Patient Monitor



Ver. 2.62

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1. BASIC

1.1 CE Standard Information

1.2 Read before Use

Warranty Period
Warning, Caution, Note
General Precaution on Environment
General Precaution on Electric Safety
Equipment Connection, Maintenance & Washing Equipment Connection

1.3 Product Components

Product Outline
Principal Characteristics of Product
Product Configuration and Option Product
Product Body Configuration

1.4 Function and Key

External Function Operation Key

1.5 Standard Power Supply Application

1.6 Battery Power Supply Application

1.7 General Menu Operation

Screen Composition Menu Selection Menu Composition

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1.1 CE Standard Information

Electromechanical safety standards met:

- EN 60601-1: 1990 + A1:1993 + A2: 1995 Medical Electrical Equipment, Part 1, General Requirements for Safety.
- IEC/EN 60601-1-2 :2001 Electromagnetic compatibility -Requirements and tests.
- EN 1060-1:1995 Non-invasive sphygmomanometers Part 1: General requirements
- EN 1060-3:1997 Non-invasive sphygmomanometers Part 3: Supplementary requirements for electro-mechanical blood pressure measuring systems
- EN ISO 9919:2005 Medical electrical equipment Particular requirements for the basic safety and essential performance of pulse oximeter equipment for medical use (ISO 9919:2005)
- EN 60601-2-27:2006 Medical electrical equipment Part 2-27: Particular requirements for the safety, including essential performance, of electrocardiographic monitoring equipment
- EN 60601-2-30:2000 Medical electrical equipment Part 2-30: Particular requirements for the safety, including essential performance, of automatic cycling non-invasive blood pressure monitoring equipment
- EN 12470-4:2000 Clinical thermometers Part 4: Performance of electrical thermometers for continuous measurement
- EN 60601-2-49:2001 Medical electrical equipment Part 2-49: Particular requirements for the safety of multifunction patient monitoring equipment

1.2 Read before Use

BIONET services are always available to you.

The followings are address and phone number for contacting information, services, and product supplies.

How to Contact Us

Product Supply Information

Bionet America Inc. 2691 Dow Ave Ste B Tustin, CA 92780 Phone: 714-734-1760 Fax: 714-734-1761

E-mail: sales@bionetUS.com,

Consumable Supplies

Request

Phone: 714-734-1760 - USA

Phone: 82-2-6300-6477 - Intenational

A/S and Technical Support

For any technical inquiries or repair on the equipment,

Phone: 714 -734-1760

E-mail: service@bionetUS.com

Home Page

URL: http://wwwbionetUS.com

X In the event of malfunction or failure, contact us along with the model name, serial number, and product name of the equipment.

※ If you need the supply circuit diagram, component list, description and calibration instruction
etc. you can contact us we will provide you with it.

The information in this manual only applies to BM3 patient monitor software version 1.10F. Due to continuing product innovation, specifications in this manual are subject to change without notice.

Warranty Period

- This product is manufactured and passed through strict quality control and through inspection.
- We provide a 1-year warranty period for main body, with 6 months warranty for accessories.
- We will repair or replace any part of the BM3 found to be defective in usual operating circumstance for free to you.
- This warranty does not apply to any defect caused by improper use, misuse or abuse.

Warning, Caution, Note

For special emphasis on agreement, terms are defined as listed below in user's manual. Users should operate the equipment according to all the warnings and cautions.

Warning

To inform that it may cause serious injury or death to the patient, property damage, material losses against the "warning" sign

Caution

To inform that it may cause no harm in life but lead to injury against the "caution" sign

Note

To inform that it is not dangerous but important "note" sign for proper installation, operation, and maintenance of the equipment.

General Precaution on Environment

- Do not keep or operate the equipment in the environment listed below.

Bono	r keep of operate the equipment	it iii tilo oliviioliiiloli	t notoa bolow.
	Avoid placing in an area exposed to moist. Do not touch the equipment with wet hand.		Avoid exposure to direct sunlight
	Avoid placing in an area where there is a high variation of temperature. Operating temperature ranges from 10(C to 40(C. Operating humidity ranges from 30% to 85%.		Avoid in the vicinity of Electric heater
	Avoid placing in an area where there is an excessive humidity rise or ventilation problem.		Avoid placing in an area where there is an excessive shock or vibration.
	Avoid placing in an area where chemicals are stored or where there is danger of gas leakage.		Avoid being inserted dust and especially metal material into the equipment
60 h	Do not disjoint or disassemble the equipment. We take no responsibility for it.	ST COLOR	Power off when the equipment is not fully installed. Otherwise, equipment could be damaged.

CAUTIONS

Before Installation

Compatibility is critical to safe and effective use of this device. Please contact your local sales or service representative prior to installation to verify equipment compatibility.

Defibrillator Precaution

Patient signal inputs labeled with the CF and BF symbols with paddles are protected against damage resulting from defibrillation voltages. To ensure proper defibrillator protection, use only the recommended cables and lead wires.

Proper placement of defibrillator paddles in relation to the electrodes is required to ensure successful defibrillation.

Disposables

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

Disposal of your old appliance



- 1. When this crossed out wheeled bin symbol is attached to a product it means the product is covered by the European Directive 2002/96/EC.
- All electrical and electronic products should be disposed of separately from the municipal waste stream via designated collection facilities appointed by the government or the local authorities.
- 3. The correct disposal of your old appliance will help prevent potential negative consequences for the environment and human health.
- 4. For more detailed information about disposal of your old appliance, please contact your city office, waste disposal service or the shop where you purchased the product.

WARNING

This product contains a chemical known to the State of California to cause cancer, birth defects, or other reproductive harm.

Electrocute Precautions

To prevent skin burns, apply electrocute electrodes as far as possible from all other electrodes, a distance of at 15 cm/6 in, is recommended.

EMC

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

Also, keep cellular phones to other telecommunication equipment away from the monitor.

CAUTIONS

Instruction for Use

For continued safe use of this equipment, it is necessary that the instructions are followed. However, instructions listed in this in no way supersede established medical practices concerning patient care.

Loss of Data

Should the monitor at any time temporarily lose patient data, the potential exists that active monitoring is not being done. Close patient observation or alternate monitoring devices should be used until monitor function is restored.

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

Maintenance

Regular preventive maintenance should be carried out annually (Technical inspections). You are responsible for any requirements specific to your country.

MPSO

The use of a multiple portable socket outlet (MPSO) for a system will result in an enclosure leakage current equal to the sum of all individual earth leakage currents of the system if there is an interruption of the MPSO protective earth conductor. Do not use an additional extension cable with the MPSO as it will increase the chance of the single protective earth conductor interruption.

Negligence

BIONET does not assume responsibility for damage to the equipment caused by improperly vented cabinets, improper or faulty power, or insufficient wall strength to support equipment mounted on such walls.

NOTES

Power Requirements

Before connecting the device to the power line, check that the voltage and frequency. Ratings of the power line are the same as those indicated on the unit's label. If this is not the case, do not connect the system to the power line until you adjust the unit to match the power source. In U.S.A, if the installation of this equipment will use 240V rather than 120V, the source must be a center-tapped, 240V, single-phase circuit.

Restricted Sale

U.S.A federal law restricts this device to sale by or on the order of a physician.

Supervised Use

This equipment is intended for use under the direct supervision of a licensed health care practitioner.

Ventilation Requirements

Set up the device in a location which affords sufficient ventilation. The ventilation openings of the device must not be obstructed. The ambient conditions specified in the technical specifications must be ensured at all times.

- •Put the monitor in a location where you can easily see the screen and access the operating controls.
- •This product is protected against the effects of cardiac defibrillator discharges to ensure proper recovery, as required by test standards. (the screen may blank during a defibrillator discharge but recovers within second as required by test standards.)

Reference Literature

Medical Device Directive 93/42/EEC

EN 60601-1/1990 +A1: 1993 +A2: 1995: Medical electrical equipment.

General requirements for safety

EN 60601-1-1/9. 1994 +A1 12.95: General requirements for safety.

General Precaution on Electric Safety

Warning

Check the item listed below before operating the equipment.

- 1. Be sure that AC power supply line is appropriate to use. (AC100 240V)
- 2. Be sure that the power source is the one supplied from Bionet. (DC18V, 2.8A, MW160KA1803)
- 3. Be sure that the entire connection cable of the system is properly and firmly fixed.
- 4. Be sure that the equipment is completely grounded. (If not, there might be the problem occur in the product.)
- 5. The equipment should not be placed in the vicinity of electric generator, X-ray, broadcasting apparatus to eliminate the electric noise during operation. Otherwise, it may cause incorrect result.

Note

The Equipment should be placed far from generator, X-ray equipment, broadcasting equipment or transmitting wires, so as to prevent the electrical noises from being generated during the operation, When these devices are near the Equipment, it can produce inaccurate measurements. For BM3, both independent circuit and stable grounding are essentially required. In the event that the same power source is shared with other electronic equipment, it can also produce inaccurate output.

Warning

Do not contacts with the patient while operate the machine It may cause serious danger to the users. Use only the provided cable.

Warning

In case the Equipment does not operate as usual or damaged, do not use on patient, and contact to the medical equipment technician of the hospital or the equipment supply division.

Note

BM3 is classified as follows:

- BM3 classifies as Class ${\bf I}$, BF ${\bf \&}$ CF concerning electric shock. It is not proper to operate this Equipment around combustible anesthetic or dissolvent.
- Noise level is B class regarding IEC/EN 60601-1 and the subject of Nose is B level concerning IEC/EN60601-1-2.

Equipment Connection

Caution

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

Manufacturer's declaration - electromagnetic emission

The BM3 system is intended for use in the electromagnetic environment specified below. The customer or the user of BM3 system should assure that it is used in such an environment				
Emission test	Compliance	Electromagnetic environment - guidance		
RF emissions CISPR 11	Group 1	The BM3 system uses RF energy only for its internal function. Therefore. Its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment		
RF emissions CISPR 11	Class B	The BM3 system is suitable for use in all establishm ents other than domestic and those directly connect		
Harmonics emission IEC 61000-3-2	А	ed to the public low-voltage power supplies building s used for domestic purposes.		
Voltage fluctuation IEC 61000-3-3	Complies			

Manufacturer's declaration - electromagnetic immunity The BM3 system is intended for use in the electromagnetic environment

The BM3 system is intended for use in the electromagnetic environment specified below.					
The customer or the user of the BM3 system should assure that it is used in such an environment					
Immunity test	IEC 60601 Test level	Compliance level	Electromagnetic Environment -guidance		
Electrostatic disc harge (ESD) IEC 61000-4-2	6 kV Contact 8 kV Air	6 kV Contact 8 kV Air	Floors should be wood, con crete or ceramic tile. If floor s are covered with synthetic material, the relative humidit y should be at least 30 %		
Electrical fast Transient / burst IEC 61000-4-4	2kV for power supply lines 1kV for input/output lines	s 1kV for input/output lines	Mains power quality should be that of a typical commerc ial or hospital environment.		
Surge IEC 61000-4-5	1 kV differential mode 2 kV common mode	1 kV differential mode 2 kV common mode	Mains power quality should be that of a typical commer cial or hospital environment.		
Power frequency (50/60Hz) Magnetic field IEC 61000-4-8	3.0 A/m	3.0 A/m	Power frequency magnetic fi elds should be at levels cha racteristic of a typical locatio n in a typical commercial or hospital environment.		
s on power supply input lines IEC 61000-4-11	for 0.5cycle 40% <i>U</i> τ (60% dip in <i>U</i> τ) for 5 cycle 70% <i>U</i> τ (30% dip in <i>U</i> τ) for 25 cycle <5% <i>U</i> τ (<95% dip in <i>U</i> τ) for 5 s) for 5 s	Mains power quality should be that of a typical commerc ial or hospital environment. If the user of the BM3 system requires continued op eration during power mains interruptions, it is recommend ed that the BM3 system be powered from an uninterruptible power supply or a battery		
Note: Ut is the a.c. mains voltage prior to application of the test level.					

The BM3 system is intended for use in the electromagnetic environment specified below.				
The customer or the user of the BM3 system should assure that it is used in such an environment				
		Electromagnetic environment -guidance		
0 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	Test level	0.1/	D (11 1 1 1 DE 1 1 1	
Conducted RF IEC 61000-4-6	3 Vrms 150 kHz to 80 MH z	3 Vrms 150 kHz to 80 MHz	Portable and mobile RF communications e quipment should be used no closer to any part of the BM3 system, including cables, t han the recommended separation distance calculated from the equation applicable to t he frequency of the transmitter.	
			Recommended separation distance	
			$d = \left[\frac{3.5}{V_1}\right] \sqrt{P}$	
Radiated RF	3 V/m	3 V/m	Recommended separation distance	
IEC 61000-4-3	80.0 MHz to 2.5 G Hz	80.0 MHz to 2.5 G Hz	$d = [\frac{3.5}{E_1}]\sqrt{P}$ 80 MHz to 800 MHz $d = [\frac{7}{E_1}]\sqrt{P}$ 800 MHz to 2,5 GHz	
			$d = [\frac{r}{E_1}] \sqrt{P}$ 800 MHz to 2,5 GHz	
			Where <i>P</i> is the maximum output power rat ing of the transmitter in watts (W) according to the transmitter manufacturer and <i>d</i> is the recommended separation distance in meters (m). Field strengths from fixed RF transmitters, as deter-mined by an electromagnetic site survey, (a) Should be less than the compliance level in each frequency range (b).	
			Interference may occur in the vicinity of equipment marked with the following symbol:	

Note 1) Ut is the A.C. mains voltage prior to application of the test level.

Note 2) At 80 MHz and 800 MHz, the higher frequency range applies.

Note 3) These guidelines may not apply in all situations. Electromagnetic propagation is affected by a bsorption and reflection from structures, objects and people.

a Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be pred icted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitt ers, an electromagnetic site survey should be considered. If the measured field strength in the location in which the EUT is used exceeds the applicable RF compliance level above, the EUT should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as re-orienting or relocating the EUT.

b Over the frequency range 150 kHz to 80 MHz, field strengths should be less than [V1] V / m.

Recommended Separation Distances Between Portable and Mobile RF Communications Equipment and the **BM3** system.

The **BM3** system is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The user of the **BM3** system can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the BM3 system as recommended below, according to the maximum output power of the communications equipment.

Rated maximum output power (W) of transmitter	Separation distance (m) according to frequency of transmitter			
	150 kHz to 80 MHz	80 MHz to 800 MHz	800 MHz to 2.5 GHz	
0.01	0.12	0.12	0.23	
0.1	0.37	0.37	0.74	
1	1.17	1.17	2.33	
10	3.70	3.70	7.37	
100	11.70	11.70	23.30	

For transmitters rated at a maximum output power not listed above, the recommended separation dist ance (d) in meters (m) can be estimated using the equation applicable to the frequency of the transmitter, where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer.

Note 1: At 80 MHz and 800 MHz, the separation distance for the higher frequency range applies

Note 2: These guidelines may not apply in all situations. Electromagnetic propagation is affected by a bsorption and reflection from structures, objects, and people.

Immunity and Compliance Level				
Immunity test	IEC 60601 Test Level	Actual Immunity Level	Compliance Level	
Conducted RF	3 Vrms, 150 kHz to 80	3 Vrms, 150 kHz to 80	3 Vrms, 150 kHz to 80	
IEC 61000-4-6	MHz	MHz	MHz	
Radiated RF	3 V/m, 80 MHz to 2.5	3 V/m, 80 MHz to 2.5	3 V/m, 80 MHz to 2.5	
IEC 61000-4-3	GHz	GHz	GHz	

Guidance and manufacturer's declaration - electromagnetic immunity

The **BM3** system is intended for use in the electromagnetic environment specified below. The customer or the user of the **BM3** system should assure that it is used in such an environment

Immunity test	IEC 60601 Test level	Compliance level	Electromagnetic environment -guidance
Conducted RF	3 Vrms	3 Vrms	BM3 system must be used only in a shield
IEC 61000-4-6		150 kHz to 80 MHz	ed location with a minimum RF shielding ef
	z		fectiveness and, for each cable that enters
			the shielded location with a minimum RF s
			hielding effectiveness and, for each cable t
5 " 55	0.14	0.1//	hat enters the shielded location
Radiated RF	3 V/m	3 V/m	Field strengths outside the shielded locatio
IEC 61000-4-3			n from fixed RF transmitters, as determine
	Hz	Hz	d by an electromagnetic site survey, should
			be less than 3V/m.a
			Interference may occur in the vicinity of eq
			uipment marked with the following symbol:
			4 4
			[((•))]
			(<u> </u>
			=

Note 1) These guidelines may not apply in all situations. Electromagnetic propagation is affected by a bsorption and reflection from structures, objects and people.

Note 2) It is essential that the actual shielding effectiveness and filter attenuation of the shielded loc ation be verified to assure that they meet the minimum specification.

a- Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephone s and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be pr edicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF trans mitters, an electromagnetic site survey should be considered. If the measured field strength outside th e shielded location in which the EUT is used exceeds 3V/m, the EUT should be observed to verify n ormal operation.

If abnormal performance is observed, additional measures may be necessary, such as relocating the EUT or using a shielded location with a higher RF shielding effectiveness and filter attenuation.

Note

For Type A Professional ME Equipment intended for use in domestic establishment instructions for use includes a warning:

This ME equipment is intended for use by professional healthcare personnel only.

Biocompatibility

When used as intended, the parts of the product described in this operator manual, including accessories that come in contact with the patient during the intended use, fulfill the biocompatibility requirements of the applicable standards. If you have questions about this matter, please contact BIONET or its representatives.

Maintenance and Washing Equipment Connection

Using various methods can clean BM3 and its accessories. Please follow the methods mentioned below to avoid unnecessary damage or contamination to the Equipment.

We do not repair with free of charge regardless of warranty period if it is contaminated or damaged with using dangerous material not designated for washing.

Cleaning Applied Parts

Do not permit any liquid to enter the monitor case and avoid pouring it on the monitor while cleaning. Do not allow water or cleaning solution to enter the connectors of jack cover.

Recommended cleaning agents:

Alcohol (Ethanol 70%, losopropanol 70%, Window cleaner)

Ammonias (Dilution of ammonia <3%, Window cleaner)

Tensides (dishwasher detergents) (Edisonite schnellreiniger[®], Alconox[®])

Cables and Leadwires

CAUTION

Do not use acetone or ketone solvents for cleaning; do not use an autoclave or steam cleaner.

Cables and leadwires can be cleaned with a warm, damp cloth and mild soap, or isopropyl alcohol wipes. For more intensive disinfecting (near sterile) Ethylene Oxide (ETO) is acceptable but will reduce the useful lifetime of the cable or leadwire.

CAUTION

The decision to sterilize must be made per your institution's requirements with an awareness of the effect on the integrity of the cable or leadwire.

Note

The Equipment needs safety inspection once a year. Please refer to user's guide or service manual for the examine specifications.

Please check carefully both frame and sensor, after cleaning the Equipment, Do not use the equipment that is worn out or damaged.

At least once a month, clean and wipe off the frame by using the soft cloth after wetting it with water and alcohol. Do not use lacquer, thinner, ethylene, and oxidizer which may leads damage to the equipment.

Make sure both cables and accessories are free of dust or contaminants, and wipe them off with soft cloth wetted with warm water (40°), and at least once a week, clean them by using the clinical alcohol.

Do not submerge the accessories under any liquid or detergent. Also, make sure any liquid not to penetrate into the Equipment or probe.

Disinfecting

Do not mix disinfecting solutions (such as bleach and ammonia) as hazardous gases may result. Clean equipment before disinfecting.

Recommended disinfecting agents:

Aldehyde based (Cidex[®] activated dialdehyde solution, Gigasept)

Alcohol base (Ethanol 70%, Isopropanol 70%, Spitacid[®], Streilium fluid[®], Cutasept[®], Hospisept[®], Tinktur forte, Sagrosept[®], Kodan[®])

Caution

Do not dispose single use probe to any hazard place, Always think about environmental contamination.

Caution

There is back-up battery on board inside system. When users dispose this battery, Please waste proper place for environmental protection.

Warning

Check the electrodes of batteries before changing them.

- Operate BM3 with internal electric power supply when unsure of external ground connection or installation occur.
- · Remove the 1st Battery when not using equipment for a while without any damage.

For other applied parts such as temperature sensors, pulse oximetry probes, and NBP cuffs, you must consult the manufacturer for cleaning, sterilization, or disinfecting methods.

1.3 Product Components

Product Outline

BM3 monitor is a product used for monitoring biological information and occurrence of a patient. Main function ns of the product include displaying information such as ECG, respiration,EtCO2, SpO2, NIBP and temperature on its LCD screen and monitoring parameter, and alarming. It also prints out waves and parameters via a printer.

Principal Characters of Product

BM3 is a small-size multifunctional monitoring equipment for a patient designed to an easy usage during movement. It features devices for DC power supply (DC 18V, MW160) as well as installing its handle to the patient's bed. The equipment also measures major parameters such as ECG, SpO2, NIBP, temperature and pulse, displaying it on a 7-inch color LCD screen. It also enables users to check waves and parameters and other vital signs of a patient via the 58mm thermal printer and monitor the patient by the remote-controlled alarm system. It also enables to build a central monitoring system by linking devices used for separate patients so that one can monitor several patients at a time.

Warning

Use only the supplement accessories provided by us. Otherwise, patient and user may be exposed to danger.

We strongly recommend you use only the authorized accessories which we supply.

Warning

BEFORE USE — Before putting the system into operation visually inspect all connecting cables for signs of damage. Damaged cables and connectors must be replaced immediately. Before using the system, the operator must verify that it is in correct working order and operating condition. Periodically, and whenever the integrity of the product is in doubt, test all functions.

Product Configuration

1. Main body of BM3 Monitor	1 EA
2. 3-Lead Patient Cable	1EA
3. Disposable electrodes	10 EA
4. NIBP extension hose (3M long)	1EA
5. Adult cuff (27.5~36.5cm)	1EA
6. SpO2 extension cable (2M)	1EA
7. Reusable Adult SpO ₂ Probe	1 EA
8. DC Adaptor (MW160 made in Bridge power Co., Ltd.)	1 EA

Option Product

- 1. Temperature
- 2. 5-Lead Patient Cable
- 3. Thermal printer and Thermal Paper
- 4. EtCO2 Module

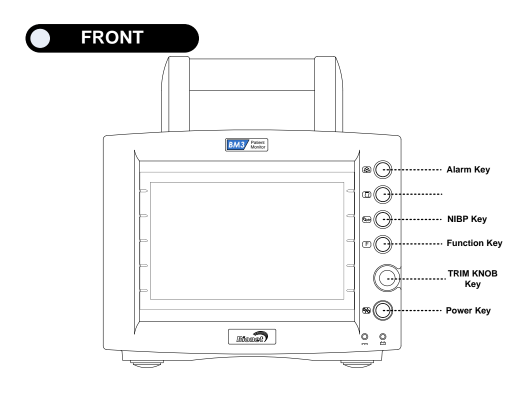
Warning

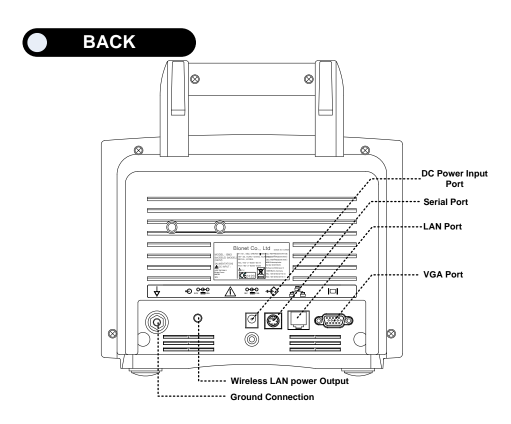
In order to avoid electrical shock, do not open the cover. Disassembling of the equipment should be done only by the service personnel authorized by BIONET

Warning

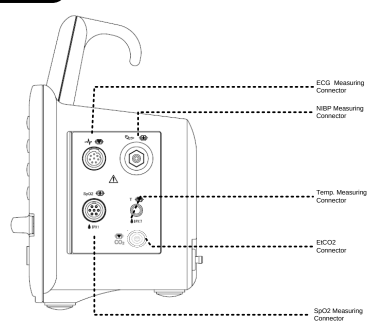
Users must pay attention on connection any auxiliary device via LAN port or nurse calling. Always consider about summation of leakage current, please check if the auxiliary device is qualified by IEC 60601-1, or consult your hospital biomedical engineer.

Product Body Configuration

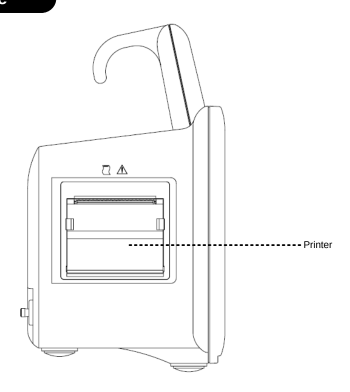




Right Side



Left Side



Accessories

ECG Cable + Extension Cable



SpO₂ Cable + Extension Cable



NIBP Cuff+ Extension cable



Temperature sensor (Option)



Equipment Sign



ATTENTION:

Consult accompanying documents



Defibrillator-proof type CF applied part :

Insulated (floating) applied part suitable for intentional external and internal application to the patient including direct cardiac application. "Paddles" outside the box indicate the applied part is defibrillator proof.

Medical Standard Definition:

F-type applied part(floating/insulated) complying with the specified requirements of IEC 60601-1/UL 2601-1/CSA 601.1

Medical Standards to provide a higher degree of protection against electric shock tan that provided by type BF applied parts.



Defibrillator-proof type BF applied part :

Insulated (floating) applied part suitable for intentional external and internal application to the patient excluding direct cardiac application. "Paddles" outside the box indicate the applied part is defibrillator proof.

Medical Standard Definition:

F-type applied part (floating/insulated) complying with the specified requirements of IEC 60601-1/UL 2601-1/CSA 601.1

Medical Standards to provide a higher degree of protection against electric shock than that provided by type B applied parts.

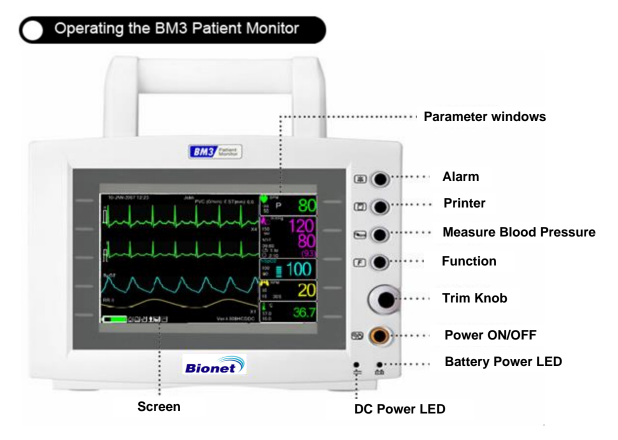
	Ground
O	Output port
5V === 0.9A	DC Power Output
	Printer
	VGA Output
	Serial Port
	LAN Port
\longleftrightarrow	AUX Connector Port
	DC Input Indicator
- +	Battery Operation Indicator
18V == 2.8A	DC Power Input port

	NIBP
Т	Temperature
PR	Pulse Rate
14	Respiration
√√~	ECG
	Heart Pulse
	Alarm Off
F	Function
•	Power On
·	Power Off

1.4 Function and Key

External Function

The front panel of this product consists of an LCD screen and five function keys and one trim knob.



Operation Key

- 1. Power: Switches on and off the Power.
- 2. Function Key
- 3. Blood Pressure: Manually completes measuring blood pressure.
- 4. Printer: Prints out the waves selected from the menu until the key is pressed to stop.
- 5. Alarm: Stop alarm sound.

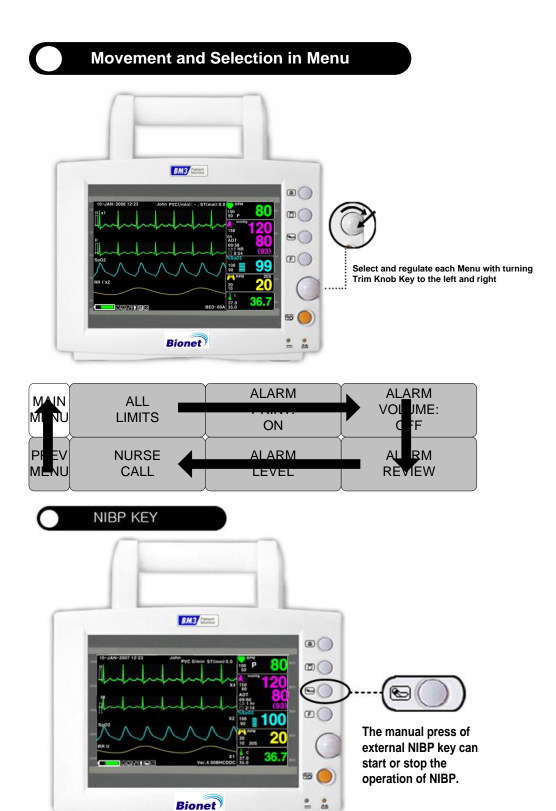
First press stops the current alarm for one minute

Second press stops the all alarm for two minutes.

Third press stops the all alarm off.

Fourth press makes the alarm back to the original setting.

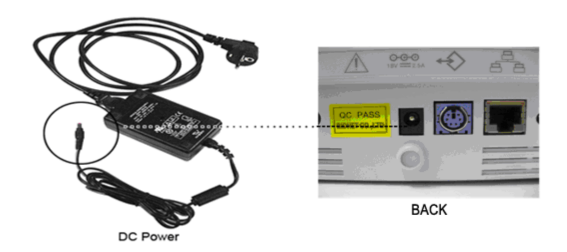
6. Trim Knob: This key is used to select menu by turning it clock or anticlockwise to move cursors.

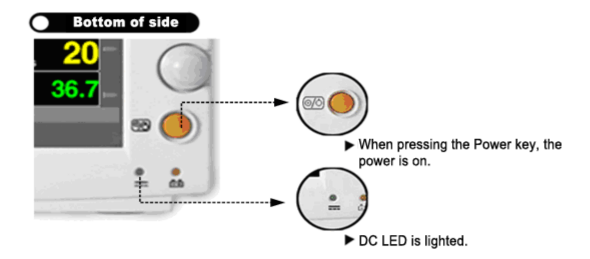


1.5 Standard Power Supply Application

DC Power

DC Power LED is lighted on when the DC Power is plugged into the inlet at the back of the product. A press of power key makes the machine ready for use.





Warning

This equipment must only be connected to a supply mains with protected earth.

1.6 Battery Power Supply Application

Battery power can be supplied for enabling a portable use or a use during DC power failure.

Operation

- 1. Battery Power LED is lighted on when the machine is in use.
- 2. The DC/battery power is only sustainable for 1 hour.
- 3. Battery is automatically charged when the machine is connected to DC Power Supply. Battery LED is lighted on after blinking.
- 4. The charging status of the batteries is displayed with 5 green boxes, each indicating a different charging
- . (0% -> 25% -> 50% -> 75% -> 100%)
 - Battery: 031PpTC(3ICR19/65)(10.8V, 2150mAh/23.22Wh)

The Lithium-Ion battery is a rechargeable battery containing Lithium-Ion cells. Each battery contains an integrated electronic fuel gauge and a safety protection circuit.







5. The discharge condition of battery is indicated with on of 5 yellow boxes, each box showing a different level of charge available.

When remained battery is less than 25%, the battery icon box is turned to red one with blink. The device will be turned off automatically after 5 minutes from that warning sign. In case of that warning sign with red and blink at icon box, charge the device immediately with DC power adaptor which is provided from BIONET.



- -Battery charging time: More than 6 hours
- -Continuous battery use time: Lowest 1 hour to highest 2 hours continuous use (buffering)

Warning Check the electrodes of batteries before charging them.

6. Battery status indication: When battery is apart from equipment and out of order, it is shown by a red `X' as shown below.



7. Low power supply: When you use the power of less than 16V, the battery indication disappears and the "LOW" indication is active.



Display of Low power supply

Note	
Battery is not charged when the LOW power is used.	

The Impact of Lithium-Ion Battery Technology on the Battery

The following are the key points you should know about Lithium-Ion battery technology:

The battery will discharge on its own, even when it is not installed in a monitor. This discharge is the result of the Lithium-Ion cells and the bias current required for the integrated electronics.

By the nature of Lithium-Ion cells, the battery will self-discharge.

The self-discharge rate doubles for every 10°C (18°F) rise in temperature.

The capacity loss of the battery degrades significantly at higher temperatures.

As the battery ages, the full-charge capacity of the battery will degrade and be permanently lost. As a result, the amount of charge that is stored and available for use is reduced.

Conditioning Guideline

the battery in the monitor full charged and discharged every six months and condition it using the battery charger.

Storage Guideline

Store the battery outside of the monitor at a temperature between 20°C to 25°C (68°F to 77°F). When the battery is stored inside a monitor that is powered by an AC power source, the battery cell temperature increases by 15°C to 20°C (59°F to 68°F) above the room's ambient temperature. This reduces the life of the battery.

When the battery is stored inside a monitor that is continuously powered by an AC power source and is not powered by battery on a regular basis, the life of the battery may be less than 12 months. BIONET recommends that you remove the battery and store it near the monitor until it is needed for transport.

How to Recycle the Battery

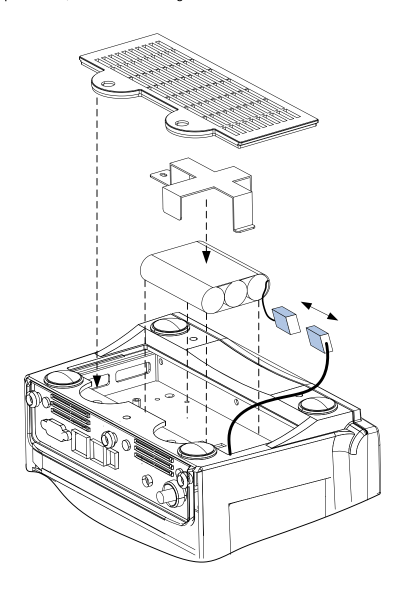
When the battery no longer holds a charge, it should be replaced. The battery is recyclables. Remove the old battery from the monitor and follow your local recycling guidelines.

WARNING

EXPLOSION HAZARD —

DO NOT incinerate the battery or store at high temperatures. Serious injury or death could result.

To insert and remove the battery pack.
Assembly or replacement, as shown in the figure below.



1.7 DISPLAY MODE (MONITOR OR SPOT)

You can selected display mode in user service (Monitoring and Spot). MONITOR: Refer to the Monitor chapter of this manual for details. SPOT: Refer to the SPOT chapter of this manual for details.

MAIN MENU	SET UNIT NAME	SET BED NUMBER : 00A	DISPLAY MODE: MONITOR
PREV	SYSTEM	AC FILTER:	W-LAN:
MENU		50HZ	OFF
MAIN MENU	SET UNIT NAME	SET BED NUMBER : 00A	DISPLAY MODE: SPOT
PREV	SYSTEM	AC FILTER:	W-LAN:
MENU		50HZ	OFF

MONITORING MODE

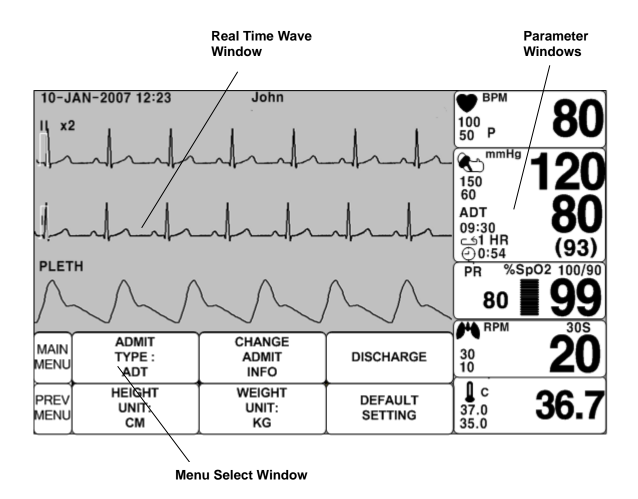
- 1. General Operation
- 2. Patient/Data Management
 - 3. Setup
 - 4. Trend
 - 5. ECG
 - 6. SpO2
 - 7. Respiration
 - 8. NIBP
 - 9. Temperature
 - 10. PRINT
 - 11. Message List
 - 12. Default Setting Value

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1. General Operation

1.1 General Manu Operation

Screen Composition

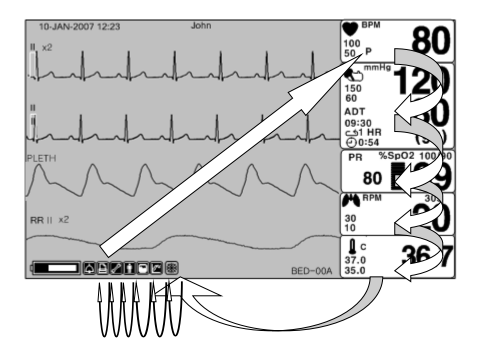


Real Time Wave Window: Displays measured results by up to three waves.

Menu Select Window: Menus appear when they are activated..

Parameter Window: Measured and setup data are displayed in five windows.

Menu Selection





Turn or press the knob.

When the Trim Knob Key is turned, menus are selected in the order indicated above. The above screen shows that the MORE menus is selected. The menus move to the right in the order of MORE MENU \rightarrow ECG \rightarrow NIBP \rightarrow SpO2 \rightarrow (RESP/EtCO2) \rightarrow TEMP. An inactivated window is jumped off.

Menu Composition

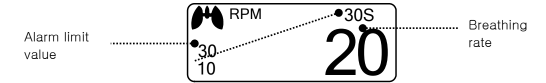
More Menu Window

When the additional menu is selected it will set and cancel the functions.

MAIN MENU	DISPLAY		USER SERVICE
PREV	SOUND:	DEMO:	MAKER
MENU		ON	SERVICE

Numerical value sign widow

This window displays a measured parameter, function setup, and the boundary of parameter values.



Menu selection by using Trim Knob key

As the key is turn to the right, the menu selection moves clockwise. As the key is turn to the left, the menu selection moves counterclockwise. The menu selection is activated when you depress Trim Knob key.

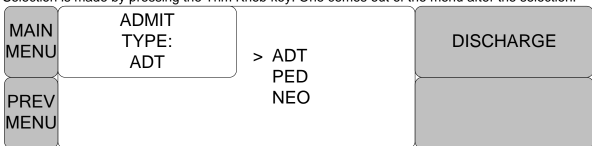
MAIN MENU	DISPLAY		USER SERVICE
PREV	KEY SOUND: ON	DEMO : ON	MAKER SERVICE

Menu selection with arrows

Upward Movement: Turns the Trim Knob key to the left.

Downward Movement: Turns the Trim Knob key to the right.

Selection is made by pressing the Trim Knob key. One comes out of the menu after the selection.

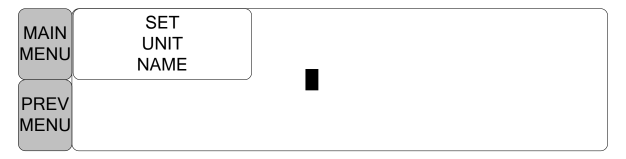


When moving the within quadrilateral, the letter reverses, and the numeric value reflects immediately.

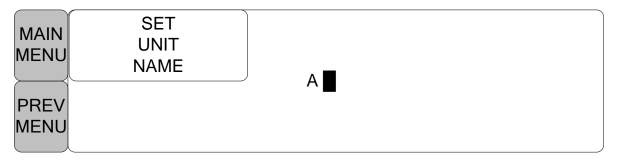
MAIN MENU	QRS VOLUME : OFF	>	OFF 10% 20%	60% 70% 80%	
PREV			30%	90%	
			40%	100%	
MENU			50%		

Word feature menu

The following figure shows the screen where the word sequence menu is activated within the word sequence correction menu. Here, the cursor moves over the words when the Trim Knob key is turned in the clockwise direction.



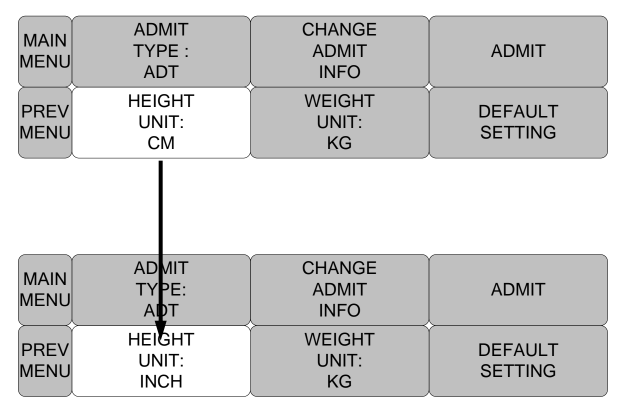
The above figure shows how the cursor moves on the screen. The cursor moves according to the direction the Trim Knob Key is turned. Press the Trim Knob key if you want to change a letter currently on the screen.



The above figure shows how the cursor is selected to change a letter. Right-hand turning of the Trim Knob Key makes it possible to select in the order of 0-9,A-Z, and a blank, while left-hand turning makes the movement in the opposite direction. Once a letter or a number is selected, the screen comes back to the condition where the same process of selection can be made. One may move to the menu item in the left of the screen to end the process, which is completed by pressing Trim Knob Key. After completion, the screen comes back to the earlier picture.

Operation menu

The setup value changes without a selection when the menu is moved.



2. PATIENT/DATA MANAGEMENT

2.1 ADMIT

ADMIT TYPE CHANGE ADMIT INFO DISCHARGE **ADMIT HEIGHT** WEIGHT **DEFAULT SETTING**

2.2 ALARM

ALL LIMITS ALARM PRINT ALARM VOLUME PARAMETER LEVEL ARRHYTH LEVEL ALARM REVIEW **ALARM LIST** SAVE CONDITION **NURSE CALL**

2.1 ADMIT



ADMIT TYPE: Set the exercise environment of equipment in discharge status.

CHANGE ADMIT INFO: The CHANGE ADMIT INFO option allows you to change or enter information pertinent to the monitored patient.

ADMIT: Depending on how your monitor is set up, you will see either ADMIT patient or new case DISCHARGE: This menu option indicates that patient is admitted. You select it to discharge the patient.

HEIGHT, WEIGHT UNIT: these options change the units of measure for height and weight. DEFAULT SETTING: Configure alarms, set alarm limits, and establish display defaults to be recalled whenever a discharge is performed.

MAIN MENU	ADMIT TYPE: ADT	CHANGE ADMIT INFO	DISCHARGE
PREV MENU	HMIT:	WEIGHT UNIT: KG	DEFAULT SETTING

ADMIT TYPE

Set the exercise environment of equipment in discharge status. ADUL: ADULT // PED: PEDIATRIC // NEO: NEONATE

100.10	ADO . ADOLT // FED. FEDIATRIC // INEO . NEONATE				
MAIN MENU	ADMIT TYPE: ADT	CHANGE ADMIT INFO	ADMIT		
PREV	HEIGHT UNIT: CM	WEIGHT UNIT: KG	DEFAULT SETTING		
MAIN MENU	ADMIT TYPE: ADT	> ADT PED	ADMIT		
PREV MENU		NEO	DEFAULT SETTING		

CHANGE ADMIT INFO

Last and first name (11 letters for each), sex (male or female), date of birth, weight, height, and patient ID (11 characters)

	ADMIT	CHANGE	
MAIN	TVPF.	ADMIT	DISCHARGE
MENU	ADT	INFO	
PREV	HEIGHT	WEIGHT	DEFAULT
MENU	UNII:	UNIT:	SETTING
	CM	KG	

	CHANGE ADMIT INFORMATION			
RETURN	CONTENTS			
LAST NAME	JOHN			
FIRST NAME	WASHINGTON			
PATIENT ID	APC001			
SEX	MALE			
BIRTH DATE	27 – JAN - 1978			
AGE	31			
HEIGHT	177.0 CM			
WEIGHT	62.0KG			
l				

DISCHARGE (Discharge Patient)

Patient information and all numbers change to standard, and the screen displays, "ALL ALARMS OFF ADMIT PATIENT TO ACTIVE ALARMS."

MAIN MENU	ADMIT TYPE: ADT	CHANGE ADMIT INFO	DISCHARGE
PREV MENU	HEIGHT UNIT: CM	WEIGHT UNIT: KG	DEFAULT SETTING
MAIN MENU	ADMIT TYPE: ADT	DISCHARGE	> NO
PREV MENU	HEIGHT UNIT: CM		YES

ADMIT(Admit patient)Depending on how your monitor is set up, you will see either ADMIT patient or new case.

MAIN MENU	ADMIT TYPE: ADT	CHANGE ADMIT INFO	ADMIT
PREV MENU	HEIGHT UNIT: CM	WEIGHT UNIT: KG	DEFAULT SETTING
MAIN MENU	ADMIT TYPE: ADT	ADMIT	> NO
PREV MENU	HEIGHT UNIT: CM		YES

HEIGHT

Unit of height is set as Cm / Inches.

MAIN MENU	ADMIT TYPE : ADT	CHANGE ADMIT INFO	ADMIT
PREV MENU	HEIGHT UNIT: CM	WEIGHT UNIT: KG	DEFAULT SETTING
MAIN MENU	ADMIT TYPE : ADT	CHANGE ADMIT INFO	ADMIT
PREV MENU	HEIGHT UNIT: INCH	WEIGHT UNIT: KG	DEFAULT SETTING

WEIGHT

Unit of weight is set as Kg / LBS.

MAIN MENU	ADMIT TYPE : ADT	CHANGE ADMIT INFO	ADMIT		
PREV MENU	HEIGHT UNIT: INCH	WEIGHT UNIT: KG	DEFAULT SETTING		
MAIN MENU	ADMIT TYPE : ADT	CHANGE ADMIT INFO	ADMIT		
PREV MENU	HEIGHT UNIT: INCH	WEIGHT UNIT: LBS	DEFAULT SETTING		

DEFAULT SETTING

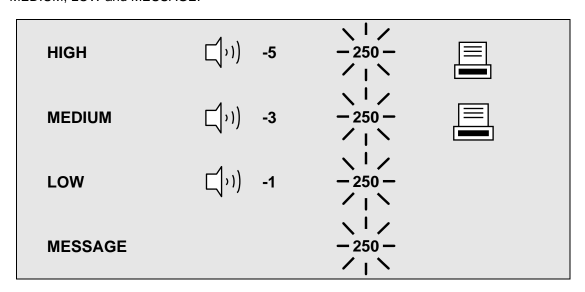
Resets the Alarm Limit settings to factory defaults as in "12.DEFAULT SETTING VALUE" section.

MAIN MENU	ADMIT TYPE: ADT	CHANGE ADMIT INFO	DISCHARGE
PREV MENU	HEIGHT UNIT: CM	WEIGHT UNIT: KG	DEAULT SETTING

2.2 ALARM



Alarm is divided into two, alarm for the patient's condition and for the product's condition. The patient's alarm sounds when the diagnostic functions (ASYSTOLE, VTAC/VFIB, and VTAC) are detected. Each alarm sound differs in order in order and volume according to the levels of HIGH, MEDIUM, LOW and MESSAGE.



: Alarm sounds

-250 - : Number flashes

: Waves are printed out

ALARM LIMITS: The machine enables one to see and change the limits of alarm for all parameter functions.

ALARM PRINT: with an ON/OFF setup, the related information is printed out whenever an alarm is given.

ALARM VOLUME: volume of each alarm can be adjusted in 10 step. ALARM LEVEL: Priority of each parameter alarm can be set up.

ALARM REVIEW: Shows the priority order information for all alarms of each measurement.

NURSE CALL: Set the feature of the NURSE CALL.

MAIN MENU	ALL LIMITS	ALARM PRINT: ON	ALARM VOLUME: OFF
PREV	NURSE	ALARM	ALARM
MENU	CALL	LEVEL	REVIEW

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ALL LIMITS

It is able to see all the alarm range and change of measurement function.

MAIN MENU	ALL LIMITS	ALARM PRINT: ON	ALARM VOLUME: OFF
PREV	NURSE	ALARM	ALARM
MENU	CALL	LEVEL	REVIEW

ALL LIMITS			
RETURN	UNITS	LOW	HIGH
HR	BPM	50	150
SPO2-%	%	90	100
SPO2-R	ВРМ	50	150
NIBP-S	mmHg	80	200
NIBP-M	mmHg	60	140
NIBP-D	mmHg	20	120
TEMP	°C	30.0	42.0
RESP	RPM	10	30
EtCO2	mmHg	25	50
FiCO2	mmHg	0	5
AWRR	RPM	10	30

ALARM PRINT

Set ON/OFF functions automatically. When the alarm is activated the corresponding information is printed on heat sensitive paper. Alarm level upper than MEDIUM Level. But, LEAD FAULT AND LOW BATTERY Alarm isn't activated the alarm print when alarm is set.

MAIN MENU	ALL LIMITS	ALARM PRINT: ON	ALARM VOLUME: OFF
PREV	NURSE	ALARM	ALARM
	CALL	LEVEL	REVIEW

ALARM VOLUME

Set the alarm volume to be set at 10 grades.

MAIN MENU	ALL LIMITS	ALARM PRINT: OFF	ALARM VOLUME: OFF
PREV MENU	NURSE CALL	ALARM LEVEL	ALARM REVIEW
MAIN MENU PREV MENU	ALARM VOLUME: OFF	> OFF 10% 20% 30% 40% 50%	60% 70% 80% 90% 100%

ALARM LEVEL

Set the order of priority in each alarm.

MAIN ALL LIMITS	ALARM PRINT: ON	ALARM VOLUME: OFF
PREV NURSE CALL	ALARM LEVEL	ALARM REVIEW

PARAMETER LEVEL

MAIN MENU	PARAMETER LEVEL	
PREV		
MENU		

PARAMET	TER ALARM LEVELS
RETURN	ALARM LEVEL
HR	MEDIUM
SPO2-%	MEDIUM
SPO2-R	LOW
RESP	MESSAGE
NIBP	MEDIUM
TEMP	MESSAGE
EtCO2	MEDIUM
FiCO2	MEDIUM
AWRR	LOW
LOW BATTERY	MEDIUM

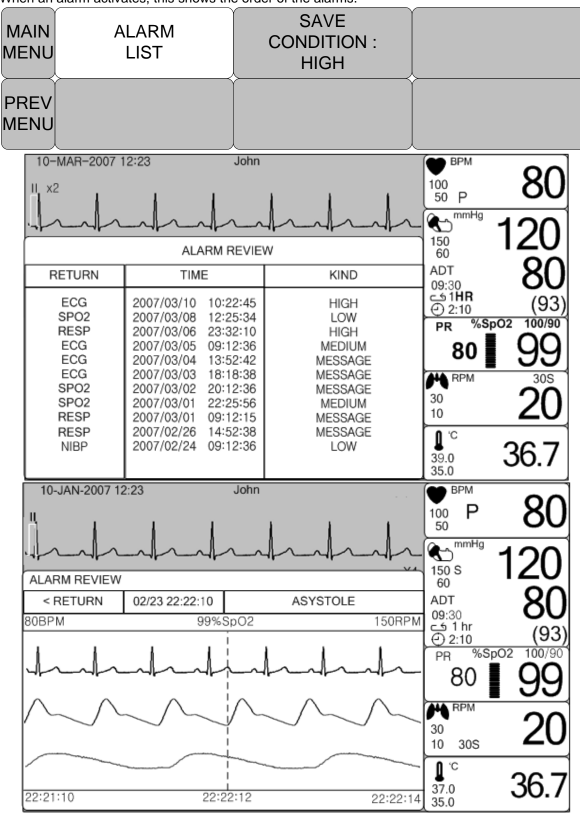
ALARM REVIEW

After an alarm is triggered the alarms and data wave pattern can be reviewed. Set up for priority of each parameter alarm.

<u> </u>					
MAIN MENU	ALL LIMITS	ALARM PRINT: ON	ALARM VOLUME: OFF		
PREV MENU	NURSE CALL	ALARM LEVEL	ALARM REVIEW		
MAIN MENU	ALARM LIST	SAVE CONDITION : HIGH			
PREV MENU					

ALARM LIST

When an alarm activates, this shows the order of the alarms.



SAVE CONDITION

This determines the order in which triggered alarms are saved.

MAIN MENU	ALARM LIST	SAVE CONDITION : HIGH	
PREV MENU			
MAIN MENU	ALARM LIST	SAVE CONDITION : HIGH	MESSAGE LOW
PREV MENU			MEDIUM > HIGH

NURSE CALL

When an alarm is triggered, this activated the NURSE CALL function.

MAIN MENU	ALL LIMITS	ALARM PRINT: ON	ALARM VOLUME: OFF
PREV	NURSE	ALARM	ALARM
MENU	CALL	LEVEL	REVIEW

NURSE CALL SETUP		
RETURN	CONTENTS	
NURSE CALL	OFF	
NORMAL MODE	NORMAL OPEN	
CALL MODE	ONE TIME	

- 1. NURSE CALL: ON/OFF
 - The nurse call function is enable or disable.
- 2. NORMAL MODE
 - NORMAL OPEN: Select this option when the hospital's call system is set to NORMAL OPEN.
 - NORMAL CLOSE: Select this option when the hospital's call system is set to NORMAL CLOSE.
- 3. CALL MODE
 - ONE TIME: When ONE TIME is selected, a nurse call signal is a pulse signal lasting 3s. When multiple alarms occur simultaneously, only one pulse signal will be output..
 - CYCLING: When CYCLING is selected, the duration of a nurse call signal is the same with the alarm, namely, from the time that the alarm occurs to the time it disappears. On and off repeatedly at intervals of 1 second.
 - CONTINUE: When CONTINUE is selected, the duration of a nurse call signal is the same with the alarm, namely, from the time that the alarm occurs to the time it disappears. However, lasts only one minute, then stops.

3. SETUP

3.1 SETUP

DISPLAY **SET PARA WAVE SELECT** SET DATE & TIME HR SOURCE **SWEEP SPEED KEY SOUND** DEMO **USER SERVICE** SET UNIT NAME SET BED NUMBER **AC FILTER SYSTEM** W-LAN **DISPLAY MODE** MAKER SERVICE FREEZING AND UNFREEZING

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3.1 SETUP



DISPLAY: screen set menu

USER SERVICE: This is the menu to set the connection used to interface with an external

computer

MAKER SERVICE: This is the basic adjustment menu used to adjust the features of this product.

MAIN MENU	DISPLAY		USER SERVICE
PREV MENU	KEY SOUND: ON	DEMO: ON	MAKER SERVICE

DISPLAY

SET PARA: Measurement function selected.

WAVE SELECT: Set wave pattern source at the bottom of the WINDOW with LARGE

SET DATE & TIME: Set and change date and time.

HR SOURCE : Set and select ECG(HR) / SpO2(PR) source. SWEEP SPEED : Set speed of ECG, SpO2 WAVE DISPLAY

MAIN MENU	SET PARA	WAVE SELECT: ECG	SET DATE & TIME
PREV MENU	SWEEP SPEED: 25mm/s		HR SOURCE: ECG

SET PARA

Select measurement function to use

MAIN MENU	SET PARA	WAVE SELECT: ECG	SET DATE & TIME
PREV MENU	SWEEP SPEED: 25mm/s		HR SOURCE: ECG

PARAMETER WINDOW SET		
RETURN	WINDOW ON/OFF	
ECG	ON	
SPO2	ON	
RESP	OFF	
NIBP	OFF	
TEMP	ON	
EtCO2	ON	

WAVE SELECT

Select waveform to display in large parameter display.

MAIN MENU	SET PARA	WAVE SELECT: ECG	SET DATE & TIME
PREV MENU	SWEEP SPEED: 25mm/s		HR SOURCE: ECG
MAIN MENU	SET PARA	WAVE SELECT: ECG	> ECG SPO2
PREV MENU	SWEEP SPEED: 25mm/s		RESP

SET DATE & TIME

It has sub menu to set date and time.

MAIN MENU	SET PARA	WAVE SELECT: ECG	SET DATE & TIME
PREV MENU	SWEEP SPEED: 25mm/s		HR SOURCE: ECG

SET TIME

Set time of equipment.

MAIN MENU	SET TIME	SET DATE	
PREV MENU			
MAIN MENU PREV MENU	SET TIME:	10:58:01	

SET DATE

Set date of equipment

MAIN SET SET DATE PREV MENU	20. 30.0	oquipmont		
MAIN SET DATE: O6-MAR-2007 PREV MENU	MENU		06-MAR-2007	

HR SOURCE

This menu is used to set the source that detects heart and pulse rate.

The source can select among ECG and SPO2.

	oc dan scicot among 200 an		
MAIN MENU	SET PARA	WAVE SELECT: ECG	SET DATE & TIME
PREV MENU	SPEED.		HR SOURCE: ECG
MAIN MENU	SET PARA	HR SOURCE: ECG	> ECG SPO2
PREV MENU	SPEED		3FU2

SWEEP SPEED

Set speed of drawing wave signal pattern in this widow.

MAIN MENU	SET PARA	WAVE SELECT: ECG	SET DATE & TIME
PREV MENU	SWEEP SPEED: 25mm/s		HR SOURCE: ECG
MAIN MENU	SWEEP SPEED: 25mm/s	> 6.25 mm/s 12.5 mm/s	SET DATE & TIME
PREV MENU		25 mm/s 50 mm/s	HR SOURCE: ECG

KEY SOUND

Set ON/OFF Key Sound of equipment.

MAIN MENU	DISPLAY		USER SERVICE
PREV MENU	KEY SOUND: OFF	DEMO: OFF	MAKER SERVICE

DEMO

Set ON/OFF DEMONSTRATION of equipment.

MAIN MENU	DISPLAY		USER SERVICE
PREV MENU	KEY SOUND: ON	DEMO: ON	MAKER SERVICE

USER SERVICE

The user is able to set the set UNIT NAME, BED NUMBER, external Wireless equipment power, communication parameters, display mode, and power supply filter.

MAIN MENU	DISPLAY		USER SERVICE
PREV MENU	KEY SOUND: ON	DEMO : ON	MAKER SERVICE
MAIN MENU	SET UNIT NAME	SET BED NUMBER : 00A	DISPLAY MODE: MONITOR
PREV MENU	SYSTEM	AC FILTER: 50HZ	W-LAN: OFF

SET UNIT NAME

Set up for UNIT(CCU,ICU,ER,etc.) name.

MAIN MENU	SET UNIT NAME	SET BED NUMBER : 00A	DISPLAY MODE: MONITOR
PREV MENU	SYSTEM	AC FILTER: 50HZ	W-LAN: OFF
MAIN MENU PREV MENU	SET UNIT NAME	ICU T	

SET BED NUMBER

Set up for patient bed number.

Allowable setters are from $0 \sim 9$, $A \sim Z$.

MAIN MENU	SET UNIT NAME	SET BED NUMBER : 00A	DISPLAY MODE: MONITOR
PREV MENU	SYSTEM	AC FILTER: 50HZ	W-LAN: OFF
MAIN MENU	SET UNIT NAME	SET BED NUMBER : 00A	0 0 A
PREV MENU	SYSTEM		0 0 A

AC FILTER

AC FILTER is function where you can set power supply frequency. This feature is required because power supply frequency can be different from one country to another. . (The selectable frequencies are 50Hz and 60Hz.)

MAIN MENU	SET UNIT NAME	SET BED NUMBER : 00A	DISPLAY MODE: MONITOR
PREV MENU	SYSTEM	AC FILTER: 50HZ	W-LAN: OFF
MAIN MENU	SET UNIT NAME	SET BED NUMBER : 00A	DISPLAY MODE: MONITOR
PREV MENU	SYSTEM	AC FILTER: 60HZ	W-LAN: OFF

SYSTEM

System able to change and verify Equipment version information and system information

SYSTEM INFO SET				
RETURN	CONTENTS			
MAIN VER	1.10.BHCDDCA			
CENTRAL	ON			
HOST IP	192 . 168 . 030 . 077			
DEVICE IP	192 . 168 . 030 . 100			
SUBNET	255 . 255 . 255 . 000			
GATEWAY	192 . 168 . 030 . 001			
MAC ADDR 00 : 02 : A8 : 80 : CB : 00				
VGA OUTPUT	OFF			
DHCP	OFF			
HL7	OFF			
HL7 SERVER IP	192 . 168 . 030 . 200			
HL7 SERVER PORT	T 04200			
EXPORT INTERVAL 5 Min				
NAK	OFF			

VGA OUTPUT: VGA output on the output board provides.

CENTRAL: ON / OFF function of the network system used to set.

HL7: ON/OFF function of HL7 network protocol.

Will turn ON after setting the equipment off and connected to the Central system or HL7 system

NAK: ON/OFF function of transmission control of HL7 protocol

DHCP: ON/OFF function of allocation IP address automatically.

HOST IP, DEVICE IP, SUBNET and GATEWAY: Set the information for connecting to the Central system.

Warning

We recommend to use a static IP outside the DHCP range

W-LAN

W-LAN power can be supplied for enabling a External wireless LAN equipment use.

MAIN MENU	SET UNIT NAME	SET BED NUMBER : 00A	DISPLAY MODE: MONITOR
PREV MENU	SYSTEM	AC FILTER: 60HZ	W-LAN: OFF

DISPLAY MODE (MONITOR or SPOT)

You can selected Monitoring display mode or Spot display mode.

MAIN MENU	SET UNIT NAME	SET BED NUMBER : 00A	DISPLAY MODE: MONITOR
PREV	SYSTEM	AC FILTER: 60HZ	W-LAN: OFF

MAKER SERVICE

Maker service is a menu is used by manufacturers.

MAIN MENU	DISPLAY		USER SERVICE
PREV MENU	KEY SOUND: ON	DEMO : ON	MAKER SERVICE

Freezing and Unfreezing



This icon is pressed to freeze waveforms. All displayed waveforms are frozen.
 The waveforms are frozen for 1 minutes or until they are unfrozen.

Press the "F" key is unfrozen with large parameter mode.

When the waveforms are frozen, the "FREEZE" message appears with the frozen time.



 Press the FREEZE icon (Unfreezing icon) on the control panel again. After exiting the frozen screen, new real time waveforms are displayed.

4. TREND

4.1 TREND

GRAPHIC TREND TABULAR TREND TREND WINDOW SETUP

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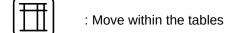
4.1 TREND

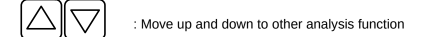


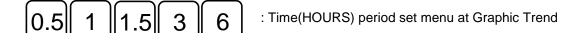
TREND shows saved data graphically displayed with numeric values.

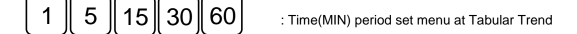
Real-time data recording duration is 1 minute. Amount of saving time is for this data will be saving for 128hours.

MAIN MENU	GRAPHIC TREND	TABULAR TREND	TREND WINDOW SETUP
PREV MENU			



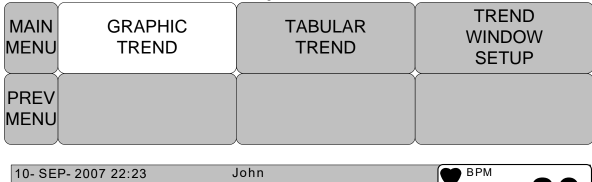


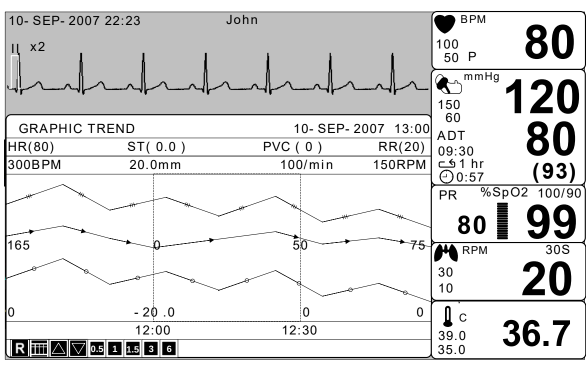




GRAPHIC TREND

Wave Data can be stored and seen according to section.





TIME PERIOD

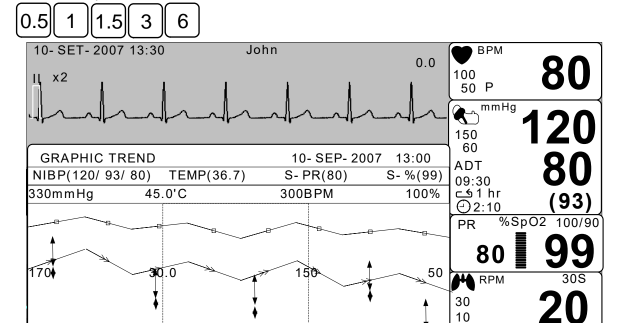
10

15.0

12:00

R III 🛆 💟 0.5 1 1.5 3 6

One can set up and store data and time that one can see in a screen.



12:30

1 c

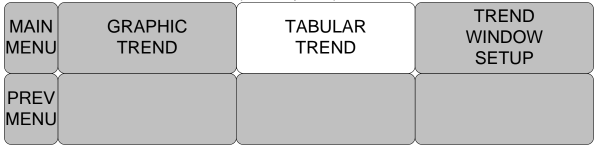
39.0

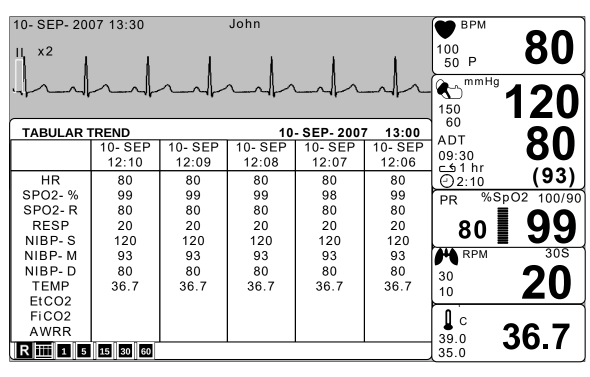
35.0

36.7

TABULAR TREND

One can see the stored data at the time previously set up.

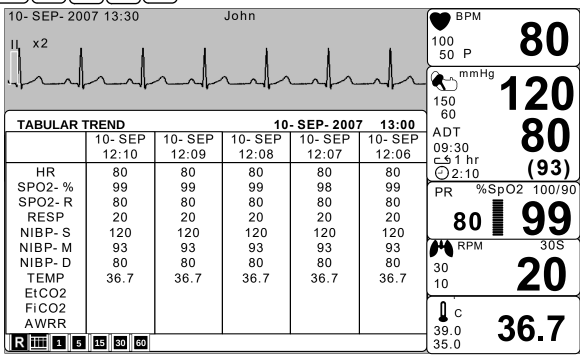




TIME INTERVAL

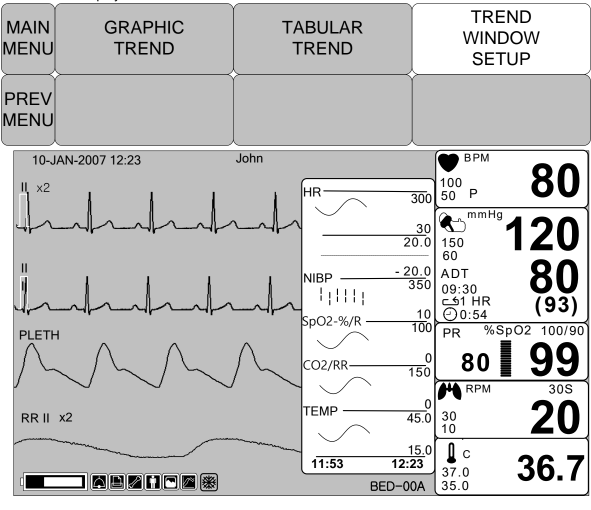
One can store data and set up time.





TREND WINDOW SETUP

Set the trend display window that will show the real time wave window.



TIME PERIOD

Set visible time period in a screen.

MAIN MENU	TIME PERIOD: 30MINS	SET TREND PARA	
PREV MENU			
MAIN MENU	TIME PERIOD: 30MINS	> 30MINS 60MINS 90MINS	
PREV MENU		3HOUR 6HOUR 12HOUR	

SET TREND

Set parameter for display in a screen.

MAIN MENU	TIME PERIOD: 30MINS	SET TREND	
PREV MENU			

PARAMETER WINDOW SET		
RETURN	ON / OFF	
HR	ON	
ST	ON	
SPO2	ON	
PR	ON	
RESP	ON	
NIBP	ON	
TEMP	ON	

TREND PRINT

Graphic: select the number which selects a graphic trend and press print to prints the selected trend. Table: select the table number to be print and press print to receive print all the data in the selected patient admit (Admit) table.

5. ECG

5.1 Outline

Color and Standards of Cables
Position of ECG Connector and Measurement Cable
Attaching Electrodes to the Patient
Choosing an ECG lead for Arrhythmia Monitoring
Information on the ECG waveform
5-Lead Electrode Placement
3-Lead Electrode Placement
Electrode Placement for Neonates

5.2 ECG Data Window

5.3 ECG Setup

LEAD SELECT
ALARM LIMIT
ALARM LEVEL
ALARM SOUND
QRS VOLUME
DISPLAY
ECG SWEEP SPEED
ECG SIZE
HR SOURCE
ANALYSIS SETTING

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5.1 Outline

It calculates the heart rate with 3 or 5 leads ECG signal acquisition and perform the alarm according to the setting value.

Colors and Standards of Cables

Leadwire	AHA Color code	AHA Label	IEC Color code	IEC Label
Right arm	White	RA	Red	R
Left arm	Black	LA	Yellow	L
Right leg	Green	RL	Black	N
Left leg	Red	LL	Green	F
V1(precordial)	Brown	V1	White	C1

AHA: American Heart Association (U.S.A. standard)

IEC : International Electro technical Commission (Europe standard)

Position of ECG Connector and Measuring Cable ECG connecter +detect cable



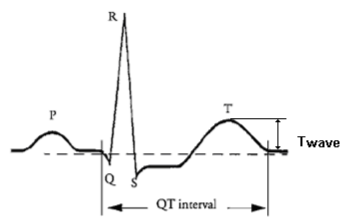
Attaching Electrodes to the Patient

- 1. Shave excess hair. With a piece of cotton pad moistened with alcohol, clean the patient's skin where the electrodes should be mounted. Avoid wrinkled or uneven skin areas. Wipe off the alcohol with a dry cotton pad.
- 2. Open the electrode package and take out the electrode.
- 3. Remove the backing paper from the electrode. Be careful not to touch the adhesive side.
- 4. Attach the disposable electrode to the previously cleaned skin. Avoid wrinkled and uneven skin areas.
- 5. The electrode lead which is connected to the monitor onto the electrode.
- 6. Fasten the electrode lead to the skin with surgical tape with an extra length of wire between the tape and the electrode. This prevents body movement from moving the electrode lead.

Note

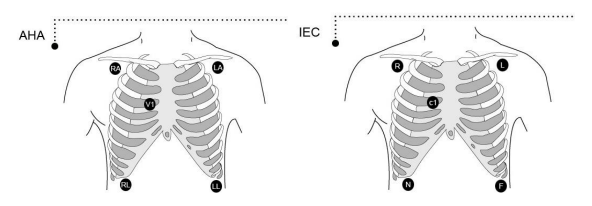
- ✓ To maintain good contact between the electrode and skin, check that the paste of the disposable electrode is not dry.
- When contact of the disposable electrode becomes poor, replace the electrode with a new one immediately. Otherwise, contact impedance between the skin and electrode increase and the correct ECG cannot be obtained.
- ✓ If the contact is bed before the expiration date on the package, replace the electrode with a new one.
- ✓ To obtain a stable ECG waveform rub the skin with "skin Pure" skin preparation gel or tincture of Benzion.
- ✓ Shall use only the CE certified disposable electrode.

Information on the ECG waveform

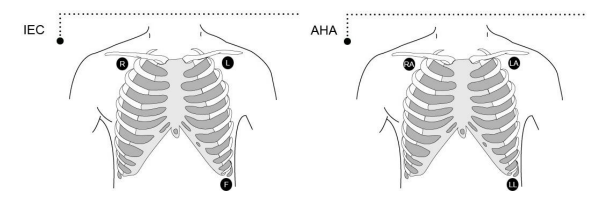


When ECG signal is 80bpm T-wave duration is 180ms, and the QT interval is 350ms.

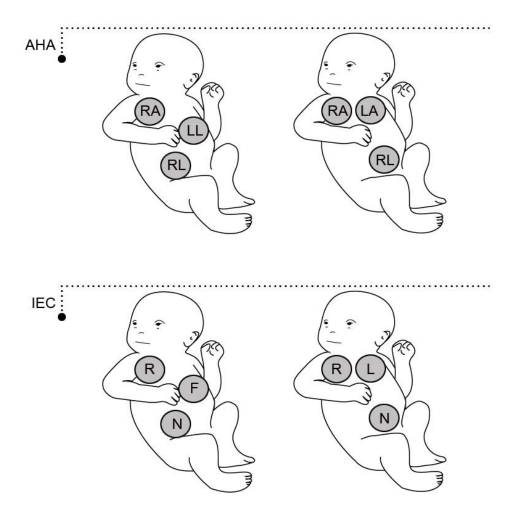
5-Leadwire Electrode Placement



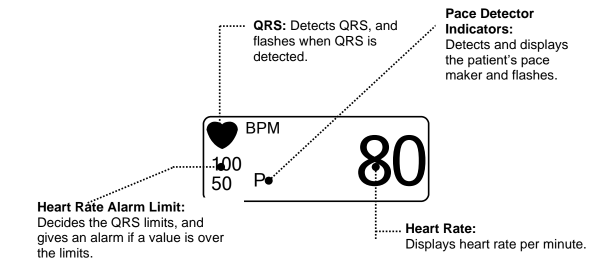
3-Leadwire Electrode Placement



Electrode Placement for Neonates



5.2 ECG Data Window



Note

ECG Wave Display is always on when the cable is connected.

The heart rate is calculated by a moving average. The monitor detects 8 consecutive beats, averages the R-R intervals of the latest 8 beats and uses this average to calculate the current heart rate. When a new beat is detected, the heart rate is recalculated using the latest 8beats. The heart rate display is updated every 3 seconds.

Heart rate meter updates a new heart rate for a step increase or decrease in 10 seconds maximum. When ventricular tachycardia is detected, the alarm set in 5 seconds maximum.

Check that the delay time of the output signal (alarm trigger 80ms maximum) is within the range of the connected equipment.

Safety Precautions

Warning

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

CONDUCTIVE CONNECTIONS — Extreme care must be exercised when applying medical electrical equipment. Many parts of the human/machine circuit are conductive, such as the patient, connectors, electrodes, transducers. It is very important that these conductive parts do not come into contact with other grounded, conductive parts when connected to the isolated patient input of the device. Such contact would bridge the patient's isolation and cancel the protection provided by the isolated input. In particular, there must be no contact of the neutral electrode and ground.

DEFIBRILLATION — Do not come into contact with patients during defibrillation. Otherwise serious injury or death could result.

To avoid the risk of serious electrical burn, shock, or other injury during defibrillation, all persons must keep clear of the bed and must not touch the patient or any equipment connected to the patient.

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

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

The peak of the synchronized defibrillator discharge should be delivered within 60ms of the peak of the R wave. The signal at the ECG output on the patient monitors is delayed by a maximum of 30ms.

If the ECG waveform on the screen is too unstable to synchronize with the patient's heart beat because of the following reason, remove the cause of an alarm, message, or unstable ECG, and then use a stable ECG lead for synchronization.

- ✓ ECG electrode is detached or broken. Lead wire is detached or broken.
- ✓ Lead wire moves. AC interference, EMG noise or noise from ESU is superimposed.
- ✓ Connection cable is broken or has a short circuit. Connector has poor contact.

INTERFACING OTHER EQUIPMENT — Devices may only be interconnected with each other or to parts of the system when it has been determined by qualified biomedical engineering personnel that there is no danger to the patient, the operator, or the environment as a result. In those instances where there is any element of doubt concerning the safety of connected devices, the user must contact the manufacturers concerned (or other informed experts) for proper use. In all cases, safe and proper operation should be verified with the applicable Manufacturer's instructions for use, and system standards IEC 60601-1-1/EN 60601-1-1 must be complied with.

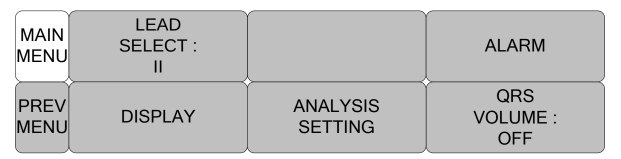
Electrosurgery Unit

- ✓ Electrosurgical units(ESU) emit a lot of RF interference. If the monitor is used with an ESU,RF interference may affect the monitor operation.
- ✓ Locate the monitor as far as possible from the ESU. Locate them on opssite sides of the operating table, if possible.
- ✓ Connect the monitor and ESU to different AC outlets located as far as possible from each other
- ✓ When using this monitor with an electrosurgical unit, its return plate and the electrodes for monitoring must be firmly attached to the patient. If the return plate is not attached correctly,it may burn the patient's skin where the electrodes are attached.

5.3 ECG Setup

A setup window appears at lower part of the screen when the Trim Knob Key is pressed in the ECG Parameter Window.

Selection is made by pressing the Trim Knob Key, while movement across the menu is performed by turning the key either clock or anticlockwise.



LEAD SELECT

Select channels from I to V in ECG

MAIN MENU	LEAD SELECT : II		ALARM
PREV MENU	DISPLAY	ANALYSIS SETTING	QRS VOLUME : OFF
MAIN MENU PREV MENU	LEAD SELECT : II	> 	aVR aVL aVF V

ALARM LIMIT

Alarm Limit is 0 ~ 300.

	11.10 0 000.		
MAIN MENU	LEAD SELECT : II		ALARM
PREV MENU	DISPLAY	ANALYSIS SETTING	QRS VOLUME : OFF
MAIN MENU	ALARM LIMIT	ALARM SOUND	
PREV MENU	ALARM LEVEL		

	ECG ALARM LIMIT			
RETURN	UNITS	LOW	HIGH	
HR	BPM	60	120	

ALARM LEVEL

Set the order of priority in each alarm.

MAIN	ALARM	ALARM	
MENU	LIMIT	SOUND	
PREV MENU	ALARM LEVEL		

ALARM LEVELS			
RETURN	ALARM LEVEL		
HR LEAD FAULT	MEDIUM MESSAGE		

ALARM SOUND

Set ON/OFF of ECG alarm sound.

001014/0	Set ON/OFF of ECG alaim sound.			
MAIN MENU	LEAD SELECT : II		ALARM	
PREV MENU	DISPLAY	ANALYSIS SETTING	QRS VOLUME : OFF	
MAIN MENU	ALARM LIMIT	ALARM SOUND		
PREV MENU				
ECG ALARM SOUND				
1				

ECG ALARM SOUND		
RETURN ECG ALARM SOUND		
HR	ON	

QRS VOLUME

Move the Key to select a volume rate from OFF, 10% to 100%.

MAIN MENU	LEAD SELECT : II		ALARM
PREV MENU	DISPLAY	ANALYSIS SETTING	QRS VOLUME : OFF
MAIN MENU PREV MENU	QRS VOLUME : OFF	> OFF 10% 20% 30% 40% 50%	60% 70% 80% 90% 100%

DISPLAY

Set the sweep speed and waveform size.

MAIN MENU	LEAD SELECT : II		ALARM
PREV MENU	DISPLAY	ANALYSIS SETTING	QRS VOLUME : OFF

ECG SWEEP SPEED

ECG speed is 25 mm/s.

Speed is changeable to 6.25, 12.5, 25, 50mm/s.

MAIN MENU	SWEEP SPEED : 25 mm/s	ECG SIZE : X1	HR SOURCE: ECG
PREV MENU			
MAIN MENU	SWEEP SPEED : 25 mm/s	6.25 mm/s 12.5 mm/s	HR SOURCE: ECG
PREV MENU		> 25 mm/s 50 mm/s	

ECG SIZE

The size is changeable to X0.25, X0.5, X1, X2, X4.

MAIN MENU	SWEEP SPEED : 25 mm/s	ECG SIZE : X1	HR SOURCE: ECG
PREV MENU			
MAIN MENU	SWEEP SPEED : 25 mm/s	ECG SIZE : X1	x 0.25 x 0.5 > x 1
PREV MENU			x 2 x 4

HR SOURCE

MAIN MENU	SWEEP SPEED : 25 mm/s	ECG SIZE : X1	HR SOURCE: ECG
PREV MENU			
MAIN MENU	SWEEP SPEED : 25 mm/s	HR SOURCE: ECG	> ECG SPO2
PREV MENU			31 02

ANALYSIS SETTING

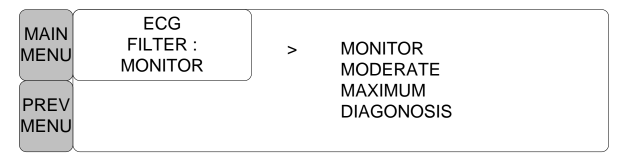
Analysis setting divided to 3 menus.

ECG FILTER: One may select from three frequency types for WAVE FILTER.

MONITÓR 0.5Hz ~ 40Hz MODERATE 0.5Hz ~ 25Hz MAXIMUM 5Hz ~ 25Hz

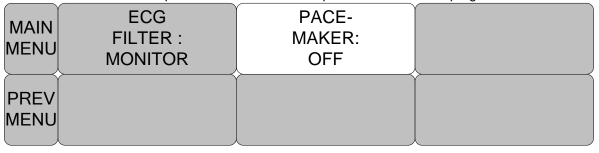
DIAGONOSIS 0.05Hz ~ 150Hz

MAIN MENU	LEAD SELECT : II		ALARM
PREV MENU	DISPLAY	ANALYSIS SETTING	QRS VOLUME : OFF
MAIN MENU	ECG FILTER : MONITOR	PACE - MAKER: OFF	
PREV MENU			



PACE MAKER: Sets up ON/OFF to indicate that the patient has PACE MAKER.

The PACE MAKER menu option enables/disables the pacemaker detection program.



Be aware of the following when monitoring a patient with a pacemaker.

Warning

FALSE CALLS—False low heart rate indicators or false asystole calls may result with certain pacemakers because of electrical overshoots.

MONITORING PACEMAKER PATIENTS—Monitoring of pacemaker patients can only occur with the pace program activated.

PACEMAKER SPIKE—An artificial pacemaker spike is displayed in place of the actual pacemaker spike. All pacemaker spikes appear uniform. Do not diagnostically interpret pacemaker spike size and shape.

PATIENT HAZARD—A pacemaker pulse can be counted as a QRS during asystole in either pace mode. Keep pacemaker patients under close observation.

PACEMAKER PATIENTS. Rate meters may continue to count the pacemaker rate during occurrences of cardiac arrest or some arrhythmias. Do not rely entirely upon rate

meter ALARMS. Keep pacemaker patients under close surveillance.

CAUTION

FDA POSTMARKET SAFETY ALERT

The United States FDA Center for Device and Radiological Health issued a safety bulletin October 14, 1998. this bulletin states "that minute ventilation rate-adaptive implantable pacemakers can occasionally interact with certain cardiac monitoring and diagnostic programmed rate."

The FDA further recommends precautions to take into consideration for patients with these types of pacemakers. These precaution for patients with these types of pacemakers. These precautions include disabling the rate responsive mode and enabling an alternate pace mode. For more information contact:

Office of Surveillance and Biometrics, CDRH, FDA 1350 Packard Drive, Mail Stop HFZ-510 Rockville, MD 20850 U.S.A

NOTE

ECG monitoring with patients in non-invasive trans coetaneous pacemakers may not be possible due to large amounts of energy produced by these devices. Monitoring ECG with an external device may be needed.

WARNINGS

VENTRICULAR ARRHYTHMISAS

The arrhythmia analysis program is intended to detect ventricular arrhythmias. It is not designed to detect a trial or supra ventricular arrhythmias. Occasionally it may incorrect identify the presence or absence of an arrhythmia. Therefore, a physician must analyze the arrhythmia information in conjunction with other clinical findings.

SUSPENDED ANALYSIS

Certain conditions suspend arrhythmia analysis. When suspended, arrhythmia conditions are not detected and alarms associated with arrhythmias do not occur. The messages which alert you to the conditions causing suspended arrhythmia analysis are: ARR OFF, ARRHYSUSPEND, LEADS FAIL, ALARM PAUSE, ALL ALARMS OFF, and DISCHARGED.

Trouble shooting

Problem:

Inaccurate heart rate and/or false a systole.

Solution:

Check ECG signal from patient:

- 1. Check/adjust lead placement.
- 2. Check/perform skin preparation.
- 3. Check/replace electrodes.

Check amplitude of ECG waveform:

- 1. Select ECG parameter label.
- 2. Select DISPLAY LEAD,
- 3. Scroll through all ECG leads and check for 0.5mV amplitude at normal (1X) size. (at least 0.5mV amplitude is required for QRS detection.) for borderline signals, validate on a graph.
- 4. If amplitudes are low, electrodes may need to be repositioned or replaced.

Problem:

False ventricular calls.

Solution:

Check ECG signal from patient: (the chest lead may exhibit polarity changes which may occasionally cause an inaccurate call.)

- 1. Check/adjust lead placement.
- 2. Check/perform skin preparation.
- 3. Check/replace electrodes. (if chest lead is a problem, move the chest lead to another chest position or leg position.)

Problem:

Inaccurate pacemaker detection

Solution:

Use pacemaker processing:

- 1. Select ECG parameter label.
- 2. Display the lead of ECG with the greatest amplitude in the top waveform position.
- 3. Select ANALYSIS SETTINGS.
- 4. SELECT DETECT PACE.

6. SpO₂

6.1 Outline

SpO₂ Connector Location and Measuring Cable

6.2 SpO2 Data Window

Signal and Data Validity Stability of SPO2 Values

6.3 SpO2 SetupRATE VOLUME ALARM ALARM LIMIT ALARM LEVEL **ALARM SOUND**

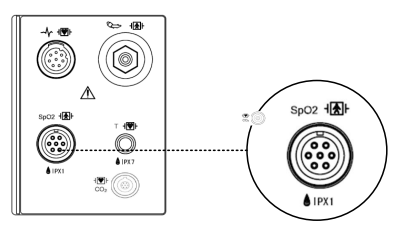
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6.1 Outline

SPO2 monitoring is a noninvasive technique used to measure the amount of oxygenated hemoglobin and pulse rate by measuring the absorption of selected wavelengths of light. The light generated in the probe passes through the tissue and is converted into an electrical signal by the photodetector in the probe. The monitor processes the electrical signal and displays on the screen a waveform and digital values for SpO2 and pulse rate. It detects SpO2 in the way of transmitting the red and infrared rays into the capillary vessel to take the pulsation. Also perform the alarm function according to the setting value.

SpO2 Connector Location and Measuring Cable

SpO₂ connector



SpO₂ Measuring Cable

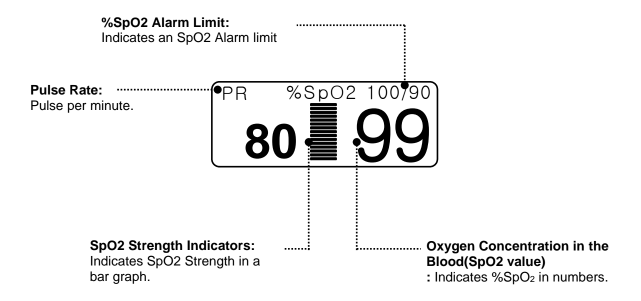


Note

The signal input is a high-insulation port and it is defibrillator proof (

The insulated input ensures patient safety and protects the device during defibrillation and electrosurgery.

6.2 SpO₂ Data Window



The current SPO2 value and the derived pulse rate (RATE) are displayed. The block sets indicate the strength of the signal (twenty block bars indicate the strongest signal). The SPO2 measurements are averaged over a 6-second period of time.

The monitor display is updated every second.

The SPO2 monitoring features are found in the SPO2 menu. These features include alarm limit adjustment, display of RATE, and RATE volume.

Note
SpO₂ WAVE SIZE is changed automatically.

Signal and Data Validity

It is extremely important to determine that the probe is attached to the patient correctly and the data is verifiable. To make this determination, three indications from the monitor are of assistance—signal strength bar, quality of the SPO2 waveform, and the stability of the SPO2 values. It is critical to observe all three indications simultaneously when ascertaining signal and data validity.

Signal Strength Bar

The signal strength bar is displayed within the SPO2 values window. This bar consists of 20 blocks set depending on the strength of the signal. Proper environmental conditions and probe attachment will help to ensure a strong signal.

Quality of SPO2 Waveform

Under normal conditions, the SPO2 waveform corresponds to (but is not proportional to) the arterial pressure waveform. The typical SPO2 waveform indicates not only a good waveform, but helps the user find a probe placement with the least noise spikes present. The figure below represents an SPO2 waveform of good quality.



If noise (artifact) is seen on the waveform because of poor probe placement, the photodetector may not be flush with the tissue. Check that the probe is secured and the tissue sample is not too thick. Pulse rate is determined from the SPO2 waveform which can be disrupted by a cough or other hemodynamic pressure disturbances. Motion at the probe site is indicated by noise spikes in the normal waveform. (See the figure below.) It has been noted that letting the patient view the SPO2 waveform enables them to assist in reducing motion artifact.



SPO2 Waveform with Artifact

Stability of SPO2 Values

The stability of the displayed SPO2 values can also be used as an indication of signal validity. Although stability is a relative term, with a small amount of practice one can get a good feeling for changes that are artifactual or physiological and the speed of each. Messages are provided in the SPO2 values window to aid you in successful SPO2 monitoring.

WARNING

In the monitoring of patients the coincidence of adverse conditions may lead to a disturbed signal going unnoticed. In this situation artifacts are capable of simulating a plausible parameter reading, so that the monitor fails to sound an alarm. In order to ensure reliable patient monitoring, the proper application of the probe and the signal quality must be checked at regular intervals.

6.3 SpO₂ Setup

ALARM : Menu in which SpO₂ alarm are set up.

RATE VOLUME: Menu in which RATE VOLUME is set up

MAIN	ALARM	RATE VOLUME: OFF

RATE VOLUME

Move the KEY to select the volume from OFF to 100%. When the ECG volume rate is set, it turns OFF automatically.

MAIN MENU	ALARM		RATE VOLUME: OFF
MAIN MENU	RATE VOLUME: OFF	> OFF 10% 20% 30%	60% 70% 80% 90%
		40% 50%	100%

ALARM

Two menus: ALARM LIMIT, LEVEL and ALARM SOUND provided in the alarm menu

MAIN MENU	ALARM	RATE VOLUME: OFF

ALARM LIMIT

Number setting of alarm value of %SpO2 is 0 ~ 100

- 1. Move the ightharpoonup mark to select from RETURN, SpO₂ or SpO₂-R, and press.
- 2. After pressing at SpO₂, move the cursor right or left to LOW, and press.
- 3. Once the color is changed, move the cursor again to the selected value and press.
- 4. Place the cursor to HIGH and press, when the color changes, move the cursor again to select the targeted value, and press. Finally move to SpO₂ and press.

(You may decide to perform the process in the opposite order, LOW to HIGH, to have the same result.)

- 5. After pressing at SpO₂-R, move the cursor right or left to LOW, and press.
- 6. Once the color is changed, move the cursor again to the selected value and press.
- 7. Place the cursor to HIGH and press, when the color changes, move the cursor again to select the targeted value, and press. Finally move to SpO₂-R and press.
- 8. With the selection of RETURN the user gets out of the menu.

MAIN MENU	ALARM LIMIT	ALARM SOUND: ON	
PREV	ALARM LEVEL		

SPO2 ALARM LIMIT					
RETURN	UNITS	LOW	HIGH		
SPO2-%	%	90	100		
SPO2-R	BPM	50	150		

ALARM LEVEL

Set the order of priority in each alarm.

MAIN	ALARM	ALARM	
MENU	LIMIT	SOUND	
PREV MENU	ALARM LEVEL		

ALARM LEVELS				
RETURN	ALARM LEVEL			
SPO2-%	LOW			
SPO2-R	MESSAGE			
PROBE OFF	MESSAGE			
CHECK PROBE	MESSAGE			
LOST PULSE	LOW			
POOR SIGNAL	LOW			
ARTIFACT	LOW			

ALARM SOUND

Warning sound or message displays configuration menu when an alarm is triggered.

MAIN MENU	ALARM LIMIT	ALARM SOUND: ON	
PREV MENU	ALARM LEVEL		
MAIN MENU	ALARM LIMIT	ALARM SOUND: OFF	
PREV MENU	ALARM LEVEL		

Probe Off Condition

When using a reusable finger probe, there is a system alarm to alert you when the probe is off the Monitor. The monitor defaults this "PROBE OFF" condition as a System Warning alarm. however, You can set it as a System ALARM LEVEL in Monitor Defaults.

SPO2 Messages

Below is a list of system status alarm messages which may be displayed in the SPO2 parameter window during monitoring.

CHECK PROBE

Reusable finger probe is off the patient. Check the probe. *The factory default for this alarm is MESSAGE ALARM.*

PULSE SEARCH

Detection by the monitor of a repeatable pulse has ceased. Check the patient and the probe site.

POOR SIGNAL

The SPO2 signal is too low. No SPO2 data is displayed. This can be due to a low patient pulse, patient motion, or some other interference. Check the patient and the probe.

LOST SIGNAL

SPO2 data continues to be displayed, but the quality of the signal is questionable. Check the patient and the probe.

ARTIFACT

The SPO2 signal is patient's motion artifact and noise

No SpO2 data is displayed. One of the following conditions is indicated:

- defective or damaged probe,
- defective or damaged cable
- probe is off the patient, or
- Detection of a repeatable pulse has ceased.
- Check the probe and cable: reposition or replace as needed.

WARNING

- Federal Laws of the United States restrict the use and sale of this device by, or on the order of a physician.
- Do not use this device in the presence of flammable anesthetics.
- Do not use this device in the presence of magnetic resonance imaging (MR or MRI) equipments.
- This device must be used in conjunction with clinical signs and symptoms. This device is only intended to be an adjunct in patient assessment.
- Operation of this device may be adversely affected in the presence of strong electromagnetic sources, such as electro-surgery equipments.
- Operation of this device may be adversely affected in the presence of computed tomography (CT) equipments.
- Misuse or improper handling of the device could result in damage to the device. This may cause inaccurate readings.

CAUTIONS

- Do not autoclave, ethylene oxide sterilize, or immerse the device in liquid.
- This device is intended for use by persons trained in professional health care who have a complete understanding of pulse Oximetry. The operator must be thoroughly familiar with the information in this manual before using the device.
- %SpO2 measurements may be adversely affected in the presence of high ambient light. Avoid exposure to high ambient light.
- Dyes introduced into the bloodstream, such as methylene blue, indocyanine green, indigo carmine, fluorescein, and patent blue V may adversely affect the accuracy of the %SpO2 reading.
- Any condition that restricts blood flow, such as use of blood pressure cuff, extremes in systemic vascular resistance, or reduction of peripheral circulation caused by hypothermia, may cause an inability to determine accurate pulse rate and %SpO2 readings.

7. RESPIRATION

7.1 Outline

Respiration Connector and Measuring Cable

7.2 RESPIRATION Data Window

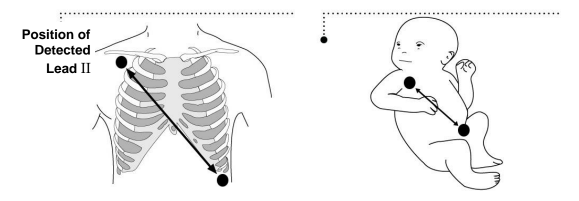
7.3 RESPIRATION Setup

Respiration SWEEP SPEED
Respiration SIZE
APNEA DETECT
ALARM
ALARM LIMIT
ALARM LEVEL
ALARM SOUND

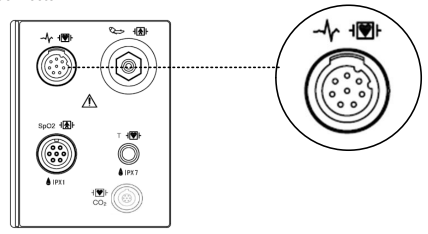
BM3OM-2.62

7.1 Outline

Respiration via ECG Lead II electrode makes the skin area of the chest enlarged, causing changes in the resistance of skin. Through this it calculates respiration value per minutes and performs the alarm function according to limit value.



Respiration Connector and Measuring Cable Respiration Connecter

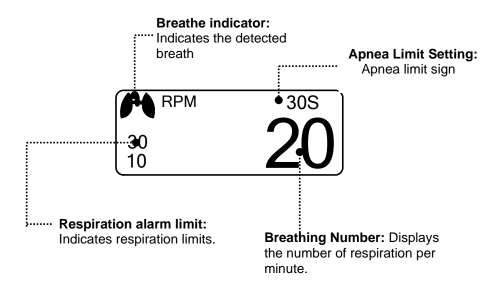


Respiration Measuring Cable



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7.2 Respiration Data Window



7.3 Respiration Setup

ALARM: Respiration alarm setting menu RESP SIZE: A menu to setup Wave Display

SWEEP SPEED: A menu to setup Wave Display of speed

MAIN MENU	ALARM	SWEEP SPEED : 25mm/s	RESP SIZE : X 2

RESPIRATION SPEED

Wave pattern speed is 6.25, 12.5, 25 mm/s.

MAIN MENU	ALARM	SWEEP SPEED: 12.5mm/s	RESP SIZE : X 2
MAIN MENU	ALARM	SWEEP SPEED: 12.5mm/s	6.25 mm/s > 12.5 mm/s
			25 mm/s

RESPIRATION SIZE

Set wave pattern size X2~ X10.

MAIN MENU	ALARM	SWEEP SPEED: 12.5mm/s	RESP SIZE: X2
MAIN MENU	ALARM	RESP SIZE: X 2	> X 2 X 4 X 6
			X 8 X10

ALARM

Alarm menu provide ALARM LIMIT, LEVEL and ALARM SOUND.

MAIN MENU	ALARM	SWEEP SPEED: 12.5mm/s	RESP SIZE : X 2
	APNEA DETECT : ON		

ALARM LIMIT

Alarm Limit of Respiration Numeric Value is 5 ~ 150bpm

MAIN MENU	ALARM LIMIT	ALARM SOUND : ON	
PREV	ALARM LEVEL		

- 1. Move the mark to select RETURN, RESP and press.
- 2. After a press in RESP, move the cursor right or left to LOW, and press.
- 3. After the color changed, move the cursor right or left to the selected value, and press.
- 4. Place the cursor to HIGH, and press. When the color has changed, move the cursor again to select the value and press. Move to the RESP and press again. (You may decide to perform the process in the opposite order, LOW to HIGH, to have the same result.)
- 5. Select RETURN to get out of the window.

	RESP ALARM LIMIT				
RETURN	UNITS	LOW	HIGH		
RESP	RPM	10	30		

ALARM LEVEL

Set the order of priority in each alarm.

MAIN	ALARM	ALARM	
MENU	LIMIT	SOUND	
PREV MENU	ALARM LEVEL		

ALARM LEVELS			
RETURN	ALARM LEVEL		
RESP	MESSAGE		

ALARM SOUND

Warning sound or message displays activation setting when Respiration ALRAM occurs.

MAIN MENU	ALARM LIMIT	ALARM SOUND : ON	
PREV MENU	ALARM LEVEL		
MAIN MENU	ALARM LIMIT	ALARM SOUND : OFF	
PREV MENU	ALARM LEVEL		

8. NIBP

8.1 Outline

Position of NIBP Connector and Cuff

8.2 NIBP Data Window

8.3 NIBP Setup

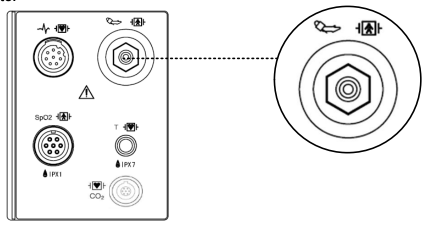
ALARM
ALARM LIMIT
ALARM SOUND
NIBP STAT
CUFF SIZE
UNIT SELECT
INTERVAL
INFLATION SET

BM3OM-2.62

8.1 Outline

This function is to measure minimum, Maximum and average blood pressure by using Oscillometric method

Position of NIBP Connecter and cuff NIBP Connector



ADULT CUFF



Note

As the value of NIBP can vary according to the age and sex of a patient, the user needs to set up right data in Parameter Menu before measurement.

WARNING

Noninvasive blood pressure monitoring is not recommended for patients with hypotension, hypertension, arrhythmias or extremely high or low heart rate. The software algorithm cannot accurately compute NIBP or patients with these conditions.

Note

As the value of NIBP can vary according to the age and sex of a patient, the user needs to set up right data in parameter Menu before measurement. Tubes between the cuff and the monitor are not kinked or blocked.

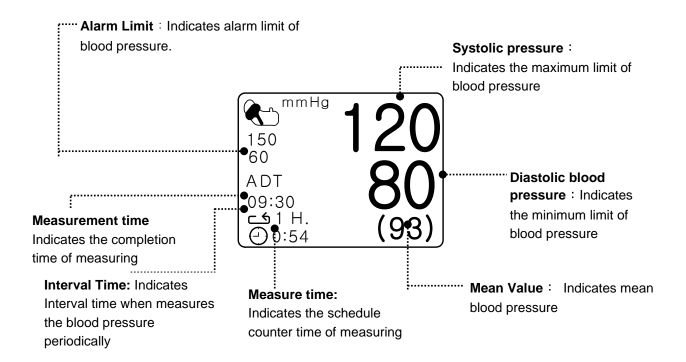
The air pad should be exactly over the branchial artery. Tubing is immediately to the right or left of the branchial artery to prevent kinking when elbow is bent.

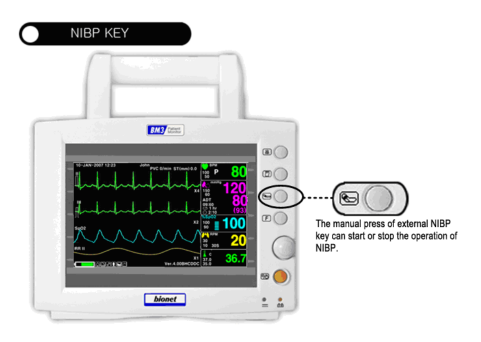
The maintenance is performed every 2 years.

Check the following list devise to operates properly and safety at all times.

- 1. Check for proper cuff size.
- 2. Check for residual air left in the cuff from a previous measurement.
- 3. Make sure cuff is not too tight or too loose.
- 4. Make sure cuff and heart are at same level, otherwise hydrostatic pressure will offset the NIBP value.
- 5. Minimize patient movement during measurement.
- 6. Watch for pulses paradox us.
- 7. Check for leak in cuff or tubing.
- 8. Patient may have a weak pulse.

8.2 NIBP Data Window





8.3 NIBP Setup

ALARM: A menu to set the Alarm

STAT: Start 5 minutes of continuous, sequential NIBP measurements.

CUFF SIZE: A menu to select cuff size

UNIT SELECT: A menu to select the pressure unit

INTERVAL : A menu to set Interval time when measures the blood pressure periodically

INFLATION: Initial Pressurization setting menu

MAIN MENU	ALARM	NIBP STAT: OFF	CUFF SIZE: ADT
	UNIT SELECT: mmHg	INFLATION SET: 170mmHg	INTERVAL: OFF

ALARM

The alarm provides ALARM LIMIT, LEVEL and ALARM SOUND.

The diam provides ALARWI LIWIT, LEVEL and ALARWI SOOND.			
MAIN MENU	ALARM	NIBP STAT: OFF	CUFF SIZE: ADT
	UNIT SELECT: mmHg	INFLATION SET: 170mmHg	INTERVAL: OFF
MAIN MENU	ALARM LIMIT	ALARM SOUND: ON	
PREV MENU	ALARM LEVEL		

ALARM LIMIT

Alarm setting Numeric Value of Systolic, Diastolic, and mean pressure is 10 ~ 360mmHg.

- 1. Move the ▶ mark to select one from RETURN, NIBP-S, NIBP-M, or NIBP-D, and press.
- 2. Press the key at NIBP-S, and move to LOW, and press again.(The user gets the same result regardless of the LOW-HIGH, or HIGH-LOW order.)
- 3. When the color has changed, move it again to select a target value, and press.
- 4. Press the key at HIGH. When the color has changed, move to the right to select a target value, and press.
- 5. Set up or revise the values of NIBP-M and NIBP in the same way as above.
- 6. With the selection of RETURN, the user can get out of the window.

MAIN MENU	ALARM LIMIT	ALARM SOUND: ON	
PREV MENU	ALARM LEVEL		

	NIBP ALARM LIMIT		
RETURN	UNITS	LOW	HIGH
NIBP-S	mmHg	80	200
NIBP-M	mmHg	40	140
NIBP-D	mmHg	20	120
			J

ALARM LEVEL

Set the order of priority in alarm.

MAIN	ALARM	ALARM	
MENU	LIMIT	SOUND	
PREV MENU	ALARM LEVEL		

ALARM LEVELS		
RETURN	ALARM LEVEL	
NIBP	MEDIUM	

ALARM SOUND

The menu which decide activate of warning sign and message display when the respiration alarm is on.

MAIN MENU	ALARM LIMIT	ALARM SOUND: ON	
PREV MENU	ALARM LEVEL		

NIBP STAT

Start 5 minutes of continuous, sequential NIBP measurements.

MAIN MENU	ALARM	NIBP STAT: OFF	CUFF SIZE: ADT
	UNIT SELECT: mmHg	INFLATION SET: 170mmHg	INTERVAL: OFF

CUFF SIZE

The user can select a CUF between ADULT, PEDIATRIC and NEONATAL.

MAIN MENU	ALARM	NIBP STAT: OFF	CUFF SIZE: ADT
	UNIT SELECT: mmHg	INFLATION SET: 170mmHg	INTERVAL: OFF
MAIN MENU	ALARM	CUFF SIZE:	> ADT PED
	UNIT SELECT: mmHg		NEO

UNIT SELECT

It is a function to set blood pressure measurement unit.

The blood pressure measurement unit provides mmHg and kPa.

MAIN MENU	ALARM	NIBP STAT: OFF	CUFF SIZE: ADT
	UNIT SELECT: mmHg	INFLATION SET: 170mmHg	INTERVAL: OFF
MAIN MENU	ALARM	NIBP STAT: OFF	CUFF SIZE: ADT
	UNIT SELECT: kPa	INFLATION SET: 170mmHg	INTERVAL: OFF

INTERVAL

This menu is used for selecting intervals when measures the blood pressure automatically. Select a target interval from 1min, 2, 3, 4, 5, 10, 15, 20, 30, 1hour, 2, 4, 8.

		1, 0, 10, 10, 20, 00, 111001, 2	
ΜΔΙΝΙ	MAIN	NIBP	CUFF
	ALARM	STAT:	SIZE:
MENU		OFF	ADT
	UNIT SELECT:	INFLATION SET:	INTERVAL: OFF
	mmHg	170mmHg	
MAIN MENU	INTERVAL: OFF	> OFF 1MIN. 2MIN. 3MIN.	15MIN. 20MIN. 30MIN. 1H
		4MIN.	2H
		5MIN.	4H
		10MIN.	8H

INFLATION SET

It is a function for pressurization pressure.

Numeric value is 80, 100, 120, 140, 160, 180, 200, 220, and 240.

MAIN MENU	ALARM	NIBP STAT: OFF	CUFF SIZE: ADT
	UNIT SELECT: mmHg	INFLATION SET: 80mmHg	INTERVAL: OFF
MAIN MENU	ALARM	NIBP STAT: OFF	CUFF SIZE: ADT
	UNIT SELECT: mmHg	INFLATION SET: 240mmHg	INTERVAL: OFF

Warning

Periodically check patient limb circulation distal to the cuff. Check frequently when using auto NBP in 1 and 2 minute intervals. Intervals below 10 minutes are not recommended for extended periods of time.

Warning

Pay attention to not to block connecting hose when you put cuff on patient.

NIBP Status Messages

Below is a list of system status alarm messages which may be displayed in the NIBP parameter window during monitoring.

Status Message	Monitor Response	Solution
OVER	System status alarm.	Remove cuff and contact
PRESSURE	Auto mode will shut off after ONE message.	service.
INFLATION FAIL. CHECK CUFF	System status alarm.	Check cuff, connections, and tubing.
DEFLATION FAIL. CHECK CUFF	System status alarm. Auto mode will shut off after ONE message.	Remove cuff and contact service.
PULSE TOO WEAK	System status alarm. Auto mode will shut off after ONE message.	Check patient and cuff placement.
EXCESSIVE MOTION	System status alarm. Auto mode will shut off after ONE message.	Possible excessive patient movement. Check patient.
MEASUREMENT System status alarm. Auto mode will shut off after ONE message.		Possible excessive patient movement or arrhythmia condition. Check patient.

Erroneous NIBP measurement

- Check for proper cuff size
 - 1. Too small a cuff can give an erroneously high value.
 - 2. Too large a cuff can give an erroneously low value.
- Check for residual air left in the cuff from a previous measurement.
- Make sure cuff is not too tight or too loose.
- Make sure cuff and heart are at same level, otherwise hydrostatic pressure will offset the NIBP value.
- Minimize patient movement during measurement.
- Check for leak in cuff or tubing.
- Patient may have a weak pulse.

9. EtCO2

9.1 INTRODUCTION

Position of EtCO₂ Connector and Accessory EtCO₂ ACCESSORY

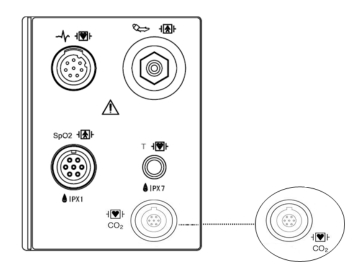
9.2 EtCO₂ Parameter Window

9.3 EtCO₂ Parameter Setting Menu

9.1 Introduction

ETCO2(End-Tidal CO2) is a device to see the concentration of end-tidal carbon dioxide, which uses a method of measurement based on the non-dispersed IR absorption of CO2 using IR ray by sampling a certain part of respiration through pipe during respiration.

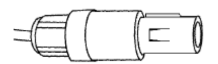
EtCO2 connector position and accessory (Sidestream, Respironics) EtCO2 Connector



LoFlo sidestream CO2 sensor and connector









Sidestream sensor connector

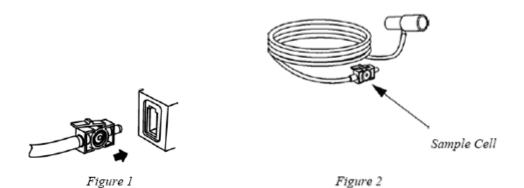
EtCO2 accessories for sidestream applications

EtCO2 monitoring accessory uses the accessories for LoFlo™ sidestream module of Respironics Company.

The airway adap	The airway adapters for sidestream intubated applications		
3473ADU-00		Airway Adapter	Weight: 4.5 grams
	1507	Kit w/	Deadspace – adds approximately 7
		Dehumidification	cc of deadspace
		Tubing	Intended for use when
			monitoring patients with ET
			Tube sizes >4.0 mm
3473INF-00		Airway Adapter	Weight: 5.8 grams
		Kit w/	Deadspace – adds approximately 1
		Dehumidification	cc of deadspace
		Tubing	Intended for use when
			monitoring patients with ET
			Tube sizes <=4.0 mm

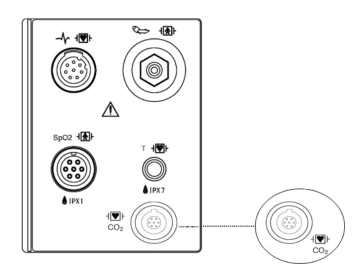
Connecting the LoFlo Sample Kit

1. The sample cell of the sampling kit must be inserted into the sample cell receptacle of the LoFlo CO₂ Module as shown in Figure 1. A "click" will be heard when the sample cell is properly inserted.



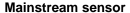
- 2. Inserting the sample cell into the receptacle automatically starts the sampling pump. Removal of the sample cell turns the sample pump off.
- 3. To remove the sampling kit sample cell from the sample cell receptacle, press down on the locking tab and pull the sample cell from the sample cell receptacle.

EtCO2 connector position and accessory (Mainstream, Respironics) **EtCO2** Connector



CAPNOSTAT 5 mainstream CO2 sensor and connector









Mainstream sensor connector

EtCO2 accessories for mainstream applications

EtCO2 monitoring accessory uses the accessories for CapnoStat 5 microstream sensor of Respironics Company.

6063-00	Single-Patient Use Airway Adapter
6312-00	Single-Patient Use Airway Adapter
7007-00	Reusable Airway Adapter
7053-00	Reusable Airway Adapter

Connecting the CAPNOSTAT® 5 CO2 Sensor to the Host System

1. Insert the CAPNOSTAT 5 CO₂ Sensor connector into the receptacle of the host monitor as shown in Figure 1.

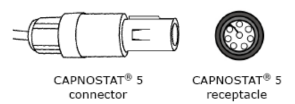
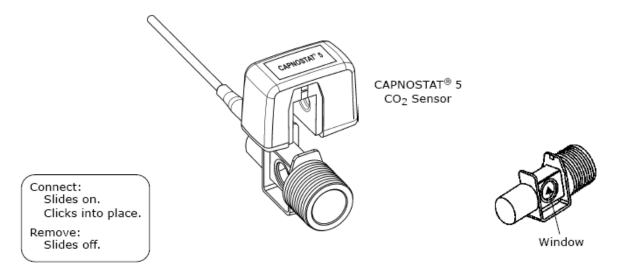


Figure 1

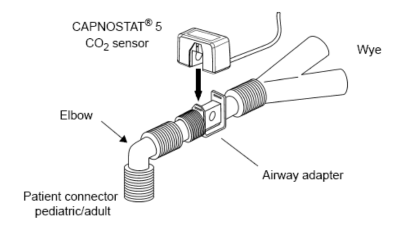
- 2. Make sure the arrows on the connector are at the top of the connector and line up the two keys of the connector with the receptacle and insert.
- 3. To remove the connector, grasp the body portion of the connector back and remove.

Note: Do not remove by pulling cable.

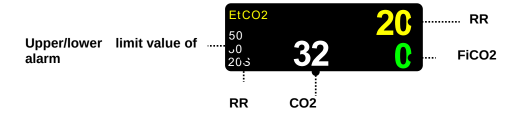
Shown below is the CAPNOSTAT 5 CO₂ Sensor connection to a Respironics Novametrix CO₂ adapter



Shown below is the CAPNOSTAT 5 CO2 Sensor with a patient circuit:



9.2 EtCO2 Parameter Window



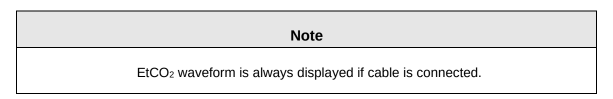
S: Display of apnea setting time in second unit

Upper/lower limit value of alarm: Display of alarm setting range value for concentration of CO2

EtCO2: Display of concentration value of carbon dioxide

RR: Display of the number of respirations per miniute

FICO2: Display of concentration value of carbon dioxide during inspiration



9.3 EtCO2 Parameter Setting Menu

ALARM LIMITS: A menu to set the alarm limit

STANDBY: A menu to set the power saving status of EtCO2 module

SCALE: A menu to set the screen scale of measured waveform

SETTINGS: A menu to handle the information of EtCO2 signal

MAIN MENU	ALARM	SWEEP SPEED: 6.25mm/s	WAVEFORM SCALE: 40mmHg
	APNEA DETECT: OFF	SETTINGS	ZERO

ART LIMIT(Upper/lower limit value of alarm)

Upper/lower limit value of alarm differs depending on the position of measurement.

The basic setting range of alarm setting value for EtCO2, FiCO2, RR, APNEA.

MAIN MENU	ΔΙ ΔΡΜ	SWEEP SPEED: 6.25mm/s	WAVEFORM SCALE: 40mmHg
	APNEA DETECT: OFF	SETTINGS	ZERO
MAIN MENU	ALARM LIMIT	ALARM SOUND: OFF	
PREV MENU			

EtCO2 ALARM LIMIT					
RETURN	UNITS	LOW	HIGH		
EtCO2	mmHg	25	50		
FiCO2	mmHg	0	5		
AWRR	RPM	10	30		
APNEA	SEC	0	20		

The following table shows standard alarm limit of parameter and setting value of scale when setting the label.

Danamatan	Adult		Neonatal			
Parameter	Low	High	Scale	Low	High	Scale
EtCO2	0	98		0	98	
FiCO2	0	20		0	20	
AWRR	0	100	40	0	100	40
APNEA	0	40		0	40	

EtCO2 SWEEP SPEED

EtCO2 speed is 6.5mm/s.

Speed is changeable to 6.25, 12.5, 25mm/s.

MAIN MENU	ALARM	SWEEP SPEED: 6.25mm/s	WAVEFORM SCALE: 40mmHg
	APNEA DETECT: OFF	SETTINGS	ZERO
MAIN MENU	ALARM	SWEEP SPEED: 6.25mm/s	> 6.25mm/s 12.5mm/s
	APNEA DETECT: OFF		25mm/s

WAVEFORM SCALE (Measured waveform scale setting)

This sets the range of measured waveform versus pressure.

Selectable numerical value means the maximum pressure range value that is shown with waveform. Pressing the knob switch key and then selecting the desired range value displays the selected pressure range value below the upper dotted line among two dotted lines in the left middle of wave window.

wiridow.			
MAIN MENU	ALARM	SWEEP SPEED: 6.25mm/s	WAVEFORM SCALE: 40mmHg
	APNEA DETECT: OFF	SETTINGS	ZERO
MAIN MENU	ALARM	WAVEFORM SCALE: 40mmHg	> 40mmHg 50mmHg 60mmHg
	APNEA DETECT: OFF		80mmHg 100mmHg

SETTINGS (Various setting)

Different menus are applied to provide menu and information for handling the EtCO2 module.

MAIN MENU	ALARM	SWEEP SPEED: 6.25mm/s	WAVEFORM SCALE: 40mmHg
	APNEA DETECT: OFF	SETTINGS	ZERO
MAIN MENU	MODULE SETUP	MODULE INFO	EtCO2 RESET
PREV MENU			

MODULE INFO SET			
RETURN	CONTENTS		
BAROMETRIC PRESSURE	760 mmHg		
GAS TEMPERATURE	0.0 🗆		
NO BREATH DETECT TIMEOUT	0 SEC		
O2 COMPENSATION	21 %		
ANESTHETIC AGENT	0.0 %		
BALANCE GAS	ROOM AIR		
CURRENT ETCO2 TIME PERIOD	0000-00-00-00		
CURRENT CO2 UNIT	mmHg		
SLEEP MODE	NORMAL OP		
ZERO GAS TYPE	ZERO ON N2		
DISABLE SAMPLING PUMP	NORMAL OP		

BAROMETRIC PRESSURE: This setting is used to set current Barometric Pressure.

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.

NO BREATH DETECT TIMEOUT: 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 Capnostat will signal no breaths detected.

O2 COMPENSATION ANESTHETIC AGENT BALANCE GAS:

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.

CURRENT ETCO2 TIME PERIOD: This setting is used to set the calculation period of the

ETCO₂ value. The end-tidal CO₂ value is the highest peak CO₂ 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 ETCO₂ value for the last

two breaths.

CURRENT CO2 UNIT: Continuous waveform mode commands (the CO2 Waveform

Mode command [command 80h] and the CO_2/O_2 Waveform Mode command [command 90h]) MUST NOT be active when this command is used otherwise this command will be ignored

and the setting will remain unchanged.

SLEEP MODE: Sleep mode is used to save power when the host monitor is in

standby mode. There are two sleep modes available for the Capnostat. Using Sleep Mode 1 maintains the heaters so the Capnostat is able to run immediately after exiting the sleep mode. Mode 2 will require the Capnostat to go through its warm

up sequence when exiting this mode and a delay will be

introduced until the system has stabilized.

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.

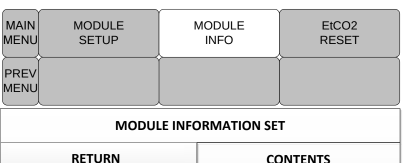
DISABLE SAMPLING PUMP: This setting allows the pump to be forced off. In Normal

Operating Mode, the pump will be turned on when the sampling cell is connected and no pneumatic system errors are detected.

In Pump Disabled Mode, the pump will remain off in all

circumstances.

MODULE INFO



RETURN	CONTENTS
SENSOR PN	01DA
OEM ID	0X57DA
SENSOR SN	SN01234
HW REVISION NUM	001.3
TOTAL USE TIME	60 MIN.
LAST ZERO TIME	5 MIN.
PUMP TOTAL USE TIME	30 MIN.
PUMP MAX USE TIME	35 MN.

APNEA ALARM: This performs a function to set the display of apnea message alarm.

This displays a "apnea" message at the center of parameter window as shown in the figure below with apnea alarm on in case of apnea until the set apnea period is passed through.



With apnea alarm off, measured values are displayed instead of message.



Warning

If defibrillation is performed while doing CO2 monitoring, remove the CO2 FilterLine from patient Getting in touch with sensor cable without removing the FilterLine can result in serious electrical burn, shock, or injury due to electric discharge energy.

Note

In the following monitoring conditions, the measured values may be inaccurate. Read the measured values carefully.

- 1. When using this in an environment of using nitrous oxide gas of high concentration
- 2. When using this in an environment where abrupt temperature change takes place
- 3. When using this in an environment with severely high humidity.

Caution

- The measured values may be inaccurate when using this equipment for patients who have very fast or irregular respiration.
- When measuring CO2 from the patient under the anesthesia, check it when gas mixture comes in. Otherwise, the measured result values may be inaccurate.
- When using a anesthesia machine that uses a volatile anesthetic, CO2 values may be inaccurate.

10. TEMPERATURE

10.1 Outline

Temperature Connector and Measuring Cable

10.2 Temperature Data Window

10.3 Temperature Setup

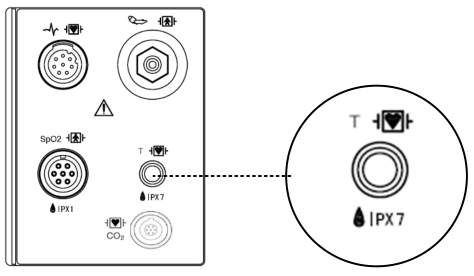
ALARM
ALARM LIMIT
ALARM LEVEL
ALARM SOUND
UNIT SELECT

BM3OM-2.62

10.1 Outline

This function is used to indicate the changes of resistance generated by the changes of temperature in numbers. The function involves the process of transferring the changes into electric signals.

Temperature Connector and Measuring Cable Temperature Connector



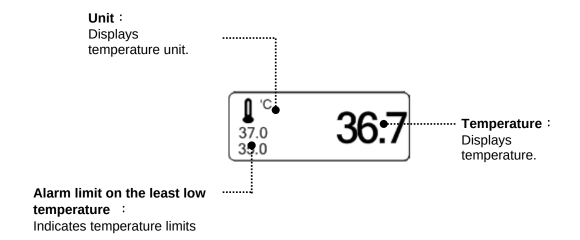
Temperature Measuring Cable



Note

Temperature probe is correctly positioned and fixed to do not disconnect on the patient. The TEMP cable connector is a high-insulation port and it is defibrillator-proof($^{\frac{1}{|\mathbf{v}|}}$).

10.2 Temperature Data Window



Note

The minimum measuring time required to obtain accurate readings at the specific body site is at least 3 minutes.

10.3 Temperature Setup

ALARM: Temperature measurement alarm set UNIT SELECT: Temperature measurement unit set

MAIN MENU	ALARM	UNIT SELECT: °C

ALARM

Alarm menu provide ALARM LIMIT, LEVEL and ALARM SOUND.

MAIN MENU	ALARM		UNIT SELECT: °C
MAIN MENU	ALARM LIMIT	ALARM SOUND : ON	
PREV MENU	ALARM LEVEL		

ALARM LIMIT

Setting numeric value is $0^{\circ}\text{C} \sim 50.0^{\circ}\text{C}$.

- 1. Move the ightharpoonup mark to select either RETURN or TEMP, and press.
- 2. After pressing the cursor at TEMP, move it to LOW, and press.
- 3. When the color has changed, move the cursor again to select a target value, and press.
- 4. Move the cursor to HIGH and press. After the color has changed, move the cursor again to select a target value, and press. (One may choose HIGH first to get the same result.)
- 5. Select RETURN to get out of the menu.

MAIN MENU	ALARM LIMIT	ALARM SOUND : ON	
PREV MENU	ALARM LEVEL		

TEMPERATURE ALARM LIMIT					
RETURN	UNITS	LOW	HIGH		
TEMP	°C	30.0	42.0		

ALARM LEVEL

Set the order of priority in alarm.

Cot the organ or phoney in claim.					
MAIN MENU	ALARM LIMIT	ALARM SOUND			
PREV MENU	ALARM LEVEL				
ALADMIT/FLS					

ALARM LEVELS				
RETURN	ALARM LEVEL			
TEMP PROBE OFF	ALARM LEVEL MESSAGE MESSAGE			

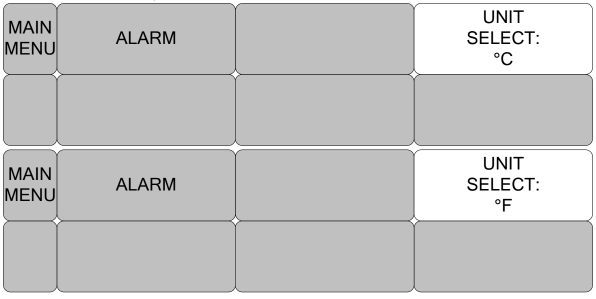
ALARM SOUND

The menu which decide activate of warning sign and message display when the respiration alarm is on.

MAIN MENU	ALARM LIMIT	ALARM SOUND : ON	
PREV MENU	ALARM LEVEL		
MAIN MENU	ALARM LIMIT	ALARM SOUND : OFF	
PREV MENU	ALARM LEVEL		

UNIT SELECT

Able to select unit with °C, °F.



Check list

- 1. The temperature probe(YSI 400 series) is correctly positioned on the patient.
- 2. Temperature cable is attached to the monitor.
- 3. Temperature setup is adjusted, if necessary. Follow detailed procedures within this chapter.

TEMP Message

If you experience some problems with temperature monitoring, one of the following messages may be displayed in the TEMP parameter window.

- PROBE OFF: Probe is not properly connected. Check the probe.
- No temperature value will be displayed . Service on the monitor is required.

Warning

To measure the peripheral temperature, attach the probe to the ankle or palm. If the patient sweats heavily or moves violently, fasten the pad with surgical tape.

NOTE

When the measuring site is exposed directly to air, the temperature may be lower than normal. It take about 20 to 30 minutes to reach the equilibrium temperature after attaching the sensor.

11. PRINT

11.1 Print

Printer and Heat Sensitivity Paper Function and Setup Menu

11.2 Paper Change

BM3OM-2.62

11.1 Print

Printer and Heat Sensitivity Paper

A printer used to print data onto thermal paper.

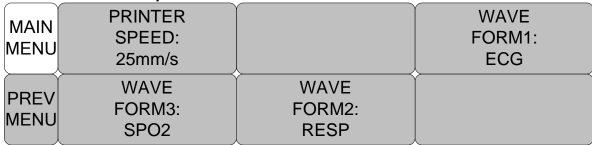
Size of the thermal paper roll: 58mm wide x 38mm in diameter any thermal paper of same size can be used for the printer.

Side View of Printer



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Function and Setup Menu



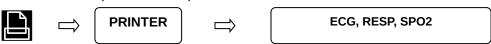
- 1. Press the PRINT Key for continuous printing.
- 2. Select Printing Speed 25, 50 mm/s.

MAIN MENU	PRINTER SPEED: 25mm/s		WAVE FORM1: ECG
PREV	WAVE FORM3: SPO2	WAVE FORM2: RESP	
MAIN MENU	PRINTER SPEED: 50mm/s		WAVE FORM1: ECG
PREV MENU	WAVE FORM3: SPO2	WAVE FORM2: RESP	

3. Set up ALARM PRINT in the MORE menu to activate ALARM during printing.



4. Data is printed in a selected wave form along with personal information of the patient. 3 channels select 3 parameters to print.



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MAIN MENU	PRINTER SPEED: 50mm/s		WAVE FORM1: ECG
PREV MENU	WAVE FORM3: SPO2	WAVE FORM2: RESP	
MAIN MENU	PRINTER SPEED: 50mm/s	WAVE FORM1: ECG	OFF > ECG SPO2
PREV MENU	WAVE FORM3: SPO2		RESP
MAIN MENU	PRINTER SPEED: 50mm/s		WAVE FORM1: ECG
PREV MENU	WAVE FORM3: SPO2	WAVE FORM2: RESP	
MAIN MENU	PRINTER SPEED: 50mm/s	WAVE FORM2: RESP	OFF ECG SPO2
PREV MENU	WAVE FORM3: SPO2		> RESP
MAIN MENU	PRINTER SPEED: 50mm/s		WAVE FORM1: ECG
PREV MENU	WAVE FORM3: SPO2	WAVE FORM2: RESP	
MAIN MENU	WAVE FORM3: SPO2	OFF ECG > SPO2	WAVE FORM1: ECG
PREV MENU		RESP	

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If there is no print sheet, no paper icon of appears.

11.2 Paper Change

1

Open the window of the printer.



2

Insert the paper roll offered with the product into the printing unit. Place the roll in a proper way so that the printed paper can roll out upwards.



3

Press the printer window until it is properly shut. Inaccurate shutting may cause failure in printing.



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12. MESSAGE LIST

Function	Message	Details
ECG	LEAD FAULT	Lead Cable is not properly connected.
SpO2	PROBE OFF CHECK PROBE PULSE SEARCH POOR SIGNAL LOST PULSE ARTIFACT	Probe is not properly connected. Patient's finger is off the probe. Detection by the monitor of a pulse has ceased. The SpO2 signal is too low. The quality of the signal is questionable. The signal is patient's motion artifact
RESP	LEAD FAULT APNEA	Lead Cable is not properly connected. APNEA gives an alarm.
NIBP	INFLATION FAILURE CHECK CUFF OVER PRESSURE DEFLATION FAILURE CHECK CUFF OVER TIME PRESSURE MEASUREMENT ERROR	Cuff hose is not properly connected. Cuff pressure is putting on excessively. Cuff is bent, preventing deflation. Measure time exceeds the preset Level. Measure signal absent
TEMP	PROBE OFF	Probe is not properly connected.
ALARM	ALARM VOL.OFF SILENCED ALARM PAUSE 2MIN ALL ALARM OFF	Alarm volume is off. Alarm key is pressed once Alarm key is pressed twice The all alarm is cleared, the alarm sounds and the lamp does not occur.
PRINT	No paper Icon	No paper in the printer
BATTERY	BATTERY LOW	The battery level is low, automatically power off within 5 minutes.

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13. DEFAULT SETTING VALUE

1. Adult-ICU Mode

Alarm level

	High	Medium	Low	Message
HR		0		
NIBP		0		
SpO ₂			0	
SpO ₂ -Rate				0
RR				0
T(°C)				0
EtCO2			0	
FiCO2				0
AWRR			0	

Parameter Limits

	Low	High
HR	50	150
NIBP-S	80	200
NIBP-M	40	140
NIBP-D	20	120
SpO ₂	90	100
SpO ₂ -Rate	50	150
RR(RESP)	10	30
T(°C/ °F)	30.0/86.0	42.0/107.6
AWRR	10	30
EtCO2	25	50
FiCO2	0	5

Display

Patient Age	Adult
Color format	Color
Primary ECG	II
Arrhythmia	Off
Detect Pace	Off
Print Waveform2	Off
Print Waveform3	Off
Alarm Print	On
NIBP Auto	Off
NIBP Cuff Size	ADT
RR(RESP) Lead	II
Alarm Volume	50%
QRS Volume	Off
Pulse Volume	Off
ECG Lead Fault	Message
SpO ₂ Probe Off	Message
Units for Height	cm
Units for Weight	kg
Temperature Units	Ů C
NIBP Limit Type	Systolic
ECG Filter	Monitoring

2. Neonate-ICU Mode

Alarm level

	High	Medium	Low	Message
HR		0		
NIBP		0		
SpO ₂			0	
SpO ₂ -Rate				0
RR				0
T(0
EtCO2			0	
FiCO2				0
AWRR			0	

Parameter Alarm Limits

	Low	High
HR	90	200
NIBP-S	40	100
NIBP-M	30	70
NIBP-D	20	60
SpO ₂	88	100
SpO ₂ -Rate	90	200
RR(RESP)	15	100
T(°C/ °F)	30.0/86.0	42.0/107.6
EtCO2	25	50
FiCO2	0	5
AWRR	15	100

Display

Display	
Patient Age	0~2 years
Color format	Color
Primary ECG	II
Arrhythmia	Off
Detect Pace	Off
Print Waveform2	Off
Print Waveform3	Off
Alarm Print	On
NIBP Auto	Off
NIBP Cuff Size	NEO
RR(RESP) Lead	II
Alarm Volume	50%
QRS Volume	Off
Pulse Volume	Off
ECG Lead Fault	Message
SpO ₂ Probe Off	Message
Units for Height	cm
Units for Weight	kg
Temperature Units	் C
NIBP Limit Type	Systolic
ECG Filter	Monitoring

3. Pediatric-ICU Mode

Alarm level

	High	Medium	Low	Message
HR		0		
NIBP		0		
SpO ₂			0	
SpO₂-Rate				0
RR				0
T(°C)				0
EtCO2			0	
FiCO2				0
AWRR			0	

Parameter Alarm Limits

	Low	High
HR	70	180
NIBP-S	60	160
NIBP-M	40	120
NIBP-D	30	100
SpO ₂	90	100
SpO ₂ -Rate	70	180
RR(RESP)	10	50
T(°C/ °F)	30.0/86.0	42.0/107.6
AWRR	10	50
EtCO2	25	50
FiCO2	0	5

Display

Patient Age	3~16years
Color format	Color
Primary ECG	II
Arrhythmia	Off
Detect Pace	Off
Print Waveform2	Off
Print Waveform3	Off
Alarm Print	On
NIBP Auto	Off
NIBP Cuff Size	PED
RR(RESP) Lead	11
Alarm Volume	50%
QRS Volume	Off
Pulse Volume	Off
ECG Lead Fault	Message
SpO ₂ Probe Off	Message
Units for Height	cm
Units for Weight	kg
Temperature Units	் C
NIBP Limit Type	Systolic
ECG Filter	Monitoring

SPOT MODE

- 1. General Operation
- 2. Patient/Data Management
 - 3. Save Record
- 4. Saved Data Management
 - 5. Setup
 - 6. NIBP
 - 7. SpO2
 - 8. Temperature
 - 9. PRINT

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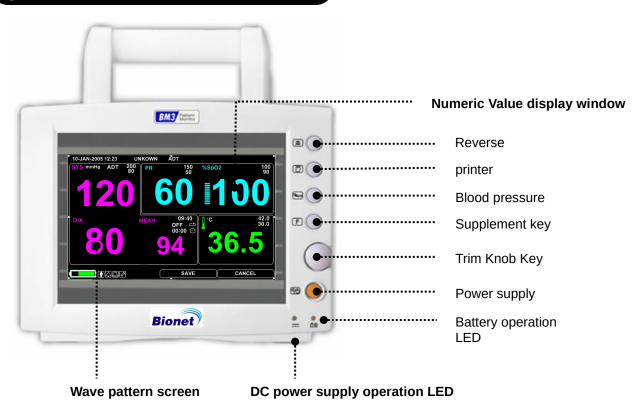
1. General Operation

1.1 Function and key

The product has LCD screen and 5 functional keys and 1 trim knob.



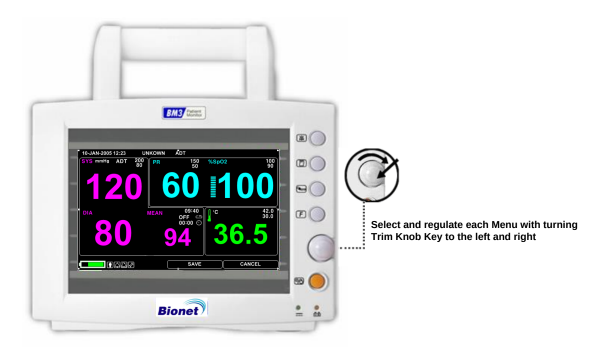
Operating the BM3 Spot Monitor



Operation Keys

- 1. Power: ON/OFF of the equipment
- Supplement Key: Using home key to bring up the menu. Adjust view mode while out of menu/list.
- 3. Blood Pressure : Able to manage blood pressure measurement with manual operation.
- 4. Print: Print selected wave pattern in the menu. It prints continuously until press the key to stop.
- 5. Alarm : Turn off the alarm when alarm rings.
 - Press once, the alarm is off for 1 minute.
 - Press twice, all alarm stops for 2 minutes.
 - Press three times, all alarm off
 - Press four times, the alarm returns.
- 6. Trim Knob Key: Move cursor turning with Trim Knob Key to the left and to the right on each menus and press it to select.





1.2 Screen Generating Power Mode

There are 3 types of screen generating power mode.

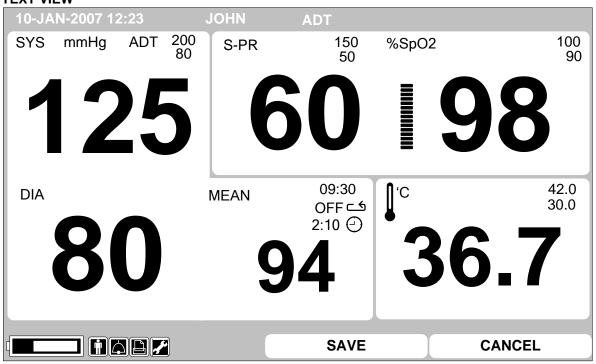
Select the screen generating mode icon or press supplement key to change the screen generating power.

TEXT VIEW (test generating mode): Display the bigger number on the screen.

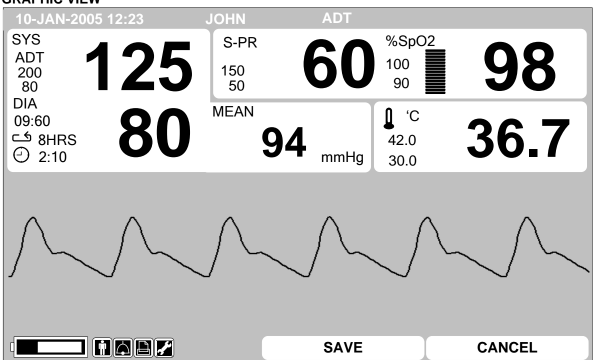
GRAPHIC VIEW (wave pattern generation mode): Generate parameter numeric value and SPO2 wave pattern together.

RECORD LIST VIEW (record list generating mode): Print Record list and parameter numeric value together.

TEXT VIEW



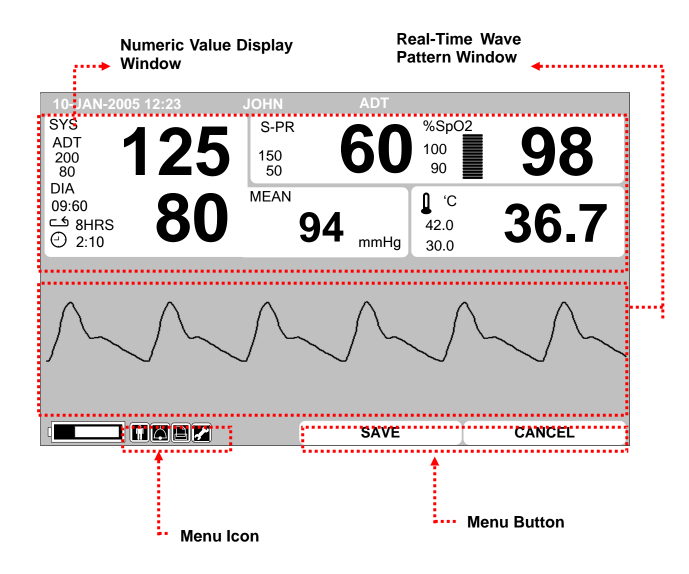
GRAPHIC VIEW



RECORD LIST VIEW

ECOND LIST VII								
10-JAN-2005	12:23		JOHN	ADT				
SYS			S-PR	01	%SpC	02		
ADT 200	17	5	150	6(100		86	
80		J	50	O	90			
DIA 09:60								
SHRS ALL Q			14	42.0	3	6.7		
② 2:10				mmH	g 30.0			
RET Pati	ient T	Date	TIME	PR	NIBP(mml	Hg) SpO2	Temp('C)	
P20072012	32 A	10-01	09:20:3	80(S)	150/90(1	15) 99	36.9	
P20071819	42 P	10-01	10:30:2	20 70(S)	132/71(9	92) 100	37.1	
Unknowr	n A	10-01	10:45:3	80(S)	164/110(1	130) 99	37.2	
Unknowr	Unknown N		11:20:2	20 75(S)	124/74(9	91) 98	36.8	
P2007081511 A		10-01	11:40:3	60(S)	128/80(9	94) 99	36.2	
					/E	CA	NCEL	

1.3 Standard Menu Operation Screen Organization



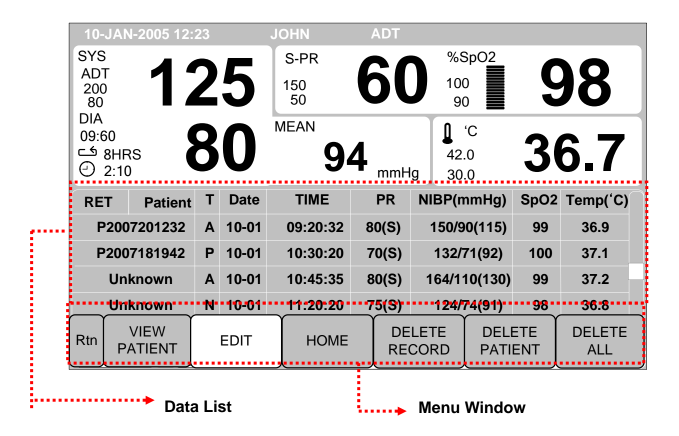
Real Time Wave Pattern Window : Print measured Wave Pattern Window

Numeric Value Window: There are 3 windows in it and each window displays analyzed data and

setting status.

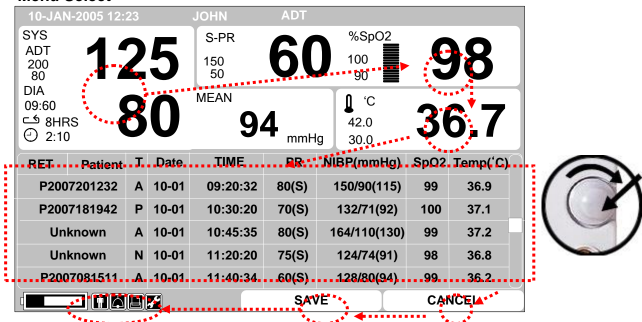
Menu Icon: The menu to select the icon.

Menu Button: A button to save the data or delete.



Menu Window: Menus appears on window. It appears when the menu activated. Data List: Display saved Data list.

Menu Select



When the Trim Knob Key is turned, Menus are selected in the order indicated above. The menus move to the right in the order of (NIBP) \rightarrow (SPO2) \rightarrow (TEMP) \rightarrow [(RECORD LIST)] \rightarrow (CANCEL) \rightarrow (SAVE) \rightarrow (SETUP) \rightarrow (PRINT) \rightarrow (ALARM) \rightarrow (PATIENT) An inactivated window is jumped off. Data list mode does not appear in the Large Parameter mode and Graphic View Mode

Menu Icon Composition



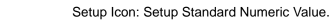
Patient Icon: Patient register and delete.



Alarm Icon: Setup alarm.

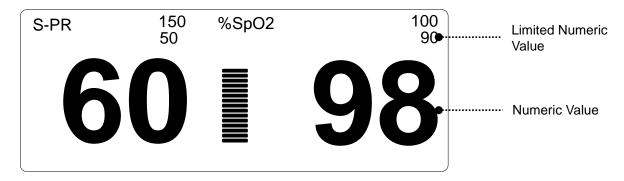


Printer Icon: Setup printer.



Numeric Value Window

It displays measured numeric value, functional setting, and limited numeric value.



Select Menu Using by Trim Knob Key

A right-hand turn makes a movement in a clockwise direction. A left-hand turn makes a movement in an anti-clockwise direction. A selection is made by pressing the Trim Knob Key.

Select Arrow Item Menu

Move to the left: Turn Trim Knob Key to the left.

Move to the right: Turn Trim Knob Key to the right.

Selection is made by pressing the Trim Knob Key. Exit out of the menu after the selection.

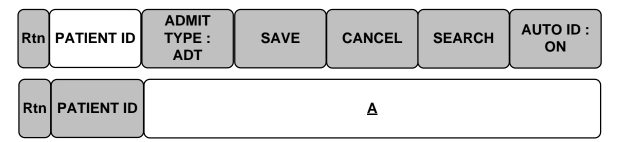
Rtn	PATIENT ID	ADMIT TYPE : ADT	SAVE	CANCEL	SEARCH	AUTO ID : ON
	ADMIT	<u> </u>	>ADT			
	TYPE :		PED.			
	ADT		NEO			

Letter Arrangement Menu

The following figure shows the screen where the word sequence menu is activated within the word sequence correction menu. Here, the cursor moves over the words when the Trim Knob Key is turned in the clockwise direction.

Rtn PATIENT ID	ADMIT TYPE : ADT	SAVE	CANCEL	SEARCH	AUTO ID : ON
Rtn PATIENT ID					

The above figure shows how the cursor moves on the screen. The cursor moves according to the direction the Trim Knob Key is turned. Press the Trim Knob Key if you want to change a letter currently on the screen.



The above figure shows how the cursor is selected to change a letter. Right-hand turning of the Trim Knob Key makes it possible to select in the order of A-Z, 0-9, and blank, while left turning makes the movement in the opposite direction.

Once a letter or a number is selected, the screen comes back to the condition where the same process of selection can be made. One may move to the menu item in the left of the screen to end the process, which is completed by pressing Trim Knob Key. After completion, the screen comes back to the earlier picture.

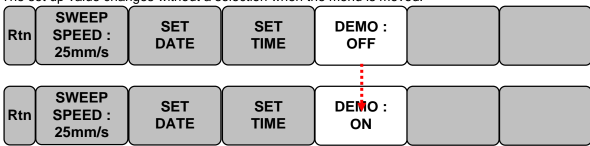
List selective menu

Whenever the square moves, a selected letter or a number is highlighted displaying its value.

RET	Patient	Т	Date	TIME	PR	NIBP(mmHg)	SpO2	Temp('C)
P2007	7201232	Α	10-01	09:20:32	80(S)	150/90(115)	99	36.9
P200	7081506	Р	10-01	10:30:20	70(S)	132/71(92)	100	37.1
Unk	nown	Α	10-01	10:45:35	80(S)	164/110(130)	99	37.2
Unk	Unknown		10-01	11:20:20	75(S)	124/74(91)	98	36.8
P2007	7081511	A	10-01	11:40:34	60(S)	128/80(94)	99	36.2

Operation Menu

The set up value changes without a selection when the menu is moved.



2. PATIENT/DATA MANAGEMENT

2.1 Outline

2.2 Admit Type

2.3 Select Patient in Admit Information

2.4 Alarm Outline

2.5 Alarm Setup

2.6 Alarm Limit Setup

2.7 Alarm Print

2.8 Alarm Volume

2.9 Alarm Level

2.10 Nurse Call

2.11 Alarm Sound

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2.1 Outline

Resister patient's ID and name to save data of each patient.

Divide to patient's ID and type.

Patient's type divided as adult, baby, and Infant.

The screen initializes after once saved patient's record in Spot mode.

Register the patient whenever you measure them or select from the patient's list to save the patient in Spot Mode.

Without registration of patient, the patient's ID is "UNKNOWN" (When selected off in AUTO ID) or "01 01 10 0000" (DD/MM/YY 0000 \sim 4000, When selected on in AUTO ID) and maintains previous numeric value in Type.

2.2 Admit Type

Select patient icon in Menu icon



Rtn PATIENT ID	ADMIT TYPE : ADT	SAVE	CANCEL	SEARCH	AUTO ID: ON
----------------	------------------------	------	--------	--------	----------------

Select ID menu in menu window and register patient ID. After the registration, select ID menu in previous menu window.



Select TYPE of menu in the menu window and register type of patient.

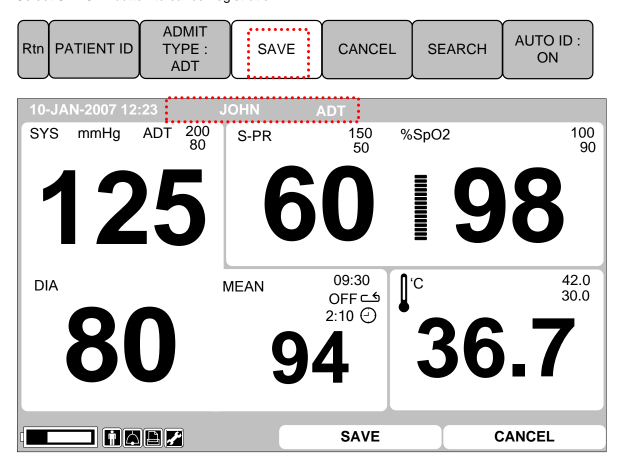
ADMIT

Rtn PATIENT ID	TYPE : ADT	SAVE	CANCEL	SEARCH	ON	
ADMIT TYPE : ADT		DT ED EO				s

elect save menu and complete patient registration.

Display registered patient's ID and Type on the top of the screen.

Select CANCEL button to cancel registration.



2.3 Select Patient in Admit Information

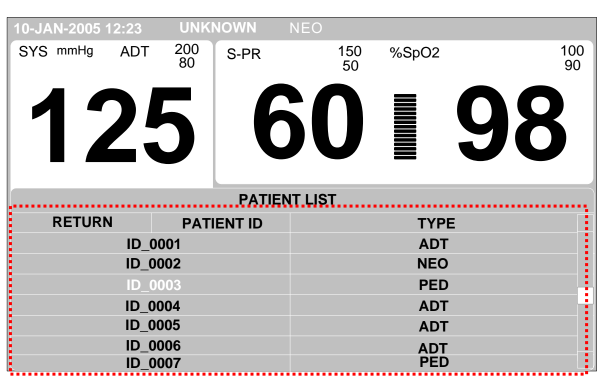
Able to select recoded patient in the patient list Select patient icon in menu icon.



Select search menu and Confirm patient list in menu window.

The patient list is the patient who already has measured data.

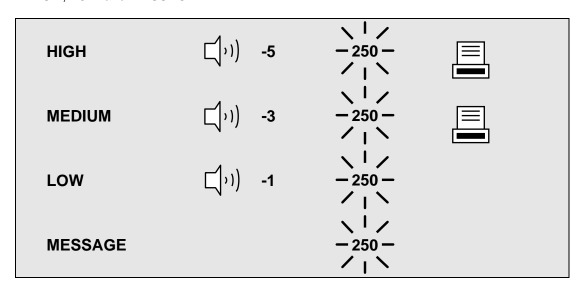




Select the patient's ID by using Trim Knob button then register. Select RETURN menu at the left top of the list to move to the top menu. Registered patient's ID and type displays on top of the screen.

2.4 Alarm Outline

Alarm is divided into two, alarm for the patient's condition and for the product's condition. The patient's alarm sounds when the diagnostic functions (ASYSTOLE, VTAC/VFIB, and VTAC) are detected. Each alarm sound differs in order in order and volume according to the levels of HIGH, MEDIUM, LOW and MESSAGE.



Display alarm sound and the number of ringing sound



Text flashes



Alarm lamp flashes



Print Wave Pattern

Product Status Alarm

The machine gives alarm sounds for its system with a related message flashing.

2.5 Alarm Setup

Select alarm icon in menu icon.





ALARM LIMITS: The machine enables one to see and change the limits of alarm for all parameter functions

ALARM PRINT: with an ON/OFF setup, the related information is printed out whenever an alarm is given.

ALARM VOLUME: volume of each alarm can be adjusted in 10 step. ALARM LEVEL: Priority of each parameter alarm can be set up.

NURSE CALL: Set the feature of the NURSE CALL.

ALARM SOUND : Set the ON/OFF feature of the ALARM SOUND.

2.6 Alarm Limit

The machine enables one to see and change the limits of alarm for all parameter functions.

10-JAN-2005 1	2:23		IOHN	ADT		·			
SYS mmHg	ADT	200 80	S-PR	6	150 50	%SpO2	98	100 90	
ALARM LIMIT									
RETURN PR		BP		LOW 50		HIGH			
SpO2-%		БР %				150 100			
•					90				
NIBP-S		mm			80		200		
NIBP-M		mmHg		40			140		
NIBP-D		mmHg		20		120			
TEMP		°C		30.0		42.0			

2.7 Alarm Print

With an ON/OFF setup, the related information is printed out whenever an alarm is given.

	ALARM ALARM VOLUME: ON OFF	ALARM LEVEL	NURSE CALL	ALARM SOUND
--	----------------------------	----------------	---------------	----------------

2.8 Alarm Volume

The volume of each alarm can be adjusted in 10 step.

Rtn	ALARM LIMIT	ALARM PRINT: ON	ALARM VOLUME: OFF	ALARM LEVEL	NURSE CALL	ALARM SOUND
Rtn	ALARM VOLUME: OFF	> OFF 10 % 20 %	30% 40% 50%		90% 00%	

2.9 Alarm Level

Priority of each parameter alarm can be set up.

10-JAN-2005	12:23	,	JOHN	ADT				
SYS mmHg	ADT	200 80	S-PR	150 50	%SpO2	100 90		
17		5	6		9	18		
		P/	ARAMETER A	ALARIM LEVI	ELS			
	RETU	RN		ALARM LEVEL				
		R		MESSAGE				
	SPC)2-%		LOW				
	NI	BP		MEDIUM				
	TE	MP		MESSAGE				
S-PROBE OFF				MESSAGE				
S-CHECK PROBE				MESSAGE				
T- PROBE OFF				MESSAGE				

2.10 Nurse Call

Set the feature of the NURSE CALL.

Rtn	ALARM LIMIT	ALARM PRINT: ON	ALARM VOLUME OFF	: ALARM LEVEL	NURSE CALL	ALARM SOUND			
	NURSE CALL SETUP								
	R	RETURN		CONTENTS					
	NU	JRSE CALL		OFF					
	NO	RMAL MODE		NORMAL OPEN					
	С	ALL MODE		ONE TIME					
						J			

- 1. NURSE CALL: ON/OFF
 - The nurse call function is enable or disable.
- 2. NORMAL MODE
 - NORMAL OPEN: Select this option when the hospital's call system is set to NORMAL OPEN.
 - NORMAL CLOSE: Select this option when the hospital's call system is set to NORMAL CLOSE.
- 3. CALL MODE
 - ONE TIME: When ONE TIME is selected, a nurse call signal is a pulse signal lasting 3s. When multiple alarms occur simultaneously, only one pulse signal will be output..
 - CYCLING: When CYCLING is selected, the duration of a nurse call signal is the same with the alarm, namely, from the time that the alarm occurs to the time it disappears. On and off repeatedly at intervals of 1 second.
 - CONTINUE: When CONTINUE is selected, the duration of a nurse call signal is the same with the alarm, namely, from the time that the alarm occurs to the time it disappears. However, lasts only one minute, then stops.

2.11 Alarm Sound

Set the ON/OFF feature of the ALARM SOUND.

ALARM ALARM

Rtn ALARM LIMIT	PRINT: ON	VOLUME: OFF	ALARM LEVEL	NURSE CALL	ALARM SOUND	
	PAF	RAMETER AL	ARM SOUND			
F	RETURN		PARAME	TER ALARM S	SOUND	
	SPO2		ON			
	NIBP		ON			
	TEMP		ON			

3. SAVE RECORD

- 3.1 Outline
- 3.2 Adjust to Record Save Mode
- 3.3 Measure with Monitor Mode
- 3.4 Measure with Manual Mode
 - **3.5 Save**
 - 3.6 Exit from Saving Mode

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3.1 Outline

There are two modes to save data. One is called MONITOR mode. It saves the patient's ID/TYPE without re-register once patient registered. The other called SPOT mode. It initializes the machine once Patient's record saved.

SPOT mode is good for measuring many patients. MONITOR mode is used to apply for monitoring only one patient's constantly.

3.2 Adjust to Record SAVE Mode

Select setup icon in icon menu.



When SAVE MODE menu selected in setup menu widow, whenever press Trim Knob Key mode switches to AUTO and MANUAL in turn.

Rtn DISPLAY	SAVE MODE : AUTO	USER SERVICE	SYSTEM	KEY SOUND : OFF	MAKER SERVICE
-------------	------------------------	-----------------	--------	-----------------------	------------------

3.3 Measure with Monitor Mode

Measure after setup mode to AUTO

Rtn	DISPLAY	SAVE MODE : AUTO	USER SERVICE	SYSTEM	KEY SOUND : OFF	MAKER SERVICE	
-----	---------	------------------------	-----------------	--------	-----------------------	------------------	--

It saves a measured data in 60 seconds.

Once NIBP measured, maintains measured data till the next measurement.

Not be able to delete measured Parameter data once save it in the machine completely Maintain ID and TYPE after complete saving.

Alarm limit numeric value does not change after saving.

If additional NIBP measure did not occur in the next 60 seconds then it is regard as NIBP measurement did not be performed.

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3.4 Measure with MANUAL Mode

Measure after setup a mode to MANUAL.



It saves as press the button after measurement.

NIBP ending spot numeric value stores when NIBP is INTERVAL mode.

When NIBP is MANUAL mode, it saves measured numeric value after 60seconds of event below.

Event: Input patient information

Measure NIBP Measure SpO2

When new event occur in 60 seconds after pervious event then it saves after 60 seconds of new event occur.

All measured parameter removes from the screen after finish with saving.

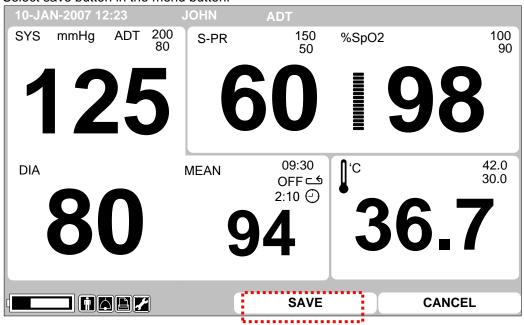
Search from record list to confirm measured result.

After saving, the patient ID initializes as UNKNOWN.

After saving, adjusted alarm limit numeric value becomes Default numeric value.

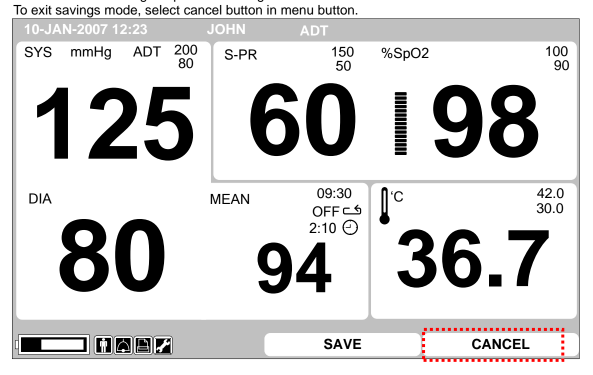
3.5 Save

It can be saved automatically by the user not only by AUTO or MANUAL mode. Select save button in the menu button.



BM3OM-2.62 3. SAVE RECORD 176

3.6 Exit from Saving ModeIt is used for exiting status of monitoring in monitor mode. It is used for initializing the patient who registered in MANUAL mode.



BM3OM-2.62 3. SAVE RECORD 177

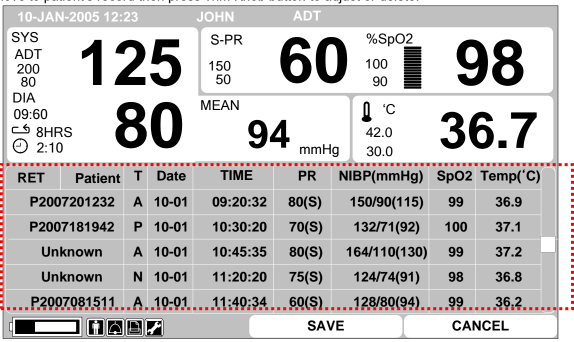
4. SAVED DATA MANAGEMENT

- 4.1 Record List View
- 4.2 Exit from Record List
- 4.3 View Specified Patients Record List
 - **4.4 View All Patients Record List**
 - 4.5 Adjust Record
 - 4.6 Delete a Record
 - 4.7 Delete a Patients Record
 - 4.8 Delete All Patients Record

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4.1 Record List View

Select in the List window and move inside of the list for Management. Turn Trim Knob button in inside of the list then move to records. Move to patient's record then press Trim Knob button to adjust or delete.



< Record List View >

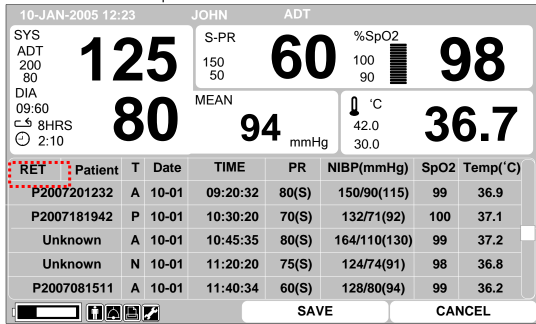
4.2 Exit from Record List

There are 4 ways to exit from Record List.

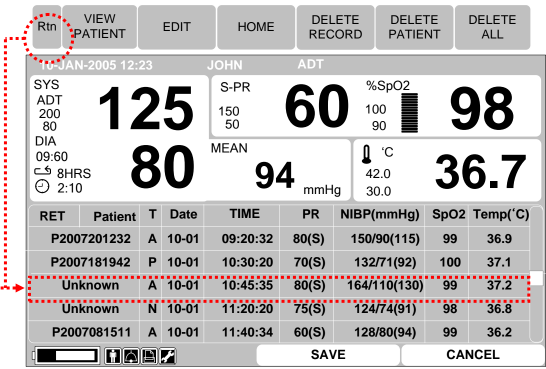
1. Press Home menu in the Menu.



2. Press return menu at the top of the record list window.



Press Rtn in the Menu. Then it will return to Record List.



4. Exit Menu simply by pressing the Supplement Key.

4.3 View Specified Patients Record List

Move to Record List window to view a patient Record List. Move to a patient's record by turning Trim Knob button.

RET	Patient	T	Date	TIME	PR	NIBP(mmHg)	SpO2	Temp('C)	
↑ P2007	7201232	Α	10-01	09:20:32	80(S)	150/90(115)	99	36.9	
P200	7081506	Р	10-01	10:30:20	70(S)	132/71(92)	100	37.1	
Unk	nown	Α	10-01	10:45:35	80(S)	164/110(130)	99	37.2	
Unk	nown	N	10-01	11:20:20	75(S)	124/74(91)	98	36.8	
▼ P2007	7081511	Α	10-01	11:40:34	60(S)	128/80(94)	99	36.2	

Press Trim Knob button on Patient's record then Menu window will pop up. Select View Patient Menu in Menu window.



4.4 View All Patients Record List

Move to Record List.

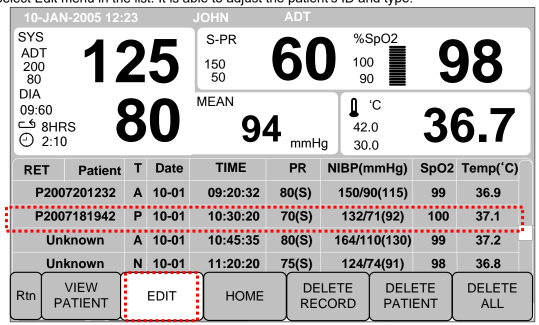
Press Trim Knob Key on Patient's record in the list then Menu window will pop up. Select View All menu in Menu window.

RET Patient	Т	Date	TIME	PR	NIBP	(mmHg)	SpO2	Temp('C)
P2007201232	Α	10-01	09:20:32	80(S)	150	/90(115)	99	36.9
P2007081506	Р	10-01	10:30:20	70(S)	132	2/71(92)	100	37.1
Unknown	Α	10-01	10:45:35	80(S)	164/	110(130)	99	37.2
Unknown	N	10-01	11:20:20	75(S)	124	4/74(91)	98	36.8
P2007081511	Α	10-01	11:40:34	60(S)	128	3/80(94)	99	36.2
Rtn VIEW ALL		EDIT	НОМЕ		DELETE DELET PATIEN			DELETE ALL

4.5 Adjust Record

Move to Record List to adjust the record.

Move to the Record where you want to adjust by turning Trim Knob Key. Select Edit menu in the list. It is able to adjust the patient's ID and type.



Select ID menu window and Adjust 1) Adjust patient's ID. DELETE **VIEW DELETE** DELETE Rtn **EDIT HOME** PATIENT **RECORD PATIENT** ALL **PATIENT ID** Rtn **TYPE** SAVE **CANCEL** PATIENT ID ABCDA_

2)Adjust patient's type. Select Type menu and Adjust

Rtn VIEW PATIENT	EDIT	НОМЕ	DELETE RECORD	DELETE PATIENT	DELETE ALL
Rtn PATIENT ID	ТҮРЕ	SAVE	CANCEL		
Rtn TYPE	> ADT NEO PED				

Alarm status will not be change as a result of excess alarm limit at the moment of measurement even though patient type changed result of alarm limit numeric value change. Select SAVE menu to save changed status.



Select CANCEL button to cancel patient information adjust



4.6 Delete a Record

Move to the Record List.

Move to the Record where you want to adjust by turning Trim Knob Key. Be cautious to delete because deleted record can not be replace.

	RET	Patient	Т	Date	TIME	PR	NIBP(mmHg)	SpO2	Temp('C)	
	P200	7201232	A	10-01	09:20:32	80(S)	150/90(115)	99	36.9	
Г	P200	7081506	Р	10-01	10:30:20	70(S)	132/71(92)	100	37.1	
•	Unl	known	Α	10-01	10:45:35	80(S)	164/110(130)	99	37.2	
	Unl	known	N	10-01	11:20:20	75(S)	124/74(91)	98	36.8	
	P200	7081511	Α	10-01	11:40:34	60(S)	128/80(94)	99	36.2	
	Rtn	VIEW PATIENT		EDIT	НОМЕ		ETE Å DEL CORD PAT	ETE IENT	DELETE ALL	
				*****	*******					
	Rtn	OK	C	ANCEL						

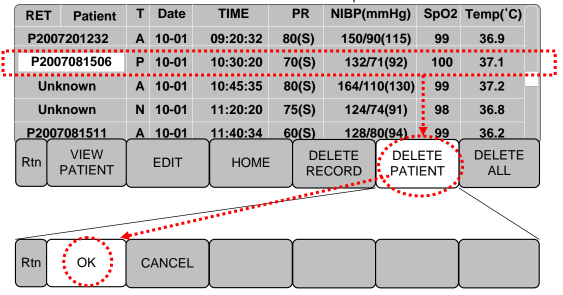
4.7 Delete a Patients Record

Move to record list in order to delete the record.

Move to the Record where you want to adjust by turning Trim Knob Key.

Press Trim Knob Key in the list and menu will pop up then select Delete Patient button.

Be cautious to delete because deleted record can not be replace.



4.8 Delete All Patients Record

Enter the record list to delete all the record.

Select Trim Knob Button in the patient's record then select Delete All.

Be cautious to delete because deleted record can not be replace.

	RET	Patient	Т	Date	TIME		PR	NIBP(m	nmHg)	SpO2	Temp('C)	
	P20	07201232	A_	10-01	09:20:32	80	0(\$)	150/9	Q(115).	99	36.9	•
	P20	07081506	Р	10-01	10:30:20	7	0(S)	132/7	71(92)	100	37.1	
•••	Ur	known	Α	10-01	10:45:35	8	0(S)	164/11	0(130)	99	37.2	
	Ur	known	N	10-01	11:20:20	7	5(S)	124/7	74(91)	98	36.8	
	P200	07081511	<u>A</u>	10-01	11:40:34	6	0(S)	128/8	30(94)	99	36.2	
	Rtn	VIEW PATIENT		EDIT	НОМЕ			CORD	DEL PATI		DELETE ALL	
					······································				******		*****	• •
	Rtn	OK		ANCEL								

5. SETUP

5.1 SETUP

5.2 DISPLAY

5.3 SAVE MODE

5.4 USER SERVICE

5.5 SYSTEM

5.6 KEY SOUND

5.7 MAKER SERVICE

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5.1 SETUP

Select setup Icon in the menu icon.



DISPLAY: A menu to set up screen

SAVE MODE: A menu to setup the record saving mode (AUTO , MANUAL)

USER SERVICE: To setup information of equipment SYSTEM: To set up connection to external computer

KEY SOUND: Set up ON/OFF of Key sound.

MAKER SERVICE: Using by manufacturer to set up and reform of the product.

Rtn	DSPLAY	SAVE MODE: MANUAL	USER SERVICE	SYSTEM	KEY SOUND: ON	MAKER SERVICE
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5.2. DISPLAY

Rtn	SWEEP SPEED:	SET	SET	DEMO:	
	25mm/s	DATE	TIME	OFF	

1. SWEEP SPEED

Set up print speed of amount of oxygen in the blood (SPO2) wave pattern.

Rtn	SWEEP SPEED: 25mm/s	SET DATE	SET TIME	DEMO: OFF	
Rtn	SWEEP SPEED: 25mm/s	> 6.25mm/s 12.5mm/s			

2. SET DATE

Setup and adjust the date.

SWEEP

Rtn	SWEEP SPEED: 25mm/s	SET DATE	SET TIME	DEMO: OFF		
Rtn	SET DATE		2	2 - DEC - 200	7	

3. SET TIME

Setup and adjust the time

Rtn	SWEEP SPEED: 25mm/s	SET DATE	SET TIME	DEMO: OFF	
Rtn	SET TIME			11:25:06	

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4. DEMO

Setup the movement to demo/action mode.

Rtn	SWEEP SPEED: 25mm/s	SET DATE	SET TIME	DEMO: OFF	
Rtn	SWEEP SPEED: 25mm/s	SET DATE	SET TIME	DEMO: ON	

5.3 SAVE MODE

Set up menu for record saving mode.

Rtn DISPLAY	SAVE MODE : AUTO	USER SERVICE	SYSTEM	KEY SOUND: OFF	MAKER SERVICE
Rtn DISPLAY	SAVE MODE: MANUAL	USER SERVICE	SYSTEM	KEY SOUND: OFF	MAKER SERVICE

AUTO mode is to save all of measured data with a same person's ID and TYPE. MANUAL mode is initializing ID whenever saving is activated.

5.4 USER SERVICE

Setup for information of the equipment

1. BED NUMBER

Setup the number for the bed which connected to the equipment.

It is able to set up $0\sim9$ and $A\sim Z$.

Rtn	SET BED NUMBER : A01	SET UNIT NAME	DISPLAY MODE : SPOT		
Rtn	SET BED NUMBER :	A0 1			

2. UNIT NAME

Set up UNIT name for connected hospital with equipment.

Rtn	SET BED NUMBER : A01	SET UNIT NAME	DISPLAY MODE : SPOT	
Rtn	SET UNIT NAME			NICU

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5.5 SYSTEM

Setup for connect to outside computer.

setup for connect to outside computer.							
10-JAN-2007 12:23		JOHN	ADT				
SYS mmHg Al	5 DT 200 80	S-PR	150 50	%SpO2	100 90		
	SYSTEM INFO SET						
RETURN			CO	NTENTS			
MAIN VER CENTRAL				1.10.BHCDDC/ OFF	A		
HOST IP			19	92 . 168 . 030 . 1	. 030 . 100		
DEVICE IP			19	92 . 168 . 030 . 1	101		
SUBNET		255 . 255 . 255 . 000					
GATEWAY			192 . 168 . 030 . 001				
MAC ADDR			00 :	02 : BD : 80 : 0	0 : 00		

5.6 KEY SOUND

Setup ON/OFF of key sound.

Rtn	DISPLAY	SAVE MODE : AUTO	USER SERVICE	SYSTEM	KEY SOUND : OFF	MAKER SERVICE
Rtn	DISPLAY	SAVE MODE : AUTO	USER SERVICE	SYSTEM	KEY SOUND : ON	MAKER SERVICE

5.7 MAKER SERVICE

A menu used by the manufacturer of the product.

Rtn	DISPLAY	SAVE MODE : AUTO	USER SERVICE	SYSTEM	KEY SOUND : ON	MAKER SERVICE	
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6. NIBP

6.1 Outline

NIBP Connector Location and Cuff

6.2 NIBP Data Window

6.3 NIBP Setup

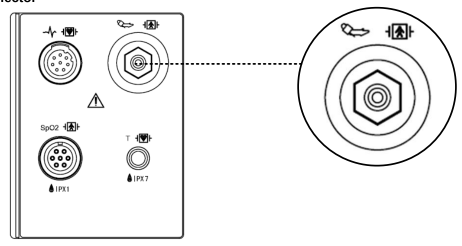
ALARM LIMIT CUFF SIZE
NIBP STAT
INFLASTION SET
UNIT SELECT
INTERVAL

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6.1 Outline

The function is to measure minimum, maximum, and average blood pressure by using oscillometric method.

NIBP Connector Location and Cuff NIBP Connector



ADULT NIBP CUFF

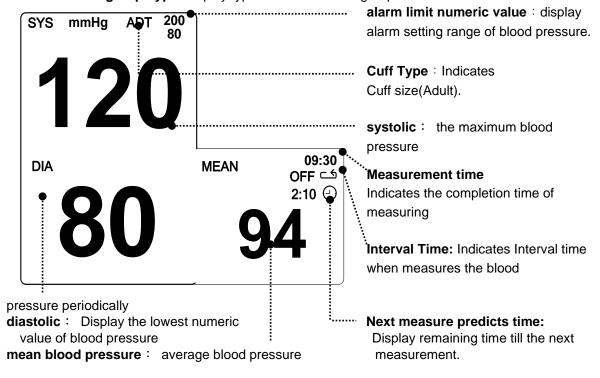


Note

As the value of NIBP can vary according to the age and sex of a patient, the user needs to set up right data in Parameter Menu before measurement.

6.2 NIBP Data Window

Measurement group Type: display type of measurement group



6.3 NIBP Setup

ALARM LIMIT: A menu to setup alarm range CUFF SIZE: A menu to select Cuff size

STAT: Start 5 minutes of continuous, sequential NIBP measurements.

INFLATION: A menu to setup INFLATION UNIT: A menu to setup blood pressure unit

INTERVAL : A menu to setup interval for blood pressure measurement

Rtn	ALARM LIMIT	CUFF SIZE: ADT	NIBP STAT: OFF	INFLATION SET: 170mmHg	UNIT SELECT: mmHq	INTERVAL: OFF
		וטא	OII	[170mming	I III III II	

ALARM LIMIT

Numeric value of Systolic, Diastolic, and mean pressure is 10 ~ 350mmHg.

10-JAN-2005 12:23	JOHN	ADT				
SYS mmHg AD	80	-PR 6 (150 50	%SpO2	98	100 90
		NIBP ALARM	LIMIT			
RETURN	UNIT		LOW		HIGH	
NIBP-S	mmHg		80		200	
NIBP-M	mmHg		40		140	
NIBP-D	mmHg		20		120	

CUFF SIZE

It is able to choose adult, baby, and children's cuff.

Rtn	ALARM LIMIT	CUFF SIZE: ADT	NIBP STAT: OFF	INFLATION SET: 170mmHg	UNIT SELECT: mmHg	INTERVAL: OFF
Rtn	CUFF SIZE: ADT			> ADT PED NEO		

NIBP STAT

Start 5 minutes of continuous, sequential NIBP measurements.

Rtn ALARM LIMIT	CUFF SIZE: ADT	NIBP STAT: OFF	INFLATION SET: 170mmHg	UNIT SELECT: mmHg	INTERVAL: OFF
-----------------	----------------------	----------------------	------------------------------	-------------------------	------------------

INFLATION SET

The function for setup of pressure at the beginning

Set numeric value is 80, 100, 120, 140, 160, 180, 200, 220, and 240.

Rtn	ALARM LIMIT	CUFF SIZE: ADT	NIBP STAT: OFF	INFLATION SET: 170mmHg	UNIT SELECT: mmHg	INTERVAL: OFF
Rtn	ALARM LIMIT	CUFF SIZE: ADT	NIBP STAT: OFF	INFLATION SET: 170mmHg	UNIT SELECT: mmHg	INTERVAL: OFF

UNIT SELECT

The function is to setup blood pressure measurement display unit.

Set unit is mmHg, kPa

Rtn	ALARM LIMIT	CUFF SIZE: ADT	NIBP STAT: OFF	INFLATION SET: 170mmHg	UNIT SELECT: mmHg	INTERVAL: OFF
Rtn	ALARM LIMIT	CUFF SIZE: ADT	NIBP STAT: OFF	INFLATION: 170mmHg	UNIT SELECT: kPa	INTERVAL: OFF

INTERVAL

The function is to setup the interval to measure blood pressure automatically Set numeric value is1min, 2, 3, 4, 5, 10, 15, 20, 30, 1hour, 2, 4, and 8.

Rtn	ALARM LIMIT	CUFF SIZE: ADT	NIBP STAT: OFF	INFLATIO SET: 170mmH	SELECT:	INTERVAL: OFF
Rtn	INTERVAL: OFF	> OFF 1MIN. 2MIN.	4MIN.	10MIN. 30N 15MIN. 1I 20MIN. 2I	. 0	

Warning

Periodically check patient limb circulation distal to the cuff. Check frequently when using auto NBP in 1 and 2 minute intervals. Intervals below 10 minutes are not recommended for extended periods of time.

Warning

Pay attention to not to block connecting hose when you put cuff on patient.

NIBP Status Messages

Below is a list of system status alarm messages which may be displayed in the NIBP parameter window during monitoring.

Status Message	Monitor Response	Solution
OVER PRESSURE	System status alarm. Auto mode will shut off after ONE message.	Remove cuff and contact service.
INFLATION FAIL. CHECK CUFF	System status alarm.	Check cuff, connections, and tubing.
DEFLATION FAIL. CHECK CUFF	System status alarm. Auto mode will shut off after ONE message.	Remove cuff and contact service.
PULSE TOO WEAK	System status alarm. Auto mode will shut off after ONE message.	Check patient and cuff placement.
EXCESSIVE MOTION	System status alarm. Auto mode will shut off after ONE message.	Possible excessive patient movement. Check patient.
MEASUREMENT ERROR	ystem status alarm. Auto mode will shut off after ONE message.	Possible excessive patient movement or arrhythmia condition. Check patient.

Erroneous NIBP measurement

- Check for proper cuff size
 - 3. Too small a cuff can give an erroneously high value.
 - 4. Too large a cuff can give an erroneously low value.
- Check for residual air left in the cuff from a previous measurement.
- Make sure cuff is not too tight or too loose.
- Make sure cuff and heart are at same level, otherwise hydrostatic pressure will offset the NIBP value.
- Minimize patient movement during measurement.
- Check for leak in cuff or tubing.
- Patient may have a weak pulse.

7. SpO₂

7.1 Outline

SpO2 Connector Location and Measuring Cable

7.2 SpO2 Data Window

7.3 SpO2 Setup ALARM LIMIT SWEEP SPEED RATE VOLUME ALARM LEVEL

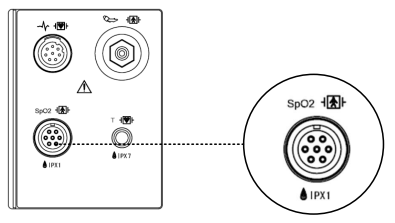
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7.1 Outline

SPO2 monitoring is a noninvasive technique used to measure the amount of oxygenated hemoglobin and pulse rate by measuring the absorption of selected wavelengths of light. The light generated in the probe passes through the tissue and is converted into an electrical signal by the photodetector in the probe. The monitor processes the electrical signal and displays on the screen a waveform and digital values for SpO2 and pulse rate. It detects SpO2 in the way of transmitting the red and infrared rays into the capillary vessel to take the pulsation. Also perform the alarm function according to the setting value.

SpO2 Connector Location and Measuring Cable

SpO₂ Connector



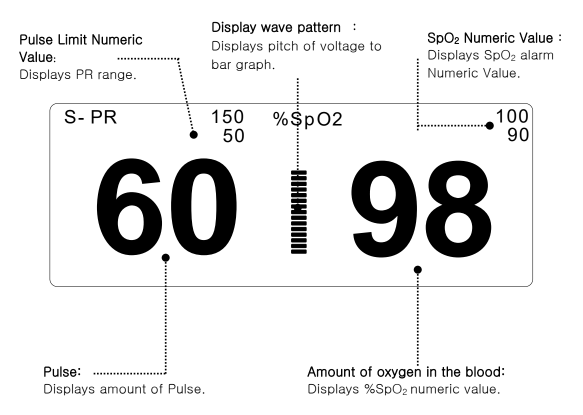
SpO₂ Measuring Cable



Note

The signal input is a high-insulation port and it is defibrillator proof († † †)
The insulated input ensures patient safety and protects the device during defibrillation and electrosurgery.

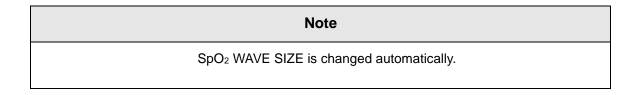
7.2 SpO₂ Data Window



The current SPO2 value and the derived pulse rate (RATE) are displayed. The block sets indicate the strength of the signal (twenty block bars indicate the strongest signal). The SPO2 measurements are averaged over a 6-second period of time.

The monitor display is updated every second.

The SPO2 monitoring features are found in the SPO2 menu. These features include alarm limit adjustment, display of RATE, and RATE volume.



7.3 SpO₂ Setup

ALARM LIMIT: A menu to set SpO₂ limit.

SWEEP SPEED: A menu to set speed of WAVE display.

RATE VOLUME: A menu to set Rate Volume.

ALARM LEVEL: A menu to set SpO2 ALARM LEVEL.

Rtn ALARM SWEEP SPEED: 6.25mm/s OFF ALARM LEVEL

ALARM LIMIT

ALAMRM Numeric Value of %SpO2 is $40 \sim 100$. Pulse numeric Value of SpO2 is $20 \sim 300$ BPM.

10-JAN-2005 12:2	3 J	OHN	ADT		
SYS mmHg AD	T 200 80	S-PR	150 50	SpO2 %	100 90
40					0
12			50	9	ਨ
		SPO2 AL	ARM LIMIT		
RETURN	UNIT	7	LOW	F	IIGH
SpO2-%	%		90		100
SPO2-R	ВРМ		50		150

SWEEP SPEED

Adjust WAVE DISPLAY speed setup as below.

Numeric value is 6.25, 12.5, 25, 50mm/s

Rtn	ALARM LIMIT	SWEEP SPEED 6.25mm/s	RATE VOLUME OFF	ALARM LEVEL		
Rtn	SWEEP SPEED		> 6.25mr 12.5mr		nm/s nm/s	

RATE VOLUME

Rate Volume can be adjusted from off and 10% to 100%.

Rtn	ALARM LIMIT	SWEEP SPEED: 6.25mm/s	RATE VOLUME: OFF	ALARM LEVEL			
Rtn	RATE VOLUME:		> OFF 10 %		60% 70%	90% 100%	
	OFF	Į	20 %	50%	80%		J

ALARM LEVEL

Set the order of priority in each alarm.

Rtn	ALARM LIMIT	SWEEP SPEED 6.25mm/s	RATE VOLUME OFF	■	ALARM LEVEL			
	PARAMETER ALARM LEVELS							
	RETURN				Al	LARM LEVEL		
		PR SPO2-%		MEDIUM LOW				
	Р	ROBE OFF			М	ESSAGE		
	CHECK PROBE				M	ESSAGE		
	LC	OST PULSE				LOW		
	PO	OR SIGNAL				LOW		
	A	ARTIFACT				LOW	J	

PROBE OFF Condition

When using a reusable finger probe, there is a system alarm to alert you when the probe is off the Monitor. The monitor defaults this "PROBE OFF" condition as a System Warning alarm. You can, however, set it as a System ALARM LEVEL in Monitor Defaults.

SPO2 Messages

Below is a list of system status alarm messages which may be displayed in the SPO2 parameter window during monitoring.

CHECK PROBE

Reusable finger probe is off the patient. Check the probe. *The factory default for this alarm is MESSAGE ALARM.*

PULSE SEARCH

Detection by the monitor of a repeatable pulse has ceased. Check the patient and the probe site.

POOR SIGNAL

The SPO2 signal is too low. No SPO2 data is displayed. This can be due to a low patient pulse, patient motion, or some other interference. Check the patient and the probe.

LOST SIGNAL

SPO2 data continues to be displayed, but the quality of the signal is questionable. Check the patient and the probe.

ARTIFACT

The SPO2 signal is patient's motion artifact and noise

No SpO2 data is displayed. One of the following conditions is indicated:

- defective or damaged probe,
- defective or damaged cable
- probe is off the patient, or
- Detection of a repeatable pulse has ceased.
- Check the probe and cable: reposition or replace as needed.

8. TEMPERATURE

8.1 Outline

Temperature Connector and Measuring Cable

8.2 Temperature Data Window

8.3 Temperature Setup

ALARM LIMIT UNIT SELECT PROBE SITE

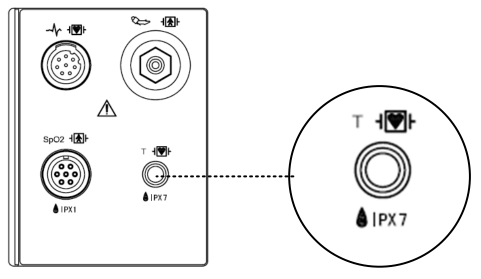
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8.1 Outline

Adjust electric signal procedure in change of resistance ingredient followed by temperature change then it shows numeric value through signal procedure.

Temperature Connector and Measuring Cable

Temperature Connector



Temperature Measuring Cable

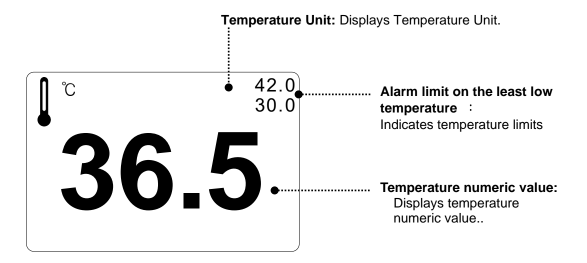


Note

Temperature probe is correctly positioned and fixed to do not disconnect on the patient. Temperature cable is attached to the monitor.

The TEMP cable connector is a high-insulation port and it is defibrillator-proof(1) .

8.2 Temperature Data Window



Note

For an accuracy measurement for human body, it takes 3 minute interval to measure.

8.3 Temperature Setup

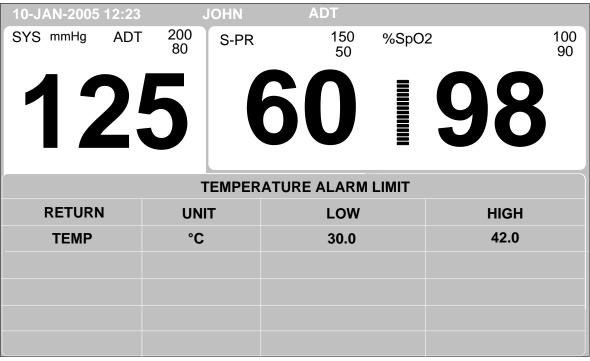
ALARM LIMIT: Sets up temperature limit. UNIT: Sets up temperature measurement unit.

PROBE SITE: Displays temperature measurement region.

Rtn ALARM SITE: ORAL	UNIT SELECT: °C		
----------------------	-----------------------	--	--

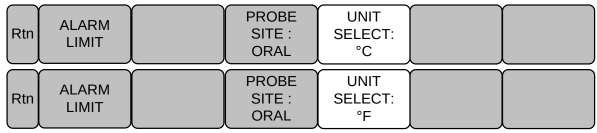
ALARM LIMIT

Numeric value is 0°C ~ 50.0°C.



UNIT SELECT

It is able to select °C and °F unit.



PROBE SITE (Measurement Position)

Set up to display temperature measurement region.

Measurement regions are ORAL, AUXILLARY, and RECTAL.

PROBE UNIT ALARM Rtn SITE: SELECT: LIMIT **ORAL** °C **PROBE** > ORAL Rtn **AXILLARY** SITE: **ORAL RECTAL**

Check list

- 4. The temperature probe(YSI 400 series) is correctly positioned on the patient.
- 5. Temperature cable is attached to the monitor.
- 6. Temperature setup is adjusted, if necessary. Follow detailed procedures within this chapter.

TEMP Message

If you experience some problems with temperature monitoring, one of the following messages may be displayed in the TEMP parameter window.

- LEAD FAULT: Probe is not properly connected. Check the probe.
- No temperature value will be displayed . Service on the monitor is required.

Warning

To measure the peripheral temperature, attach the probe to the ankle or palm.

If the patient sweats heavily or moves violently, fasten the pad with surgical tape.

NOTE

When the measuring site is exposed directly to air, the temperature may be lower than normal. It take about 20 to 30 minutes to reach the equilibrium temperature after attaching the sensor.

9. PRINT

9.1 Print

Print and Heat Sensitivity Paper Function and Setup Menu

9.2 Paper Change

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9.1 Print

Print and Heat Sensitivity Paper

A printer used to print data onto thermal paper, this product is offered as an option, Size of the thermal paper roll: width 58mm x diameter 38 mm papers can be used. Any thermal paper of same size can be used for the printer.

Side view of printer



Function and Setup Menu

	PRINT	RECORD	WAVE		
Rtn	SPEED:	NUMBER:	TIME:		
	25mm/S	RECENT	20		

- 1. Able to ON/OFF the PRINT Key in constant printing.
- 2. Able to Set up the print speed to 25, 50 mm/s.

Rtn	PRINT SPEED: 25mm/S	RECORD NUMBER: RECENT	WAVE TIME: 20		
Rtn	PRINT SPEED: 50mm/S	RECORD NUMBER: RECENT	WAVE TIME: 20		

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3. RECORD NUMBER

Able to setup print from top RECORD to RECORD NUMBER numeric value in current list while activate PRINT in RECORD LIST window.

Rtn	PRINT SPEED: 25mm/S	RECORD NUMBER: RECENT	WAVE TIME: 20		
	RECORD		RECENT	30	
Rtn	NUMBER:		> 10	50	
	RECENT	Į	20	ALL	

4. WAVE TIME

When printing in WAVEFORM VIEW

Able to setup print from current time till WAVE TIME while activate PRINT in the WAVEFORM VIEW.

Rtn	PRINT SPEED: 25mm/S	RECORD NUMBER: RECENT	WAVE TIME: 20	
Rtn	PRINT SPEED: 25mm/S	RECORD NUMBER: RECENT	WAVE TIME: CONTINUE	

5. Set up ALARM PRINT in additional menu, and then print automatically when alarm occurs.



Rtn	ALARM LIMIT	ALARM PRINT: OFF	ALARM VOLUME: 50%	ALARM LEVEL	NURSE CALL: OFF	ALARM SOUND
Rtn	ALARM LIMIT	ALARM PRINT: ON	ALARM VOLUME: 50%	ALARM LEVEL	NURSE CALL: OFF	ALARM SOUND

If there is no print sheet, no paper icon of



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9.2 Paper Change

1

Open the window of the printer.



2

Insert the paper roll offered with the product into the printing unit. Place the roll in a proper way so that the printed paper can roll out upwards.



3

Press the printer window until it is properly shut. Inaccurate shutting may cause failure in printing.

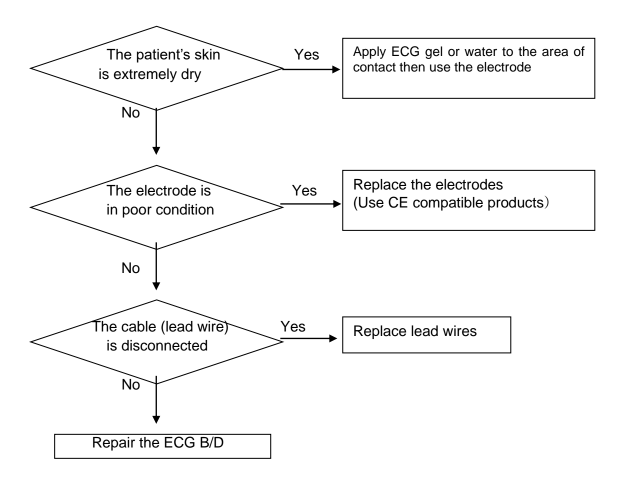


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10. TROUBLE SHOOTING

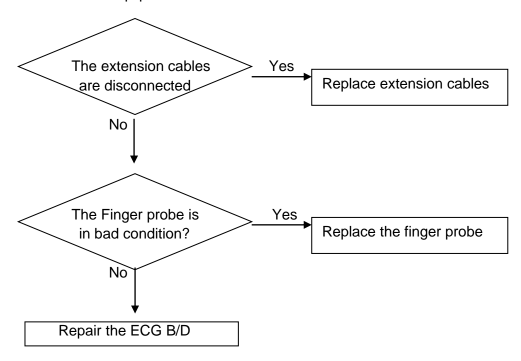
1. Noise in ECG

- Gel is dry
- Electrodes does not stick well to skin

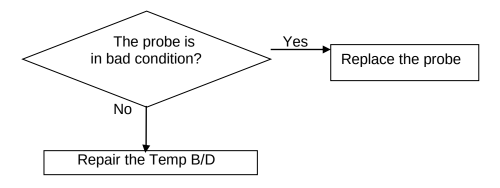


2. SpO₂ malfunction

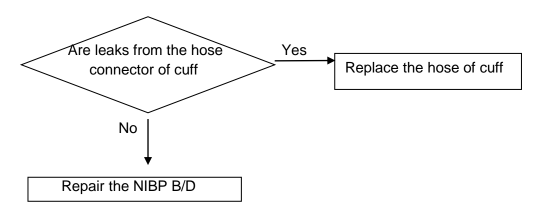
Connectors of the equipments are in bad condition?



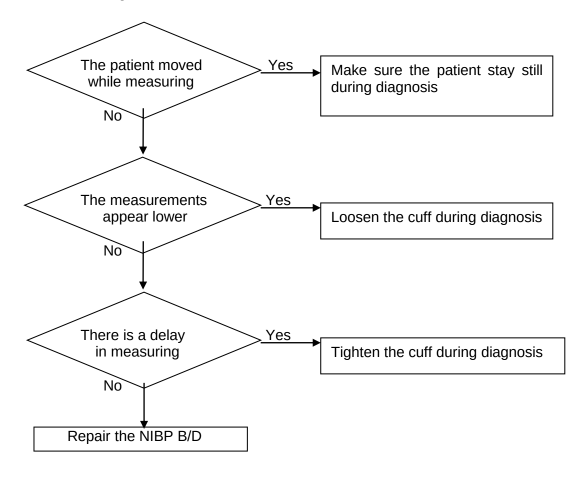
3. Temp malfunction



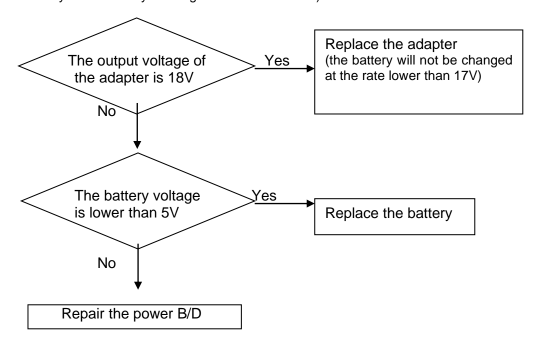
4. NIBP malfunction



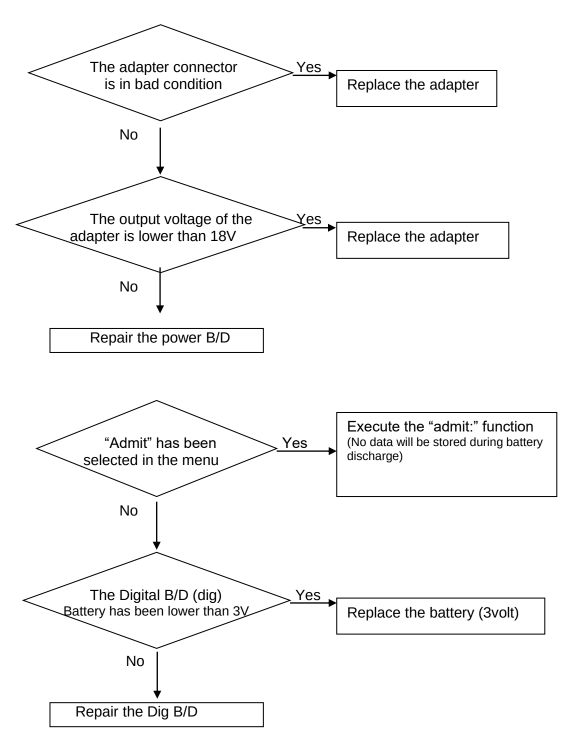
5. Abnormality in NIBP measurements



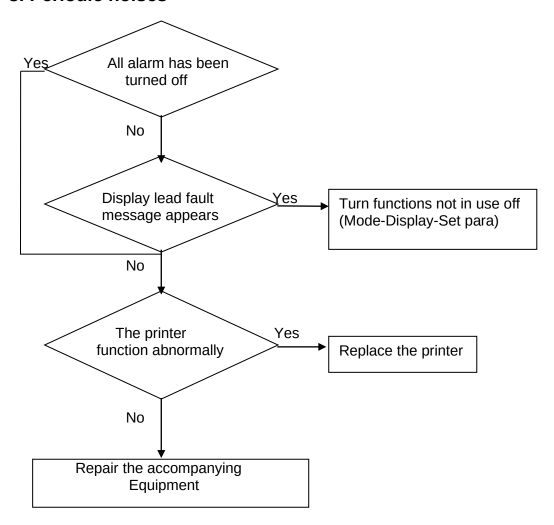
6. Failure in battery recharge (the battery does not fully recharge in 6 hours or more)



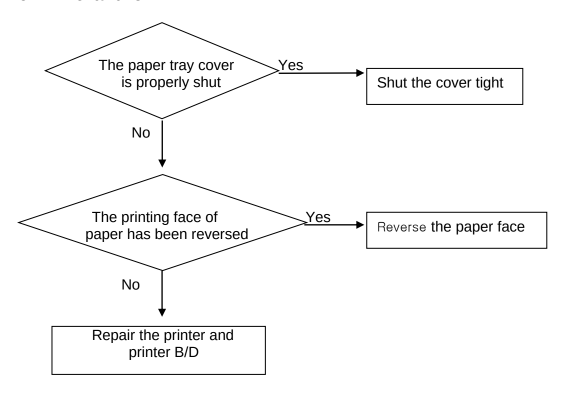
7. Power failure



8. Periodic noises



9. Print failure



SPECIFICATION

Ease of use

Customization

Special Features

Monitor Environmental Specifications

Power adaptor

Monitor Performance Specifications

Graphical and Tabular Trends

SpO2 Performance Specifications

Respirations Performance Specifications

NIBP Performance Specifications

ECG Performance Specifications

Temperature Unit Performance Specifications

Accessories included

OPTION

Ease of use

- · Battery operation
- · Attached printer
- · Table and graphic trend
- · Nellcor SpO₂ sensor compatible (OxiMax sensor exclusion)

Intended use

- The monitors are intended to be used for monitoring and recording of, and to generate alarms for, multiple physiological parameters of adults, pediatrics, and neonates.
- The monitors are intended for use by trained healthcare professionals in a hospital environment.
- The BM3 monitors are additionally intended for use in transport situations within hospital environments.
- The monitors are only for use on one patient at a time. They are not intended for home use. Not therapeutic devices.
- The monitors are for prescription use only.
- The BM3 Patient Monior is not intended for use during MRI.
- The BM3 Patient Monior monitors and displays ECG (including arrhythmia and ST segment analysis), heart/pulse rate, oscillometric non-invasive blood pressure(systolic, diastolic and mean arterial pressure), end-tidal carbon dioxide, respiration rate, temperature with a electronic thermometer for continual monitoring
 - Esophageal/Nasopharyngeal/Tympanic/Rectal/Bladder/Axillary/Skin/Airway/Room/Myocardial/C ore/Surface temperature, and functional oxygen saturation (SpO₂) and pulse rate via continuous monitoring, including monitoring during conditions of clinical patient motion or low perfusion.
- The ECG measurement is intended to be used for diagnostic recording of rhythm and detailed morphology of complex cardiac complexes.
- ST segment monitoring is intended for use with adult patients only and is not clinically validated for use with neonatal and pediatric patients.
- The gas measurement is restricted to neonatal patients only.

Indication for use

Monitoring, recording and/or alarming of patients including with arrhythmias in ICU, Post-OP and others. The target populations are adult, pediatric and neonate with the exception of:

· Arrhythmia detection and ST segment analysis, for which the target populations are adult and pediatric only.

Additional Function

· LAN Connection

Monitor Environmental Specifications

Operating Temperature : 10°C to 40°C (50°F to 104°F)

- Storage Temperature : - 10°C to 60°C (14°F to 140°F)

· Humidity: 20% to 95% RH

- Operating Attitude: 70(700) to 106Kpa(1060mbar)

Power

- AC 100-240V (50/60Hz) 1.2A
- Adapter 18 V, 2.5 A (MW160KA1803)

Monitor Performance Specifications

- Screen: 7" TFT LCD (800×480)
- · Indicators
 - Up to 3 wave patterns
 - 3 levels of alarm sound
 - Visual alarmPulse sound
 - Battery status
 - LED external power supply LED
- · Interfaces
 - Generating power for LAN, Wireless LAN: 5.0V max 0.9A
- Battery
 - Li-ion battery
 - Battery status display
 - Operating time: 1.5hours(with fully charged Battery)
- Thermal Printer : internal printer
 Speed : 25, 50 mm/sec
 Paper width : 58 mm

Graphical and Tabular Trends

- · Table Trend
 - Memory Storage : 128 hours
 - Data Interval: 1 minute
 - Display Interval: 1MIN, 5, 15, 30, 1HR
- · Graphical Trend
 - Display Period: 30MINS, 60, 90, 3HRS, 6, 12

ECG capacity

· Lead : 3,5

- Heart rate range: Adult-30 to 300 bpm, Nedonate/Pediatric-30 to 350bpm (accuracy: ±3 bpm)

Bandwidth(monitoring mode) : 0.5 Hz to 40 Hz
Display Sweep Speed : 2 5mm / sec
ECG size (Sensitivity) : 0.5, 1, 2, 4 mV/cm

· Lead-off Detection with display indicator

· Pace maker Detection Mode

- Differential Input Impedance : $> 5 M\Omega$

Common Mode Rejection Ratio : > 90 dB at 50 or 60 Hz

DC Input Range : ±5 mV
 Defibrillator Discharge : < 4s
 Defibrillation Artifact Recovery Time : < 8s

SpO₂ capacity

- Saturation Range : 0% to 100% oxygen proportion

Pulse Rate Range : 30 to 254 bpm

- SpO₂ accuracy: 70% to 100% ±2 digits, 0% to 69% unspecified

pulse accuracy : ±2 bpm

· Sensor Red 660nm, 2mW (typical)

Infrared 880nm, 2-2.4mW (typical)

 Minimum Signal: 0.05% modulation (Low perfusion level performance and Amplitude limitation validation using FLUKE Index 2 Oximetry Simulator)

Respiration Performance Specifications

Range: 5 to 120 breaths/min
Accuracy: ±3 breaths/min
Display Sweep Speeds: 25mm/sec

NIBP capacity

· Technique : Oscillometric

· Measurement mode:

- Manual : Single Measurement

- STAT: Start 5 minutes of continuous

- Auto: automatic Intervals of 1MIN., 2, 3, 4, 5, 10, 15, 20, 30, 1Hour, 2, 4, 8

Pressure Display : 0 to 300 mmHg (Accuracy ±3mmHg)

Measurement Range:

- Systolic: ADULT 40 – 260mmHg

PEDIATRIC 40 – 230mmHg NEONATE 40 – 130mmHg

- MAP(Mean Aterial Pressure): ADULT 26 - 220mmHg

PEDIATRIC 26 – 183mmHg NEONATE 26 – 110mmHg

- Diastolic : ADULT 20 – 200mmHg

PEDIATRIC 20 – 160mmHg NEONATE 20 – 100mmHg

- Pulse Rate: ADULT 30 – 220BPM

PEDIATRIC 30 – 220BPM NEONATE 30 – 220BPM

Temperature Unit Performance Specifications

• Range : 0°C to 50°C (32°F to 122°F)

• Accuracy : 25°C to 50°C ± 0.1°C, 0°C to 24°C±0.2°C

· Sensor: YSI 400 Series compatibility

Accessories Included:

· 3Lead patient cable	1 EA
· Electrodes	10 EA
· NIBP extension hose, 3m long	1 EA
- Adult cuff(27.5~36.5cm)	1 EA
- SpO ₂ extension cable 2m	1 EA
Reusable Adult SpO2 Probe	1 EA
DC adapter, 18VDC, 2.8A (MW160 Made in Bridge Power Co., Ltd.)	1 EA

Option

- · Temperature sensor (skin)
- 5 lead patient cable
- · Thermal printer (58mm) and Thermal paper roll
- Reusable Multi-site "Y" SpO2 Sensor
 Disposable Adult SpO2 Sensor (D-MDNA)
 1 EA

Abbreviations and Symbols

Abbreviations and symbols which you may encounter while reading this manual or using the monitor are listed below with their meanings.

Abbreviations

		Α
A AC ADT ARRYTHM ASYS Auto, AUTO AUX aVF aVL	amps alternating current adult arrhythmia asystole automatic Auxiliary left foot augmented lead left arm augmented lead	
aVR	right arm augmented lead	В
BPM	beats per minute	
C CAL cm, CM	Celsius calibration centimeter	С
D DC DEFIB, Defib DIA	diastolic direct current defibrillator diastolic	D
ECG EMC EMI ESU	electrocardiograph electromagnetic compatibility electromagnetic interference electrosurgical cautery unit	E
F	Fahrenheit	F
g	gram	G
HR Hz	heart rate, hour hertz	Н
ICU Inc	intensive care unit incorporated	I
kg, KG	kilogram	K

kPa kilopascal L liter, left LA left arm, left atrial **LBS** pounds LCD liquid crystal display LED light emitting diode LL left leg M M mean, minute meter MIN, min minute millimeters MM, mm MM/S millimeters per second MMHG, mmHg millimeters of mercury m۷ millivolt N **NIBP** noninvasive blood pressure NEO, Neo neonatal 0 OR operating room Р PED pediatric **PVC** premature ventricular complex **QRS** interval of ventricular depolarization R RA right arm, right atrial **RESP** respiration RL right leg RR respiration rate S S systolic sec second SpO2 arterial oxygen saturation from pulse oximetry SYNC, Sync synchronization SYS systolic T Temp, TEMP temperature U ٧ precordial lead volt V-Fib, VFIB ventricular fibrillation VTAC ventricular tachycardia

W

X

X multiplier when used with a number (2X)

Symbols

and
degree(s)
greater than
less than
minus
number
percent
plus or minus

PRODUCT WARRANTY

Product Name	Patient Monitor
Model Name	ВМЗ
Approval Number	
Approval Date	
Serial Number	
Warranty Period	1 year from date of purchase
Date of Purchase	
Customer Section	Hospital Name : Address : Name : Phone :
Sales Agency	
Manufacturer	

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^{*} Thank you for purchasing BM3.
* The product is manufactured and passed through strict quality control and through inspection.

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BIONET CO., LTD.

Product Name: BM3

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