge Time: Operating Time: ENVIRONMENTAL SPECIFICATIONS Temperature: Humidity Range:	Operating time after the first alarm of low battery: 10 minutes Manufacturer: Pilot Battery Co.,Ltd. 4 hours 1 hour Operating: 5~40 °C Storage: -10~45 °C Operating: ≤80 % Storage: ≤80 %
Charge Time: Operating Time: ENVIRONMENTAL SPECIFICATIONS Temperature: Humidity Range: RECORDER (OPTION)	Operating time after the first alarm of low battery: 10 minutes Manufacturer: Pilot Battery Co.,Ltd. 4 hours 1 hour Operating: $5\sim40$ °C Storage: $-10\sim45$ °C Operating: ≤ 80 % Storage: ≤ 80 %
Charge Time: Operating Time: ENVIRONMENTAL SPECIFICATIONS Temperature: Humidity Range: RECORDER (OPTION) Record Width: Paper Speed: Trace:	Operating time after the first alarm of low battery: 10 minutes Manufacturer: Pilot Battery Co.,Ltd. 4 hours 1 hour Operating: $5\sim40$ °C Storage: $-10\sim45$ °C Operating: ≤ 80 % Storage: ≤ 80 % 48 (mm) 25 (mm/s) 2 or 3