Patient Monitor **BP-S510** Operation Manual

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- Please contact your local distributor about the warranty period.
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 - Damage sustained to the Colin Medical Technology product(s) caused by product(s) from another company excluding products delivered by Colin Medical Technology.
 - Damage caused by mishandling and/or misuse is the responsibility of the user.
 - Maintenance and repairs utilizing maintenance components that are not specified by Colin Medical Technology.
 - Device modifications or use of accessories not recommended by Colin Medical Technology.
 - Damage caused by accidents or natural disasters (earthquakes, flooding, etc.).
 - Damage resulting from usage where caution statements and operating instructions shown in this manual have not been followed.
 - Damage due to neglect of specified maintenance checks.
- This warranty only covers the hardware of the BP-S510. The warranty does not cover the following selections:
 - Whatever damage or loss results from the attachment of accessories or their operation.
 - In the event of a defect in the product, contact our sales outlet or EU representative as noted on the back cover.
- The BP-S510 conforms to the EMC standard IEC60601-1-2.
 Note that mobile phones should not be used in the vicinity of the BP-S510.

Note, however, any device not complying to the EMC standard that is used with the BP-S510 renders the BP-S510 as non-compliable to the EMC standard.

Trademark

Product brand names shown in this manual are likely to be the trademark or registered trademark of the company concerned.

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SAFETY INFORMATION

General Safety Information

This section contains important safety information related to general use of the BP-S510 multi-parameter patient monitor. Other important safety information appears throughout the manual. The BP-S510 will be referred to as the monitor throughout this manual.

Important! Before use, carefully read this manual, accessory directions for use, all precautionary information and specifications.

Warning

Warnings are identified by the WARNING symbol shown above.

Warnings alert you to potential serious outcomes (death, injury, or adverse events) to the patient or user.

WARNING: Do not take into or use the monitor in locations where highly combustible anesthetics or flammable gases are used or in high-pressure oxygen rooms or inside oxygen tents, as this may cause a flammable explosion.
WARNING: When using the monitor with a commercial electric power source, use the monitor with an electric power wall socket with a grounding wire for medical use. Not doing so could cause electric shock.
WARNING: Do not connect grounding wire to gas pipes. This could cause fire.
WARNING: Only doctors and officially certified personnel should use this monitor. Do not allow patients to touch this monitor. Allowing patients to touch this monitor could cause accidents.
WARNING: This monitor cannot be used when MRI is in progress. If MRI is in use, keep patient attachments away from patients to prevent accidents.
WARNING: The monitor conforms to the requirements of the EMC standard (IEC60601-1-2), and may therefore be used simultaneously with pacemakers and other electrical simulators. It should, however, be noted that the BP-S510 may be affected by electrical scalpels and microwave therapeutic apparatus. Please check operation of the monitor during and after use of such equipment.
WARNING: Do not take mobile phones or transceivers into a room where this monitor is installed, as such devices may cause accidents.
WARNING: In order to avoid accidents, do not use any unauthorized accessories or options.
WARNING: Thoroughly read the instruction manuals supplied with accessories and options to ensure correct use. This instruction manual does not carry the caution selections for such equipment.
WARNING: Do not open cover or disassemble this monitor. Doing so could cause electric shock or fire. It is prohibited by law to modify the monitor without authorization.
WARNING: Do not use power source other than the specified voltage, (100-240V~50/60Hz) as this may cause fire or electric shock.

WARNING: Pre-use inspection and preventive maintenance must be performed for safe use.
WARNING: The monitor may be used with electrical surgical equipment.
Follow the instruction manuals for medical instruments – notably electrosurgical
and diathermy instruments - when used, as their high - frequency energy units
 may cause burns to patients via attachments.
WARNING: This monitor is protected against the discharge of a defibrillator. But
do not touch the monitor when a defibrillator is being discharged (electrified), as
warning: The following cautions apply when connecting he monitor with other
1. Ensure that the connected equipment is in accordance with the IEC60601-1 or
IEC safety standards, so that the system complies with IEC60601-1.
2. Employ additional protective measures (e.g. additional protective earthing) as
 necessary.
WARNING: Do not connect devices that do not meet medical safety standards
(such as commercial PCs), as they may cause electric shock. This monitor meets
monitor cannot be connected to a device that would give a combined total of
leakage current beyond the restricted level.
WARNING: Do not place anything on top of this monitor. If something is spilled
over the monitor or gets into it, such spillage may cause fire or electric shock. If
fluid spills on the monitor accidentally, disconnect power cord, wipe dry
 immediately, and have the monitor serviced to make sure that no hazard exists.
WARNING: Do not place heavy objects on the power cord, as doing so may cause
the power cord from the wall socket to prevent electric shock
 WARNING: When the following occur, turn the newer OFF immediately and unplug
the power cord from the wall socket. Continued use in such situations may cause
fire or electric shock.
 There is smoke or a strange odor leaking out of the device.
 The devices has been dropped or impacted by an object.
Liquid or foreign matter gets inside the device.
Device failure has occurred.
Also, when any of the above occurs, promptly do the following:
1. Check to see that the power cord has been unplugged from the wall socket.
2. Place an "Out of Order" sign on the device and do not use it.
WARNING: Do not connect more than one patient to the monitor. Do not connect
more than one monitor to a patient.
WARNING: The patient monitor is a prescription device and is to be operated by
qualified personnel only.

Cautions

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Cautions are identified by the CAUTION symbol shown above.

Caution statements identify conditions or practices that could result in damage to the equipment or other property.

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CAUTION: The monitor may not operate properly if it is operated or stored at conditions outside the ranges stated in this manual, or subjected to excessive shock or dropping.

CAUTION: When connecting the patient monitor to any instrument, verify proper operation before clinical use. Both the monitor and the instrument connected to it must be connected to a grounded outlet.

CAUTION: Accessory equipment connected to the monitor's data interface must be certified according to IEC60950 for data-processing equipment or IEC60601-1 for electromedical equipment. All combinations of equipment must be in compliance with IEC60601-1-1 system requirements. Anyone who connects additional equipment to the signal input or signal output port configures a medical system and is therefore responsible that the system complies with the requirements of IEC 60601-1-1 and the electromagnetic compatibility system standard IEC60601-1-2. If in doubt, consult Colin Medical Technology Technical Support Representative.

CAUTION: Risk of explosion if battery is replaced by an incorrect type.

CAUTION: Where the integrity of the external protective conductor in the installation or its arrangement is in doubt, equipment shall be operated from its internal electrical power source.

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INTRODUCTION

WARNING: Patient conditions may result in erroneous readings. If the measurements are suspect, verify the reading using another clinically accepted measurement method.

Intended Use for the BP-S510

The BP-S510 is intended to be used to monitor electrocardiography (ECG), heart rate (HR), noninvasive blood pressure (NIBP) - systolic, diastolic and mean arterial pressures, functional arterial oxygen saturation (SpO₂), pulse rate (PR), respiration (RR), temperature (Temp), invasive blood pressure (IBP) and capnography (EtCO₂ and InCO₂) for adult and neonatal patients in all areas of a hospital and hospital-type facilities. Monitor users should be skilled at the level of a technician, doctor, nurse or medical specialist.

Note: Hospital use typically includes such areas as general care floors, operating rooms, special procedure areas, intensive and critical care area, within the hospital. Hospital-type facilities include physician office-based facilities, sleep labs, skilled nursing facilities, surgical centers, and sub acute care centers.

About This Manual

This manual explains how to set up and use the BP-S510 patient monitor.

Read the entire manual including the *Safety Information* section, before you operate the monitor.

Identifying the BP-S510 monitor Configurations

The following table identities BP-S510 monitor configurations and how they are indicated. The model-option number and serial number are located on the back of the monitor. All information in this manual, including the illustrations, is based on a monitor configured with the Capnography (EtCO₂ and InCO₂), IBP and recorder. If the relevant functions do not exist, please verify your unit configuration.

Configuration	REF No.	Description	
BP-S510	112580	Standard	
		(ECG, NIBP, SpO ₂ , 2-channel Temperature, Respiration)	
BP-S510P	112581	Standard + Recorder	
BP-S510C	112582	Standard + Capnography	
BP-S510PC	112583	Standard + Capnography + Recorder	
S510-IBP	131324	2-channel IBP module	

Features for the BP-S510

Physical/Mechanical

The BP-S510 is a multi-parameter patient monitor which can be battery-operated when AC power source is not available.

Electrical

The BP-S510 is powered by an internal battery pack that typically provides one hour of monitoring from fully charged new batteries. The batteries are continuously recharged when the monitor is connected to AC power source. Refer to the **Battery Operation** section for details.

Display

The monitoring screen is a color LCD that shows all graphic and numeric patient information as well as status conditions and warning messages.

Jog dial

The jog dial provides user interaction with the display and the monitor functions. Rotating and pressing the jog dial allows the user to navigate and make changes to the display elements and monitor functions. Refer to the **Using the Monitor** section for details.

Auxiliary Outputs

The monitor provides RS-232, LAN and USB ports.

DESCRIPTION OF THE MONITOR

Front Panel Components



- 1 Visual alarm indicator
- 2 Alarm silence button
- 3 NIBP interval button
- 4 NIBP start/stop button
- 5 LCD
- 6 Jog dial
- 7 Power on indicator
- 8 Power on/off button
- 9 Battery charging indicator 1
- 10 Battery charging indicator 2
- 11 Record button
- 12 Trend button
- 13 Home button

Figure 1. Front Panel Components

Symbols	Description		
Ů∕⊙	Power on/off button turns the monitor on or off.		
\mathbb{N}	Record button prints measured data if an optional recorder is installed.		
Ķ	Alarm silence button silences the audible alarm temporarily. suspends the audible alarm by pressing over 2 seconds.		
	NIBP interval button allow you to set the NIBP auto measurement interval.		
	NIBP start/stop button toggles between starting and stopping NIBP measurements.		
	Home button exits a menu displayed on the screen and goes to the main screen.		
F	Trend button allows you to set the trend display.		
	Jog dial provides user interaction with the monitor to control the functions.		

Table 1 PB 8510 Controle

Rear Panel Components



Figure 2. Rear Panel Components

Left Panel Components



- 1 ECG connector
- Temperature channel 1 2
- Temperature channel 2 3
- SpO₂ connector 4
- 8 AC power connector 9
- Water trap (option) 10 CO₂ connector (option)

NIBP connector

- 5 IBP channel 1 (option)
- 11 CO₂ Filter (option)
- 6 IBP channel 2 (option)
- CO₂ Exhaust port (option) 12

Figure 3. Left Panel Components

7

Right Panel Components



Figure 4. Right Panel Components

Symbols	Description	Symbols	Description
\odot	Power on indicator	EC REP	EU representative
→⊡ 1	Battery charging indicator 1	● ●IP21	Dust and water resistance
→□ 2	Battery charging indicator 2	Â	Attention, consult accompanying documents
((Visual alarm indicator	CE 0434	CE mark
⊣♥	Type CF- Defibrillator proof	X	Crossed-out wheeled bin
A	ECG connector		Manufacturer
a	Temperature connector	M	Date of manufacture
SpO₂	SpO ₂ connector	REF	Reference number
	NIBP connector	SN	Serial number
	IBP connector	SIG-1060HPa	Environmental shipping/storage altitude limitations
CO ₂	CO ₂ connector	15%	Environmental shipping/storage humidity limitations
~	CO ₂ Filter	-20 C	Environmental shipping/storage temperature limitations
•	USB port		Fragile-handle with care
蛊	LAN port	ÎÎ	This way up
-Đ¢	RS-232 interface port	Ţ	Keep dry
1 12V 3.8Ah	Battery placement & rating	<u>_</u> !	Attention: consult accompanying documents
Ą	Equipotential terminal		Handle with care
AC IN ▲ 100-240V ~50/60Hz	AC power input rating	3	Stack up to 3 boxes

Displays



- 1 Alarm message area
- 2 Title of waveform parameter
- 3 Waveform
- 4 Waveform area
- 5 Informative message area
- 6 Battery status icon
- 7 Patient mode icon
- 8 Big number screen icon

- 9 Setup icon
- 10 Alarm limits icon
- 11 Time display
- 12 Numerical area
- 13 Alarm icon
- 14 Alarm limit values
- 15 Title of numeric parameter
- 16 Numeric value

Figure 5. Displays

Symbols	Description	Symbols	Description
FCG	ECC waveform icon	() 15 min	NIPD outo mode Interval
200		4 13 mm	NIBP auto mode interval
II	ECG lead pair	() 10 min	NIBP elapsed time
1mV	ECG size scale	%SpO₂	SpO_2 icon and unit
[ECG size bar		Pulse amplitude indicator
P1	P1 Label	RR/min	Respiration rate icon & unit
P2	P2 Label	Im	Respiration source icon: Im
ABP	P1 or P2 Label: Arterial Blood Pressure	Aw	Respiration source icon: Aw
СVР	P2 Label: Central Venous Pressure		Lung icon
PAP	P2 Label: Pulmonary Artery Pressure	InCO ₂	InCO ₂ icon: Inspired carbon dioxide concentration
LAP	P2 Label: Left Atrial Pressure	EtCO ₂	EtCO ₂ icon: End-tidal carbon dioxide concentration
PLETH	Plethysmograph icon	T1	Temperature channel 1 icon
CAPNO	Capnograph icon	Т2	Temperature channel 2 icon
RESP	Impedance respiration waveform icon	ΔΤ	Delta T icon T1-T2
🖤 bpm	HR/PR icon & unit	°C	Temperature unit: Celsius
ECG	HR source icon: ECG	°F	Temperature unit: Fahrenheit
P1	PR source icon: IBP1		Battery status icon
ABP	PR source icon: IBP1	A CHARACTER AND A CHARACTER AN CHARACTER AND A CHARACTER ANTER ANTER ANTER ANT	Big number screen icon
SpO ₂	PR source icon: SpO ₂		Setup icon
NIBP	PR source icon: NIBP	<u> </u>	Alarm limits icon
NIBP	NIBP icon	Ĥ	Patient mode: Adult
mmHg	NIBP, IBP or EtCO ₂ unit: mmHg	Đ	Patient mode: Neonatal
kPa	NIBP, IBP or EtCO ₂ unit: kPa	08:12	Time display
%	EtCO ₂ unit: %	100 90	Alarm limits value
S	Systolic pressure icon	•	Alarm icon
М	MAP or Mean pressure icon	×	Audible Alarm silence icon
D	Diastolic pressure icon	*	Audible Alarm suspend icon

Table 3. Display Symbols

Symbols Description		Symbols	Description
$\succ \prec$	NIBP graphical trend icon		HR/PR graphical trend icon
Δ	T2 graphical trend icon	×	SpO ₂ graphical trend icon
Т	T1 graphical trend icon	\vdash –1	IBP graphical trend icon
\$	EtCO ₂ graphical trend icon	+	Respiration graphical trend icon

Table 4. Display Colors

Function	Color
ECG and HR/PR	Green
Plethysmograph and SpO ₂	Cyan
NIBP	Yellow
Respiration	White
Temperature	Pink
Capnograph, InCO ₂ and EtCO ₂	Orange
IBP1 (P1 Label)	Purple
IBP2 (P2 Label)	Blue
General background	Black
Informative message	Black background, White font
Low priority alarm message	White background, Black font,
Medium priority alarm message	Yellow background, Black font,
High priority alarm message	Red background, Black font,
Battery status icon (normal)	Green
Battery status icon (low battery)	Yellow or Red (refer to Table 8)

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SETTING UP THE MONITOR

	WARNING: To ensure accurate performance and prevent device failure, do not
	expose the monitor to extreme moisture, including direct exposure to rain. Such
	exposure may cause inaccurate performance or device failure. Refer to
	Specification section.
	WARNING: The monitor should not be used adjacent to or stacked with other
	equipment. If adjacent or stacked use is necessary, the monitor should be
	observed to verify normal operation in the configuration it is to be used.
	WARNING: Make sure that the monitor speaker is not obstructed. Failure to do so
	could result in an inaudible alarm tone.
\wedge	CAUTION: Recharging the battery is strongly recommended when the battery has



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CAUTION: Recharging the battery is strongly recommended when the battery has not been recharged for 2 or more months.

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

Unpacking and Inspection

The monitor is shipped in one carton. Examine the carton carefully for evidence of damage. Contact Colin Medical Technology Technical Support Representative immediately if any damage is discovered. Refer to the **Maintenance** section for instructions on returning damaged items.

Note: Refer to **Performance Verification** section in the service manual for the detailed information.

List of Components

The following items are standard in the package.

	Figure	Description	REF	Q'ty
Main Unit		BP-S510	-	1
	BIP-5510 Coestion Manual	OPERATION MANUAL (English)	1731064	1
		AC POWER CORD	046111	1
		BATTERY	040074	1
	00	ROLL PAPER *Only when Recorder option is installed.	-	2
For NIBP		CUFF No.3 (12cm)	A013ZZ	1
		CUFF HOSE No.1 (3.5m)	A015ZZ	1
For ECG		ECG CABLE No.8	AY1005	1

Table 5. Standard Accessories

E 500	Figure	Description	REF	Q'ty
For ECG		ECG 3-LEAD WIRES No.5 (G,R,Y)	AG1002	1
For SpO ₂		SpO2 DURA SENSOR DS-100A	-	1
		SpO ₂ EXTENSION CABLE DOC-10	-	1
For CO ₂		SAMPLING SET (AIRWAY ADAPTER, NAFION TUBE & SAMPLING TUBE) *Only when CO ₂ option is installed.	C010ZZ	1

Optional items listed below can be ordered. Contact your local sales supplier for the detailed information.

	REF	Description	Unit	Min.Qty
NIBP	A012ZZ	CUFF No.2 (9cm)	рсе	-
-	A014ZZ	CUFF No.4(14cm)	рсе	-
-	AL021Z	CUFF No.10(2.5cm)	pce	10
-	AL022Z	CUFF No.11(3cm)	pce	10
-	AL023Z	CUFF No.12(4cm)	pce	10
	AL024Z	CUFF No.13(5cm)	рсе	10
	A016ZZ	CUFF HOSE No.2(1.5m)	pce	-
-	AL012Z	CUFF HOSE No.3(3.5m)	рсе	-
ECG	AG006Z	ECG ELECTRODES 25pcs/bag	bag	1
-	AG1003	ECG 5-LEAD WORES No.6(G,R,Y,W,B)	pce	-
SpO ₂	-	ADULT FINGER OXISENSOR MAX-A	-	-
-	-	CHILD FINGER OXISENSOR MAX-P	-	-
-	-	NEONTAL OXISENSOR MAX-N	-	-
-	-	INFANT OXISENSOR MAX-I	-	-
-	-	ADULT NASAL OXISENSOR MAX-R	-	-
-	-	MAX-FAST	-	-
CO ₂	C005ZZ	WATER TRAP	-	-
-	C003ZZ	NAFION TUBE	рсе	10
-	C004ZZ	SAMPLING TUBE	рсе	10
-	C002ZZ	AIR WAY ADAPTER	рсе	10
IBP	AS012Z	IBP DISPOSABLE TRANSDUCER DT-12	рсе	10
-	AS011Z	IBP DISPOSABLE TRANSDUCER DT-4812	рсе	10
	046432	IBP INTERFACE CABLE TC-COL-2	рсе	1
TEMP	AS004Z	BT Sensor Model 401J/Rectum, Gullet (produced by YSI)	pce	1
-	AS005Z	BT Sensor Model 402J/Rectum, Gullet/Small(produced by YSI)	pce	1
Others	1731064A	OPERATION MANUAL (English)	pce	1
-	1731065A	OPERATION MANUAL (German)	pce	1
-	1731066A	OPERATION MANUAL (French)	pce	1
-	1731093A	OPERATION MANUAL (Portugese)	pce	1
-	1731068A	OPERATION MANUAL (Spanish)	pce	1
-	1731071A	SERVICE MANUAL (English)	рсе	1
-	040074	BATTERY (BP-S510)	-	-
-	048151	ROLL PAPER BP-S510	-	-
-	A023ZZ	GROUDING WIRE TYPE1	рсе	1
	A024ZZ	GROUDING WIRE TYPE2	рсе	1

Table 6. Optional Accessories

Power Cable Connections

WARNING: Do not connect to an electrical outlet controlled by a wall switch because the device may be accidentally turned off.



CAUTION: If in doubt the integrity of the AC power source, the monitor must be operated from its internal battery.

AC Power

Make sure that the AC outlet is properly grounded and supplies the specified voltage and frequency ($100-240V \sim 50-60$ Hz).



Figure 6. AC Power connection

- 1. Connect the female connector end of the AC power cord to mains connector on the monitor's left panel.
- 2. Plug the male connector end of the AC power cord into a properly grounded mains outlet.
- 3. If necessary, connect grounding wire. Connect the grounding wire connector to the equipotential terminal on the rear panel. Now attach the clip end of the grounding wire to the medical equipment grounding terminal on the wall.
- 4. Verify that *Battery charging indicator* on the monitor's front panel is lit.
- Note: Even if the monitor is not turned on, the **Battery charging indicator** is lit when the AC power cord is connected into a mains outlet.

Note: If Battery charging indicator is not lit, check:

- the power cord
- the AC power inlet
- the power/ mains outlet
- No Battery

If the *Battery charging indicator* still is not lit although any problem is not found, contact qualified service personnel or your local supplier for assistance.

Measurement Cable Connections



Note: Both frequent checks by the operator on daily basis and more comprehensive technical checks less frequently are covered by this requirement in order to detect mechanical damage and damage to cables, etc.

ECG Cables and Leads

- 1. Connect an ECG cable to the "ECG" connector making sure that the connector arrow is pointing panel.
- 2. Attach the ECG lead wire to the end of the cable. (see Figure 3)

NIBP Hoses and Cuffs

- 1. Select an appropriate size cuff for the patient. (Refer to the NIBP Monitoring section.)
- 2. Connect the hose to the "CUFF" connector making sure to tighten the connector in the clockwise direction.
- 3. Attach the cuff to the end of the hose. (see Figure 3)

SpO₂ Cables and Sensors

- 1. Select an appropriate sensor for the patient and desired application. (Refer to the **SpO**₂ **Monitoring** section.)
- 2. Connect the extension cable DOC-10 to the "SpO₂" connector on the monitor's left panel.
- 3. Attach the sensor to the end of the cable. (see Figure 3)

Temperature Probes

- 1. Select the appropriate probe(s) for the desired application.(YSI 400 Series)
- 2. Connect the temperature probes to the "T1" and/or "T2" connector on the monitor's left panel. (see Figure 3)

IBP Transducers (option)

- 1. Connect the interface cable(s) for the transducer(s) to the IBP connector on the monitor's left panel. An interface cable for the transducer has to be selected correctly as it depends on the transducer type. (see Figure 3)
- 2. Set up the patient circuit according to the directions for use of the transducer, monitoring kit and IV set.

CO₂ Sampling set (option)

- 1. Connect a sampling tube to the "Water Trap" port on the monitor's left panel. (see Figure 3)
- 2. Connect the nafion tube to the sampling tube.
- 3. Connect the nafion tube to the airway adaptor.
- Note: If lead wires/cables, cuffs/hose cables, sensors and/or probes are not connected firmly, the monitor could lose signals from the patient.

BATTERY OPERATION

CAUTION: Recharging battery is strongly recommended when it has not been fully recharged for 2 or more months.



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CAUTION: When the voltage of the battery is very low, it is a possibility of not operating.

- Note: It is recommended that the monitor remain connected to AC power source when not in use. This will ensure a fully charged battery whenever it is needed.
- Note: As the battery is used and recharged over a period of time, the amount of time between the onset of the low battery alarm and the instrument shut-off may become shorter.

Operating the Monitor on Battery Power

The monitor has an internal battery that can be used to power the monitor when AC power source is not available. The battery status icon appears on the screen when the monitor is on battery power.



Figure 7. Battery Placement

- 1. Turn off the monitor.
- 2. Open the battery cover by pressing the latch.
- 3. Insert the battery carefully.

Table 7. Front	panel Indications	for	power	source
----------------	-------------------	-----	-------	--------

Power Connections	Front panel Indications
AC source	Battery charging indicator is lit.
Battery	Battery status icon appears on the screen.

The monitor cannot operate with a fully discharged battery. Before turning on the monitor with the battery that has been completely discharged, first plug the monitor into an AC outlet to charge the battery for minimum 3 minutes. The monitor may then be powered on.

A new, fully charged battery will provide 1 hour monitoring operation under the following conditions:

- Operation of ECG, Respiration, SpO₂, Temperature, CO₂ and IBP
- NIBP automatic measurement per 5 minutes
- No audible alarm condition
- No external communication operating
- No printing
- Ambient temperature at 25°C

Battery Status Indication

When operating on batteries, the battery status icon in the lower part of the display indicates the battery charge condition. See Table 8.

Battery Status Icons	Battery Status Icon Color	
	Green (constant)	
	Yellow (constant) ≤ 15 minutes	
	Red (flashing) ≤ 5 minutes	

Table 8. The Monitor Battery Status Icon

A low priority alarm occurs when the remaining battery power is only enough for 15 minutes of operation. The alarm message **'Low Battery'** appears on the screen and the visual alarm indicator is lit with yellow.

This alarm cannot be silenced while running on battery power. Connecting the monitor to AC power will silence the alarm.

A high priority alarm occurs for about 5 minutes before the monitor shuts off. The alarm message '*Critically Low-Battery Condition*' will appears and the visual alarm indicator will flash with red. After that, the monitor will automatically shut down. Connect the monitor to an AC power source to avoid any loss of trend data or settings.

Charging a Low Battery

- 1. Connect the monitor to AC power source to charge a low or depleted battery. (see the **Setting up the Monitor** section)
- 2. Verify that *Battery charging indicator* is lit with orange.

Table 9. Front Pane	I Indications for	Battery Status
---------------------	-------------------	-----------------------

Battery status	Battery charging indicator
Full charged	Green
Charging	Orange
Not installed	Off

Note: Even if the monitor is turned off, *Battery charging indicator* is lit while the battery is recharged.

Note: A full charge of a depleted battery takes over 12 hours per battery.

USING THE MONITOR

 Matrix
 WARNING: If the Power On Self-Test is not completed successfully, do not try to use the monitor.

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

 Matrix
 WARNING: Keep patients under close surveillance when monitoring. It is possible, although unlikely, that radiated electromagnetic signals from sources external to the patient and the monitor can cause inaccurate measurement readings. Do not rely entirely on the monitor readings for patient assessment.

Turning on the Monitor

Before using the monitor, confirm that the monitor is working properly and is safe to use as described below.



CAUTION: When power is applied, the monitor automatically starts the Power-On Self-Test (POST), which tests the monitor circuitry and functions. During POST, confirm that the monitor screen turns on. If the monitor screen does not function properly, do not use the monitor. Instead, contact qualified service personnel or your local supplier.

- Note: The post test tone sounds when the monitor completes the Power-On Self-Test (POST). This functions as an audible confirmation that the speaker is performing properly. If the speaker does not function, the alarm warning sounds cannot be heard.
- Note: If there is any problem on the speaker during the operation, the buzzer generates the sound automatically to alert the user to notice the speaker failure. If unusable sound like buzzer can be heard, do not use the monitor. Instead, please contact qualified service personnel or your local supplier.
- 1. Turn on the monitor by pressing *Power on/off button*. Confirm that the *Power on indicator* on the monitor's front panel is lit.
- 2. After the checksum for flash memory is completed, the monitor performs POST. The initial screen appears during POST. The initial screen displays the company logo, the version of system and the current time.
- 3. If there is no error, all indicators are lit for at least 2 seconds and the post pass tone sounds during POST. Confirm that the post pass tone sounds and all indicators are lit during POST.



Figure 8. Initial Screen

Note: The system version shown above is only an example.

- 4. After power-up diagnostics are successfully completed, the monitor is ready for operation.
- Note: If the monitor detects an internal problem during POST, the monitor will display an error code and will not display the monitoring screen. If an error code is displayed during POST, contact qualified service personnel or your local supplier for assistance.
- When the monitor detects valid signals, it displays real-time waveforms similar to Figure 9.



Figure 9. Typical Screen during monitoring

Setting Date and Time

You may set the date and time displayed on the screen and printed on the reports.

- 1. Rotate the jog dial to highlight *Time display*, and then press the jog dial to select *Date/Time menu*.
- 2. Rotate the jog dial to display the desired number for year, month, day, hour or minute, and then press the jog dial to select the desired number.

Note: The time format is 24 hours only. The date format can be set via Service menu.



Table 10. Date/Time Menu

Level 1 Menu	Level 2 Menu or Response
DATE/TIME MENU	
Date	Year
	Month
	Day
Time	Hour
	Min
Return	

Setting Basic Setup Parameters

This procedure will allow you to set Patient Mode, Record Speed, Wave Record Time, Wave Record Select, Record On Alarm, Auto List Record, Alarm Volume, HR/PR Tone Volume, Key Beep Volume, Sleep Mode, Main Screen and Service menu.



Table 11. Setup Menu

•		
Level 1 Menu	Level 2 Menu or Response	
SETUP MENU		
Patient Mode	Adult, Neonatal	
Record Speed*	25.0mm/s, 50.0mm/s	
Wave Record Time*	20 sec, Continuous	
Wave Record Select*	ECG 1 + ECG 2	
	ECG 1 + PLETH	
	ECG 1 + RESP	
	ECG 1 + (P1 Label)	
	ECG 1 + (P2 Label)	
	ECG 1 + CAPNO	
Record On Alarm*	On, Off	
Auto List Record*	On, Off	
Alarm Volume	1, 2, 3, 4, 5, 6, 7, 8	
HR/PR Tone Volume	Off, 1, 2, 3, 4, 5, 6, 7	
Key Beep Volume	Off, 1, 2, 3, 4, 5, 6, 7	
Sleep Mode	Off, 10, 20, 30 minutes	
Main Screen	4-ch Wave, 6-ch Wave, Big Number	
Service Menu	(Pass code required)	
Return		

Note: The menu options followed by an asterisk (*) are only displayed with an optional recorder installed.

Note: If there is no activity for 20 seconds, the monitor will return to main screen.

Patient Mode

- 1. Rotate the jog dial to highlight Setup icon. Press the jog dial to display the setup menu.
- 2. Rotate the jog dial to highlight *Patient Mode*, and then press the jog dial to select an appropriate mode: Adult or Neonatal.

Setting Record

If an optional recorder is installed, this menu will allow you to set **Record Speed**, **Wave Record Time, Wave Record Select, Record On Alarm** and **Auto List Record**. Refer to the **Printing** section for details.

Note: These menus are grayed out if no recorder installed in the monitor.

Setting Volume

Setting *Volume* allows you to adjust an audible alarm volume, Key beep volume and HR/PR tone volume. *Alarm volume* can be set level 1 to 8 and *Key beep volume* and *HR/PR tone volume* can be set level 1 to 7 or Off. (see Alarms and Limits section)

- 1. Rotate the jog dial to highlight *Alarm volume, Key beep volume* or *HR/PR tone volume*.
- 2. Press the jog dial. Levels of *Alarm volume, Key beep volume* or *HR/PR tone volume* will appear.
- 3. Rotate the jog dial to select a volume level. (see each volume level in the Table 11).
- 4. Press the jog dial to enter a desired volume into the monitor.

Sleep Mode

The monitor can be set to sleep mode for saving the power. The back light of the screen is turned on continuously when *Off* is selected, When *10 min, 20 min* or *30 min* is selected, the back light of the screen will be turned off automatically after the selected time if there is not any alarm condition and control by the user.

Setting Main Screen

You may select the number of the waveforms to be displayed; *4-ch wave*, *6-ch wave* or *Big number*.

The following are the default screens of waveforms and big numbers as per each configuration.



✓ 4-ch wave with basic configuration: ECG 1 + ECG 2 + SpO₂ + RESP

Figure 12. Basic configuration display



✓ 6-ch wave with CO₂ option: ECG 1 + ECG 2 + ECG 3 + SpO₂ + RESP + CAPNO

Figure 13. CO₂ option display

✓ 6-ch wave with IBP option: ECG 1 + ECG 2 + IBP 1 + IBP 2 + SpO₂ + RESP

	trev NBP meeting ¢oFF @····nin
ABP_150 	
^{CVP}	
	$\begin{bmatrix} 1 & 1 & 1 & 1 \\ DIA & 4 \end{bmatrix}_{33}^{133} DIA & 4 \end{bmatrix}_{33}^{133} DIA & 4 \end{bmatrix}_{33}^{133} DIA & 4 \end{bmatrix}_{33}^{123} DIA & 4 \end{bmatrix}_{33}^{123} DIA & 4 \end{bmatrix}_{33}^{123} DIA & 4 \end{bmatrix}_{33}^{133} DI$
	80 80 80 37.5 A 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
Image: marked black blac	· · · · · · · · · · · · · · · · · · ·

Figure 14. IBP option display

✓ 6-ch wave with IBP&CO₂ option: ECG 1 + ECG 2 + IBP 1 + IBP 2 + SpO₂ + CAPNO



Figure 15. IBP and CO₂ option display
Note: The user can select a desired waveform in each waveform area.

- 1. Rotate the jog dial to highlight the waveform area to be changed.
- 2. Press the jog dial to display the waveform menu.
- 3. Select *Waveform select* by rotating and pressing the jog dial.
- 4. Select the desired waveform by rotating and pressing the jog dial.
- Note: The monitor can display *Big Number Screen* by selecting *Big number icon* and return to the main screen by selecting *Big number icon* again.
- Note: The menus can be accessed in *Big Number Screen* without returning the main screen.
- Note: When an alarm condition occurs, the visual or audible alarm is activated in *Big Number Screen.*

Service Menu

This menu includes Save Settings on Power off, Audible Alarm Silence Period, Audible Alarm Suspend Period, Audible Alarm Type, AC Line Frequency, Unit Configuration, Language Setting, Date Format, Jog Dial Speed, LAN Setting, System Setting, System Test, NIBP Test and Demo Mode. Only authorized personnel is allowed to change the *Service Menu* settings. A pass code is required for access. Refer to the service manual for instructions.

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ALARMS AND LIMITS



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

General

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

- Visual alarm indication
- Audible alarm indication
- Physiological alarms including identification of out-of-limit vital signs
- Technical alarms

Note: The audible and visual alarms on the monitor, used in conjunction with clinical signs and symptoms, are the primary source for notifying medical personnel that a patient alarm condition exists.

Changing Alarm Volume

You can select an alarm volume level 1 to 8. Refer to the **Using the Monitor** section (see Figure 11, Table 11).

Taking an NIBP Measurement on Alarm

You can activate BP on Alarm via *NIBP menu*. Refer to the **Using the Monitor** section.

Initiating Print out on Alarm (Only when recorder option is installed)

You can activate Record on Alarm via Setup menu. Refer to the Printing section.

Alarm Priority and Messages

There are three possible priorities for visual and audible alarms: High, Medium, and Low. The high, medium and low priority messages are displayed in the alarm message area, and the informative messages are displayed in the informative message area. Refer to the **Troubleshooting** section for the recommended actions.

High Priority

Physiological Alarm				
Parameter	Condition	Messages		
	Over the systolic BP upper limit	NIBP: SYS upper limit violated.		
	Under the systolic BP lower limit	NIBP: SYS lower limit violated.		
	Over the diastolic BP upper limit	NIBP: DIA upper limit violated.		
NIDF	Under the diastolic BP lower limit	NIBP: DIA lower limit violated.		
	Over the map BP upper limit	NIBP: MAP upper limit violated.		
	Under the map BP lower limit	NIBP: MAP lower limit violated.		
ECG	Cardiac arrest	ECG: Asystole.		
	Over the heart rate upper limit	HR: Upper limit violated.		
	Under the heart rate lower limit	HR: Lower limit violated.		
	Over the pulse rate upper limit	PR({source}): Upper limit violated.		
	Under the pulse rate lower limit	PR({source}): Lower limit violated.		
	Respiration signal is not detected during	Resp: Loss of respiration signal.		
Posp	40sec.			
Resp	Over the respiration rate upper limit	Resp: Upper limit violated.		
	Under the respiration rate lower limit	Resp: Lower limit violated.		
	Over the systolic upper limit	{label}: SYS upper limit violated.		
	Under the systolic lower limit	{label}: SYS lower limit violated.		
IRD	Over the diastolic upper limit	{label}: DIA upper limit violated.		
IDI	Under the diastolic lower limit	{label}: DIA lower limit violated.		
	Over the mean upper limit	{label}: MEAN upper limit violated.		
	Under the mean lower limit	{label}: MEAN lower limit violated.		
%SnO2	Over the %SpO2 upper limit	SpO ₂ : Upper limit violated.		
785pO2	Under the %SpO2 upper limit	SpO2: Lower limit violated.		
Temp	Over the temperature upper limit	Temp{n}: Upper limit violated.		
Temp	Under the temperature lower limit	Temp{n}: Lower limit violated.		
Cappo	Incapable of detecting respiration for	Capno: Apnea.		
	certain period (set).			
	Over the EtCO ₂ upper limit	EtCO ₂ : Upper limit violated.		
Capilo	Under the EtCO ₂ lower limit	EtCO2: Lower limit violated.		
	Over the InCO ₂ upper limit	InCO ₂ : Upper limit violated.		
	Under the InCO ₂ lower limit	InCO2: Lower limit violated.		

Table 12. High Priority Alarm

	Technical Alarm				
Parameter	Condition	Messages			
	Inflation doesn't finish with specified time. Specified time is 30 sec for adult and 20 sec for neo. (C11)	NIBP: Check cuff (C11)			
NIBP	Measurement value can't be calculated even when inflated cuff pressure is deflated to specified pressure. Specified pressure is 10mmHg for adult and 5 mmHg for neo. (C12)	NIBP: Check cuff / Patient (C12)			

	Technical Alarm	
Parameter	Condition	Messages
	Deflation time is too long	NIBP: Cuff excessive artifact (C13)
	Deflation speed is too low (C13)	
	Maximum inflating pressure is too low to	NIBP: Cuff insufficient pressure
NIBP	calculate patient BP value. (C14)	(C14)
	Abnormal pulse due to arrhythmia and too	NIBP: Cuff irregular pulses (C15)
	much noise. (C15)	
	Collected pulse's movement is abnormal.	NIBP: Cuff motion artifact (C16)
	(C16)	
	Measurement time is beyond specified	NIBP: Cuff time-out (C17)
	time. Specified time 160sec for adult and	
	80sec for neo. (C17)	
	Number of detected pulses is beyond	NIBP: Cuff time-out, over 160 pulses
	specified number. Specified number is 160	(C18)
	for both adult and neo. (C18)	
	Cuff pressure is beyond the pressure	NIBP: Cuff pressure failure (C19)
	specified for the patient safety. Specified	
	150mmHg for non (C10)	
	The maximum value of collected pulse is	NIRD: Cuff work pulse (C20)
	too low (C20)	NIBF. Cull weak pulse (C20)
	Cuff size is inadequate to natient (C21)	NIBP: Check cuff, bose and mode
		(C21)
	NIBP module error	NIBP: Internal error (E03)
	BPM pressure sensor fault.	
	Pump operated for ten seconds, however	
	pressure does not change.	
	Check the connection of the cuff hose.	
	NIBP module error	NIBP: Internal error (E07)
	NIBP module error	NIBP: Internal error (E08)
	NIBP module error	NIBP: Internal error (E09)
	Fault detected in accordance with safety	
	monitoring to BPM IEC standards.	
	The pressure inside the cuff reaches the	
	standard pressure,	
	Standard pressure	
	Adult: 320mmHg	
	Neonatal: 15/mmHg	
		NIBP: Internal error (ROM)
F CC		
EUG Pasn	RESP module problem	Resp: Internal error
Кезр	IRE signal is not detected	
IBP	IBP module problem	IBP: Internal error
	SnO_2 signal is not detected	SnO_2 : Loss of nulse
%SpO2	SnO_2 module problem	SpO_2 : Loss of pulse.
Temn	Temperature module problem	TEMP{n}: Internal error
ТСПР	Cappo module have problem	Cappo: Internal error
0	CO2 sensor error	Capno: Sensor error
Capno		

Technical Alarm			
Parameter Condition		Messages	
	Critically low battery condition. The device will be automatically turned off within 5 minutes.	System: Critically low-battery condition.	
System	Real time clock is malfunction.	System: Real time clock error.	
	Watch dog timer is malfunction.	System M: WDT error.	
	Data memory is broken.	System M: RAM error.	
	Manufacturer use	System: Failure.	

Medium Priority

Table 13. Medium Priority Alarm

Technical Alarm Message			
Parameter	Condition	Messages	
ECG	Patient lead of cable disconnected.	ECG : Check ECG leads & electrodes.	
וסס	Transducer or interface cable is	{label}: Cable/Sensor disconnected.	
IBP	disconnected.		
%SpO2	Sensor or cable is disconnected.	SpO2: Check probe.	
Temp	Temperature probe is disconnected.	Temp{n}: Temperature probe disconnected.	
Canad	Occlusion alarm.	Capno: Occlusion.	
Caprio	Water trap full alarm.	Capno: Water trap full.	

Low Priority

Table 14. Low Priority Alarm

Technical Alarm Message			
Parameter	Condition	Messages	
ECG	ECG signal is saturated.	ECG: Signal saturation.	
Resp	Patient lead of cable disconnected.	Resp: Check Resp leads & electrodes.	
	Failed in zero calibration.	{label}: Unable to zero calibration.	
IBP	In case of following BP value -100 ≤ IBP ≤ -55, 330 ≤ IBP [mmHg]	<i>{label}</i> : Out of range.	
	Sensor disconnected from patient	SpO2: Check sensor.	
%SpO2	The sensor is broken down or defected.	SpO ₂ : Sensor failure.	
	Module is reset during operation.	SpO ₂ : Module reset.	
Temp	In case following temperature 14.0 ≤ T ≤ 45.0 °C	Temp{ <i>n</i> }: Out of range.	
	Failed in calibration	Capno: Zero calibration range error	
Conno		Capno: Zero calibration signal unstable error	
Capilo		Capno: High calibration range error	
		Capno: High calibration signal unstable error	
System	Low Battery, The device will be automatically turned off within 15 minutes.	System: Low battery.	

Informative Messages

Informative messages indicate a system condition that needs to be corrected.

Parameter	Condition	Messages	
	Measurement value can't be calculated	NIBP: Retry, Check cuff/Patient (C12)	
	even when inflated cuff pressure is		
	deflated to specified pressure. Specified		
	pressure is 10mmHg for adult and 5		
	mmHg for neo. (C12)		
	Deflation time is too long	NIBP: Retry, Cuff excessive artifact (C13)	
	Deflation speed is too low. (C13)		
	Maximum inflating pressure is too low to	NIBP: Retry, Cuff insufficient pressure	
	calculate patient BP value. (C14)	(C14)	
	Abnormal pulse due to arrhythmia and	NIBP: Retry, Cuff irregular pulses (C15)	
NIRP	too much noise. (C15)		
ND	Collected pulse's movement is	NIBP: Retry, Cuff motion artifact (C16)	
	abnormal. (C16)		
	Number of detected pulse is beyond	NIBP: Retry, Cuff time-out, over 160	
	specified number. Specified number is	pulses (C18)	
	160 for both adult and neo. (C18)		
	Cuff pressure is beyond specified	NIBP: Retry, Cuff pressure failure (C19)	
	pressure that is specified for patient		
	safety. Specified pressure is 300mmHg		
	for adult and 150mmHg for neo. (C19)		
	Cuff size is inadequate to patient. (C21)	NIBP: Retry, Check cuff, hose and mode	
ECC	Paper pulse detection is on	(C21) ECC: Depart datast is an	
ECG	Waiting for zoro calibration	LCG. Facer delect is on.	
IBP	Zero calibration in progress	{label}: No zero reading.	
	Current measurement is affected by	SpOr: Motion artifact	
%SnO2	natient motion		
700002	On pulse searching	SpO ₂ : Pulse search in progress	
	Messages about Zero calibration	Capno: Calibration in progress	
	process	Capno: Turn on CO ₂ (10%) calibration gas.	
Capno		Capno: Turn off CO ₂ (10%) calibration gas.	
		Capno: Calibration gas detected.	
	Capno module is warming up.	Capno: Warming up in progress.	
	Recorder has no paper.	System: No recorder paper.	
System	Device abnormally shut down last time	System: Abnormally shut down last time.	
	Device has no recorder module.	System: No recorder installed.	
	Alarm sound is silenced.	Audible alarm silenced.	
	Alarm sound is suspended.	Audible alarm suspended.	
Other	Alarm sound is inhibited.	Audible alarm inhibited.	
Other	Device in Demo Mode	Demo Mode.	
	Exit scrolling in trend screens	Press jog dial to exit scroll.	
	Contact your service personnel.	Contact your service personnel.	

Note: There may be other informative messages that are not listed above.

Visual Alarm Indication

Alarm Category	Color	Alarm Indicator Flashing Rate
High priority	Red	5 flashes in 3 seconds (1.7Hz)
Medium priority	Yellow	5 flashes in 8 seconds (0.6Hz)
Low priority	Yellow	Always on (non-flashing)

Table 16. Visual Alarm Characteristics

Note: *Visual alarm Indicator* on the right top of the front panel responds with the flashing rates described in Table 16 when an alarm occurs.

When the **high priority alarm** is activated, a non-flashing alarm message is displayed. The numerical area will flash red.

When a **medium priority alarm** is activated, a non-flashing alarm message is displayed. The numerical area will flash yellow.

When a **low priority alarm** activated, a non-flashing alarm message is displayed. The numerical area will change to yellow.

Audible Alarm Indication

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

... . . .

WARNING: Make sure that the monitor speaker is not obstructed. Failure to do so could result in an inaudible alarm tone.

Table	e 17.	Audible	e Alarm	Charact	teristics	

Alarm Category	Tone Pitch	Beep Rate
High priority	High	10 beeps in 10 seconds (976Hz)
Medium priority	Medium	3 beeps in 16 seconds (697Hz)
Low priority	Low	Non-beep

Note: Audible alarms may be decreased in volume as described in Table 11 or temporarily silenced.

Verifying Visual and Audible Alarm Indication

If the monitor fails to perform as specified in this test, contact qualified service personnel or Mediana Technical Support Representative for assistance.

You can verify the alarm operation for all parameters like ECG, SpO_2 , NIBP, Temp, and RESP by following the below procedures.

- 1. Connect the monitor to an AC power source.
- 2. Press *Power button* to turn on the monitor.
- 3. Connect the simulator to sensor input cable and connect cable to monitor.
- 4. Set the simulator to smaller value than the lower alarm limit on the monitor.
- 5. Verify following the monitor reaction:

- a. The monitor begins to track the physiological signal from the simulator.
- After about 10 to 20 seconds, the monitor displays the value measured as specified by simulator. Verify values are within the tolerances specified in Specification section for each parameter (ECG, SpO₂, NIBP, Temp, RESP, IBP, EtCO₂).
- c. Audible alarm sounds.
- d. Visual alarm Indicator on the front panel flashes.
- e. "Lower limit violated" message is displayed.
- f. The numerical area flashes, indicating the parameter has violated default alarm limits.

Changing Alarm Limits

A

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

You can change alarm limits from default values, if necessary.

Alarm limits or Alarm suspension may be set in two ways:

- Via interaction with HR/PR, SpO₂, NIBP, Respiration, Temperature, CO₂ and IBP menu or
- Via interaction with *Alarm limits menu* that presents the limits in all the parameters at one time

Setting Alarm Limits via Alarm Limits Menu

1. Rotate the jog dial to highlight *Alarm limits icon* on the lower of the screen, then press the jog dial to display *Alarm limits menu*



2. Press the jog dial to select *Alarm limits*. The monitor will display all alarm limits that are currently in effect for all monitored parameters. Select the alarm limits to set.



Table 18. Alarm Limits Menu

Level 1 Menu	Level 2 Menu or Response
ALARM LIMITS MENU	
Record on Alarm	On, Off
Audible Alarm Silence Period	(30, 60, 90, 120 seconds)
Audible Alarm Suspend Period	(Off,10, 20, 30, 60 minutes, Indefinite)
Alarm Limits	HR/PR, NIBP (SYS, DIA, MAP), RESP, SpO ₂
	P1 (SYS, DIA, MAP), P2 (SYS, DIA, MAP)
	InCO ₂ , EtCO ₂ , T1, T2
	Alarm Suspend for each parameter
Alarm Limits Display	On, Off
Auto Alarm	On, Off
Auto Alarm Setting	% Setting for each parameter
Return	

Note: Record on Alarm can be only set via Setup menu.

Note: Audible alarm silence period and audible alarm suspend period can be only set via *Service menu*.

Alarm Limits Ranges

Table 19 describes the possible alarm limits. The monitor is shipped with factory default settings.

Note: Authorized personnel can define the way to save the settings upon power off: custom, backup and default. The detailed information is described in the service manual.

D	Harris I have been the		Deschafter
Parameters	opper Limit, Default	Lower Limit, Default	Resolution
HR/PR (BPM)			
Adult	35 ~ 305 BPM, 180 BPM	30 ~ 300 BPM, 40 BPM	5 BPM
Neonatal	35 ~ 305 BPM, 200 BPM	30 ~ 300 BPM, 50 BPM	5 BPM
NIBP Systolic	(mmHg, kPa)		
A	60 ~ 260 mmHg, 200 mmHg	50 ~ 250 mmHg, 70 mmHg	10 mmHg
Adult	(8.0 ~ 34.7 kPa, 26.7 kPa)	(6.7 ~ 33.3 kPa, 9.3 kPa)	(1.3 kPa)
Numerated	40 ~ 130 mmHg, 130 mmHg	30 ~ 120 mmHg, 50 mmHg	10 mmHg
Neonatai	(5.3~ 17.3 kPa, 17.3 kPa)	(4.0 ~ 16.0 kPa, 6.7 kPa)	(1.3 kPa)
NIBP Diastolic	(mmHg, kPa)		
	$40 \sim 210 \text{ mmHg}$ 160 mmHg	30 ~ 200 mmHa, 30 mmHa	10 mmHa
Adult	$(5.3 \sim 28.0 \text{ kPa}, 21.3 \text{ kPa})$	$(4.0 \sim 26.7 \text{ kPa}, 4.0 \text{ kPa})$	(1.3 kPa)
	$20 \sim 100 \text{ mmHg} \ 100 \text{ mmHg}$	$10 \sim 90 \text{ mmHg}$ 10 mmHg	10 mmHg
Neonatal	$(2.7 \sim 13.3 \text{ kPa} \ 13.3 \text{ kPa})$	$(1.3 \sim 12.0 \text{ kPa} = 1.3 \text{ kPa})$	(1.3 kPa)
	mHa kPa)	(1.0 12.0 ki d, 1.0 ki d)	(1.0 Kr d)
	$50 \sim 240 \text{ mmHg} \cdot 180 \text{ mmHg}$	$40 \sim 230$ mmHg 40 mmHg	10 mmHa
Adult	$50 \sim 240$ mm/g, 160 mm/g	$40 \sim 230$ mm/g, 40 mm/g	
	$(0.7 \sim 52.0 \text{ KFd}, 24.0 \text{ KFd})$	$(5.5 \sim 50.7 \text{ KFd}, 5.5 \text{ KFd})$	(1.3 KFd)
Neonatal	$30 \sim 110 \text{ mmg}, 110 \text{ mmg}$	$20 \sim 100$ mmg, 20 mmg	
0	(4.0 ~ 14.7 KPa, 14.7 KPa)	(2.7 ~ 13.3 KPa, 2.7 KPa)	(1.3 KPa)
SpO ₂ (%)			
Adult	70 ~ 100 %, 100 %	69 ~ 99 %, 90 %	1%
Neonatal	70 ~ 100 %, 100 %	69 ~ 99 %, 85 %	1%
Respiration (E	3PM)		
Adult	5 ~ 155 BPM, 30 BPM	0 ~ 150 BPM, 0 BPM	5 BPM
Neonatal	5 ~ 155 BPM, 50 BPM	0 ~ 150 BPM, 0 BPM	5 BPM
Temperature (°C, °F)		
	15.0 ~ 45.5°C, 38.0 °C	14.5 ~ 45.0 °C, 14.5 °C	0.5°C
Adult	(59.0 ~ 113.9°F, 100.4°F)	(58.1 ~ 113°F, 58.1°F)	(0.9°F)
	15.0 ~ 45.5 °C. 39.0 °C	14.5 ~ 45.0 °C. 14.5 °C	0.5°C
Neonatal	(59.0~ 113.9°F. 102.2°F)	(58.1 ~ 113°F, 58.1°F)	(0.9°F)
EtCO ₂ (mmHa	, kPa, %)		
	$2 \sim 80 \text{ mmHa} 80 \text{ mmHa}$	$0 \sim 78 \text{ mmHg} 0 \text{ mmHg}$	2 mmHa
Adult	$(0.3 \sim 10.7 \text{ kPa} \cdot 10.7 \text{ kPa})$	$(0 \sim 10.4 \text{ kPa} \ 0 \text{ kPa})$	(0.3 kPa)
/ tault	$(0.3 \sim 10.5 \% \ 10.5 \%)$	$(0 \sim 10.3 \% 0 \%)$	(0.3 %)
	$2 \sim 80 \text{ mmHg} = 80 \text{ mmHg}$	$0 \sim 78 \text{ mmHg} 0 \text{ mmHg}$	2 mmHa
Neonatal	$(0.3 \sim 10.7 \text{ kPa} \cdot 10.7 \text{ kPa})$	$(0 \sim 10.4 \text{ kPa} \cdot 0 \text{ kPa})$	(0.3 kPa)
NCONALAI	$(0.3 \sim 10.5 \% 10.5 \%)$	$(0 \sim 10.3 \% 0 \%)$	(0.3 %)
InCO (mmHa	(0.5 × 10.5 %, 10.5 %)	(0 10:3 70; 0 70)	(0.5 70)
	, KF d , 70)	0 . 19 mmHg 0 mmHg	2 mmHa
Adult	$2 \sim 20$ mmHg, 20 mmHg	$0 \sim 10 \text{ mmg}, 0 \text{ mmg}$	∠ IIIII⊡y (0.2 kDa)
Adult	$(0.3 \sim 2.7 \text{ KPa}, 2.7 \text{ KPa})$	$(0 \sim 2.4 \text{ KPa}, 0 \text{ KPa})$	(0.3 KPa)
	(0.3 ~ 2.6 %, 2.6 %)	(0 ~ 2.4 %, 0 %)	(0.3 %)
Number	$2 \sim 20$ mmHg, 20mmHg	$0 \sim 18 \text{ mmHg}, 0 \text{ mmHg}$	2 mmHg
Neonatal	(0.3 ~ 2.7 kPa, 2.7 kPa)	(0 ~ 2.4 kPa, 0 kPa)	(0.3 kPa)
	(0.3 ~ 2.6 %, 2.6 %)	(0.3 ~ 2.6 %, 0 %)	(0.3 %)
IBP1 Systolic	(mmHg, kPa)		
Adult	-50 ~ 260 mmHg, 200 mmHg	-60 ~ 250 mmHg, 70 mmHg	10 mmHg
, laan	(-6.7 ~ 34.7 kPa, 26.7 kPa)	(-8.0 ∼ 33.3 kPa, 9.3 kPa)	(1.3 kPa)
Neonatal	-50 ~ 260 mmHg, 130 mmHg	-60 ~ 250 mmHg, 50 mmHg	10 mmHg
riconatai	(-6.7 ~ 34.7 kPa, 17.3 kPa)	(-8.0~33.3 kPa, 6.7 kPa)	(1.3 kPa)
IBP1 Diastolic			
Adult	-50 ~ 260 mmHg, 160 mmHg	-60 ~ 250 mmHg, 30 mmHg	10 mmHg
	(-6.7 ~ 34.7 kPa, 21.3 kPa)	(-8.0 ~ 33.3 kPa, 4.0 kPa)	(1.3 kPa)
Nooratal	-50 ~ 260 mmHg, 100 mmHg	-60 ~ 250 mmHg, 10 mmHg	10 mmHg
ineoriatai	(-6.7 ~ 34.7 kPa, 13.3 kPa)	(-8.0 ~ 33.3 kPa, 1.3 kPa)	(1.3 kPa)
IBP1 Mean			
	-50 ~ 260 mmHa. 180 mmHa	-60 ~ 250 mmHa. 40 mmHa	10 mmHa
Adult	(-6.7 ~ 34.7 kPa, 24.0 kPa)	(-8.0 ~ 33.3 kPa, 5.3 kPa)	(1.3 kPa)
Neonatal	-50 ~ 260 mmHa. 110 mmHa	-60 ~ 250 mmHa. 20 mmHa	10 mmHa
	,		

Table 19. Alarm Limits Ranges

Parameters	Upper Limit, Default	Lower Limit, Default	Resolution
	(-6.7 ~ 34.7 kPa, 14.7 kPa)	(-8.0 ~ 33.3 kPa, 2.7 kPa)	(1.3 kPa)
IBP2 Systolic			
Adult	-50 ~ 260 mmHg, 200 mmHg	-60 ~ 250 mmHg, 70 mmHg	10 mmHg
Adult	(-6.7 ~ 34.7 kPa, 26.7 kPa)	(-8.0 ~ 33.3 kPa, 9.3 kPa)	(1.3 kPa)
Noonatal	-50 ~ 260 mmHg, 130 mmHg	-60 ~ 250 mmHg, 50 mmHg	10 mmHg
Neonatai	(-6.7 ~ 34.7 kPa, 17.3 kPa)	(-8.0 ~ 33.3 kPa, 6.7 kPa)	(1.3 kPa)
IBP2 Diastolic			
Adult	-50 ~ 260 mmHg, 160 mmHg	-60 ~ 250 mmHg, 30 mmHg	10 mmHg
Adult	(-6.7 ~ 34.7 kPa, 21.3 kPa)	(-8.0 ~ 33.3 kPa, 4.0 kPa)	(1.3 kPa)
Noonatal	-50 to 260 mmHg, 100 mmHg	-60 to 250 mmHg, 10 mmHg	10 mmHg
Neonatai	(-6.7 ~ 34.7 kPa, 13.3 kPa)	(-8.0 ~ 33.3 kPa, 1.3 kPa)	(1.3 kPa)
IBP2 Mean			
Adult	-50 to 260 mmHg, 180 mmHg	-60 to 250 mmHg, 40 mmHg	10 mmHg
	(-6.7 ~ 34.7 kPa, 24.0 kPa)	(-8.0 ~ 33.3 kPa, 5.3 kPa)	(1.3 kPa)
Noonatal	-50 to 260 mmHg, 110 mmHg	-60 to 250 mmHg, 20 mmHg	10 mmHg
Neonalai	(-6.7 ~ 34.7 kPa, 14.7 kPa)	(-8.0 ~ 33.3 kPa, 2.7 kPa)	(1.3 kPa)

Alarm Limits Display

When the alarm limits display is **On**, the monitor displays the alarm limits value on numerical areas.

Auto Alarm

When the auto alarm is **On**, the monitor automatically sets the alarm limits upon the current measurement values by specified percentage.

Auto Alarm Setting

You can select the percentages each parameter to set automatically the alarm limits.

- 1. Select *Auto alarm setting* in the alarm limits menu by rotating the jog dial.
- 2. Select the parameter to be changed. Change the value by rotating the jog dial.



Figure 18. Auto Alarm setting menu

Audible Alarm Silence



When an alarm occurs, you can silence the audible alarm for audible alarm silence period (30, 60, 90 or 120 seconds) selected via service menu. However, visual alarms continue during this time. The factory default of audible alarm silence period is 120 seconds.

To silence an audible alarm:

- 1. Press *Alarm silence button* to immediately silence the alarm tone. The alarm resumes after the audible alarm silence period if the alarm condition has not been corrected.
- 2. Check the patient and provide appropriate care.

During the audible alarm silence period, you can press *Alarm silence button* again to reenable the audible alarm tones. Also, if another alarm occurs during the audible alarm silence period, the audible alarm tones will be automatically re-enabled.



Figure 19. Audible Alarm Silence Display

Note: The audible alarms caused by some technical errors may be canceled by pressing *Alarm silence button*. However, battery failure and physiological alarms cannot be canceled until the alarm condition is corrected.

Audible Alarm Suspend

WARNING: If an alarm condition occurs while in the Alarm Suspend state, the only alarm indication at the monitor will be visual displays related to the alarm condition.

To initiate an audible alarm suspend:

- 1. To initiate an audible alarm suspend, press *Alarm silence button* and hold it for at least 2 seconds.
- 2. To cancel the suspend condition, press *Alarm silence button* for 2 seconds again.

Note: You may disable physiological alarms of each vital sign via *HR/PR, SpO₂, NIBP, Respiration, Temperature, IBP, CO₂ or Alarm/Limits menu*.

This action disables audible alarms for a user-defined *Audible alarm suspend period* (10, 20, 30 or 60 minutes) selected via service menu.

If Audible alarm suspend period is set to other than *Off* or *Indefinite*, the audible alarm is not activated for the time interval and the message "*Audible alarm suspended*" is displayed. If *Off* is selected, the alarm suspension is not allowed to activate. If *Indefinite* is selected, the audible alarm suspension continues until canceled by pressing *Alarm silence button* again and the message "*Audible alarm inhibited*" is displayed. In this case, the monitor will remind the user every 3 minutes that the audible alarm is suspended or inhibited.

Note: The periods can only be changed by authorized personnel via Service menu.



Figure 20. Audible Alarm Suspend Display

ECG MONITORING

WARNING: For best product performance and measurement accuracy, use only accessories supplied or recommended by Colin Medical Technology. Use accessories according to the manufacturer's directions for use and your facility's standards.
WARNING: Line isolation monitor transients may resemble actual cardiac waveforms and thus inhibit heart rate alarms. Such transients may be minimized by proper electrode and cable placement, as specified in this manual and electrode directions for use.
WARNING: Do not use damaged ECG leads. Do not immerse ECG leads completely in water, solvents, or cleaning solutions. Do not sterilize ECG leads by irradiation, steam, or ethylene oxide. Follow the manufacturer's directions for use.
WARNING: Do not use ECG electrodes with expired dates. Do not use defective ECG electrodes. These might cause improper performance.
WARNING: ECG cables may be damaged if they are connected to a patient during defibrillation. Cables that have been connected to a patient during defibrillation should be checked for functionality before using again.
WARNING: It is possible for the patient to receive a burn due to an improperly connected electrosurgical unit. Additionally, the monitor could be damaged or measurement errors could occur. Place the ECG cable and leads as far as possible from the site of the electrosurgical unit and from the electrosurgical cables. This will minimize interference and the risk of burns to the patient.
WARNING: For pacemaker patients, the monitor may continue to count pacemaker rate during occurrences of cardiac arrest or some arrhythmias. To reduce the likelihood of this, ensure that the Pacer Detect setting is ON in the ECG waveform menu when monitoring such patients. Do not rely entirely upon the monitor alarms. Keep pacemaker patients under close surveillance.
WARNING: To ensure patient safety, the conductive parts of the ECG electrodes (including associated connectors) and other patient-applied parts should not contact other conductive parts, including earth ground, at any time.

General

The process of depolarization and repolarization of the myocardium generates electric potentials that are sensed by ECG electrodes on the skin surface. These electrodes are typically attached to the patient's right arm, left arm, and left leg. The monitor processes and amplifies these signals and presents the ECG waveform on the screen. In addition to the acquisition of the QRS complex, the circuitry performs a number of other functions. The monitor can display:

- Heart rate in beats per minute
- Detection of a "lead off" condition if an electrode is disconnected or poorly connected
- Detection of the presence of pacemaker signals within the ECG waveform complex

Note: Occasionally, electromagnetic interference beyond the range guaranteed from manufacture's declaration may cause the monitor to display an "Check ECG Leads & Electrodes" alarm. This occurrence is rare, and duration should be short. When the interference ceases, the monitor removes the "Check ECG Leads & Electrodes" alarm. Refer to **Specification** section

Setup Connections

- Note: Colin Medical Technology recommends the use of silver/silver chloride electrodes (Ag/AgCl). When dissimilar metals are used for different electrodes, the electrodes may be subject to large offset potentials due to polarization, which may be severe enough to prevent obtaining an ECG trace. Using dissimilar metals may also increase recovery time after defibrillation.
- 1. Select the electrodes to be used. Use only one type of electrode on the same patient to avoid variations in electrical resistance. Prepare the electrode sites according to electrode manufacturer's instructions. See Figure 21 and 22 for electrode placement configurations.



Figure 21. Standard 3 Electrode Placement



Figure 22. 5 Electrode Placement

Note: One of 5-1 to 5-6 Lead electrode placement sites for the fifth lead.

- 2. Connect the ECG lead wires and ECG cable.
- 3. Connect the ECG cable to the ECG connector on the monitor's left panel.

4. Attach the leads to the electrodes, and then apply the electrodes to the patient, using the color-code guide in Table 20. Verify that the desired Lead Selection is active in the ECG waveform area. Refer to Table 21. Lead II is best suited for most monitoring situations.

Lead	AAMI	IEC
1. Right arm	White (RA)	Red (R)
2. Left arm	Black (LA)	Yellow (L)
3. Left leg	Red (LL)	Green (F)
4. Right leg	Green (RL)	Black (N)
5-1 to 5-6. V (Chest)	Brown (V)	White (C)

Table 20. ECG Lead Colors

Table 21. ECG Lead Pairs

Lead-Selection	Electrode Differential (AAMI)	Electrode Differential (IEC)
I	RA LA	R L
I	RA LL	R F
III	LA LL	LF
V (Chest)	(RA+LA+LL)/3 Chest (V)	(R+L+F)/3 Chest (C)
aVR	– (Lead I + Lead III/2)	– (Lead I + Lead III/2)
aVL	(Lead I – Lead III)/2	(Lead I – Lead III)/2
aVF	(Lead II + Lead III)/2	(Lead II + Lead III)/2

Description of HR/PR Menu Functions

The calculated Heart rate/Pulse rate may be derived from different sources (ECG, IBP, SpO₂ or NIBP) as shown by the icon in the HR/PR numerical area.



Figure 24. HR/PR Menu

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Table 22	2. HR/	PR N	lenu
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Level 1 Menu	Level 2 Menu or Response
HR/PR MENU	
HR/PR Source	Auto (ECG>IBP>SpO ₂ >NIBP)
	HR (ECG)
	PR (IBP>SpO ₂ >NIBP)
(Alarm Limits Adjustment)	
	Upper Alarm Limit
▼	Lower Alarm Limit
HR/PR Alarm Suspend	On, Off
Return	

HR/PR Source

You may select HR, PR or Auto to decide the source of the heart rate or pulse rate. If you select Auto, the monitor automatically derives the heart rate or pulse rate from one of the monitoring parameters in this order of priority: ECG, IBP, SpO2 or NIBP. When HR is selected, the heart rate is measured from ECG. When PR is selected, the pulse rate is measured from in order of IBP, SpO2 or NIBP. The color of the source icon will be changed according to current source. If the pulse rate is derived from NIBP, the value will be displayed for only 180 minutes after the NIBP measurement, then the value will be removed from the display. The HR/PR tone volume can be adjusted in the Setup menu. Refer to the Using the Monitor section. (See Figure 11, Table 11)

Description of ECG Waveform Menu Functions



- 1 ECG waveform icon 4 ECG lead pair
 - ECG waveform 5
- ECG size bar 3 ECG size scale

2

Figure 25. ECG Waveform display

		EGG NIBP morty \$ 120/80 € 1000 €
EGG CBBIG SBICET AUTO Lead Select II Sweep Speed 25.5mm/s Size Pacer Detect Off Filter Mode Monitor Waveform Select ECG		APP mmHs 120/80 [100] ^{1990 M} (100) ▲ 1100 [100] ^{CVP} mmHs 100/1 [100] ▲ 1250 ^{CVP} mmHs 100/1 [100] ▲ 1250 ^{CVP} mmHs 100/1 [100] ▲ 1250 ^{CVP} mmHs
Return		% \$
	08:12	^{12 °C} 36.8 ▲ 38.0

Figure 26. ECG Waveform Menu

Level 1 Menu	Level 2 Menu or Response
ECG WAVEFORM MENU	
ECG cable select	Auto, 5 Leads, 3 Leads
Lead Select	Lead I, II, III, aVR, aVL, aVF, V (Chest Lead)
Sweep Speed	12.5 mm/s, 25.0 mm/s, 50.0 mm/s
Size (mm per 1mV)	×1/4, ×1/2, ×1, ×1.5, ×2
	Note: The size is selected by pressing on the up/down
	button of the screen.
Pacer Detect	On, Off
Filter Mode	Monitor, Low Extend, Filter, Respiration Rejection
Waveform Select	ECG, Pleth, Respiration, (P1 Label),(P2 Label), Capno
Return	

Table 23. ECG Waveform Menu

ECG Cable Select

When ECG cable select is set to *Auto*, the monitor sets ECG leads automatically. Also, you can select the 3 Leads or 5 Leads manually.

Lead Select

When leads are attached, the *Lead Select* menu is displayed with the available ECG lead selection. You can select the desired ECG lead. For more information about the lead selection, refer to Table 21.

Sweep Speed

The user-selectable sweep speed determines the speed at which the ECG waveform trace moves across the screen. *Sweep Speed* can be selected from 12.5 mm/s, 25.0 mm/s and 50.0 mm/s, and ECG waveform is synchronized with Pleth waveform and IBP waveform.

Size

The user-selectable ECG waveform size allows you to adjust the amplitude of an ECG waveform. The size can be selected from $\times 1/4$, $\times 1/2$, $\times 1$, $\times 1.5$ or $\times 2$. When the size is $\times 1$ selected, 1mV ECG signal is shown as 1cm on the display and on the print-out.

Pacer Detect

Pacer detect should always be **On** for patients with pacemakers (refer to the warning in this section). When **Pacer detect** is **On**, the monitor detects and filters pacemaker-generated signals so that they will not be calculated in determining a patient's heart rate. When monitoring the patient without pacemaker, Pacer detect should be set to **Off** to avoid misdiagnosis.

Filter Mode

The monitor can filter ECG waveform noise with different ranges of frequency response: **Low Extend** (0.05 Hz to 40 Hz): Expands the range to display very low frequencies down to 0.05 Hz.

Filter (0.5 Hz to 30 Hz): Generally called a filter mode it reduces ECG waveform noise. **Monitor** (0.5 Hz to 40 Hz): Choose this mode to see just the ECG waveform monitoring. **Respiration Rejection** (1 Hz to 40 Hz): Removes the respiration signal measured by impedance method.

NIBP MONITORING

	WARNING: For best product performance and measurement accuracy, use only accessories supplied or recommended by Colin Medical Technology. Use accessories according to the manufacturer's directions for use and your facility's standards.
	WARNING: Inaccurate measurements may be caused by incorrect cuff application or use. This can include placing the cuff too loosely on the patient, using the incorrect cuff size, or not placing the cuff at the same level as the heart, leaky cuff or hose or excessive patient motion.
	WARNING: In some cases, rapid, prolonged cycling of an oscillometric, noninvasive blood pressure monitor cuff has been associated with any or all of the following: ischemia, purpura, or neuropathy. Periodically observe the patient's limb to make sure that the circulation is not impaired for a prolonged period of time. Also make sure the cuff is placed according to directions in this manual and the cuff directions for use.
	WARNING: Do not place the cuff, the catheter or SpO ₂ sensor on an extremity being used for intravenous infusion or any area where circulation is compromised or has the potential to be compromised.
	WARNING: As with all automatically inflatable blood pressure devices, continual cuff measurements can cause injury to the patient being monitored. Weigh the advantages of frequent measurement and/or use of CONT mode against the risk of injury.
	WARNING: Ensure the patient is quiet with minimal movement during NIBP readings; minimize the patient's shivering.
	WARNING: Never place the cuff on extremity being used for intravenous infusion or any area where circulation is compromised or has the potential to be compromised. Never fit NIBP system with Luer Lock adapters.
	WARNING: Never use an adult monitor setting or cuff for an NIBP measurement on a neonatal patient. Adult inflation limits can be excessive for neonatal patients, even if a neonatal cuff is used.
•	CAUTION: In the automatic mode, the monitor displays results of the last blood pressure measurement until another measurement starts. If a patient's condition changes during the time interval between measurements, the monitor will not detect the change or indicate an alarm condition.
•	CAUTION: Any excessive patient motion may cause inaccurate measurements of non-invasive blood pressure. Minimize motion to improve blood pressure measurements.
•	CAUTION: Do not apply the blood pressure cuff to the same extremity as the one to which the SpO ₂ sensor or IBP catheter is attached. Cuff inflation can disrupt SpO ₂ monitoring and lead to nuisance alarms.
\diamond	CAUTION: Make sure that heavy objects are not placed on the cuff hose. Avoid crimping or undue bending, twisting, or entanglement of the hose.

Note: Blood pressure measurements can be affected by the position of the patient, the patient's physiological condition and other factors.

General

The monitor performs Non-Invasive Blood Pressure measurements using the oscillometric measuring technique. A notarized pump inflates the cuff to initially blocking the flow of blood in the extremity. Then, under monitor control, the pressure in the cuff is gradually reduced, while a pressure transducer detects air pressure and transmits a signal to the NIBP circuitry.

When the cuff pressure is still above systolic pressure, small pulses or oscillations in the cuff pressure begin to be sensed by the transducer. As the cuff continues to deflate, oscillation amplitude increases to a maximum and then decreases. When maximum oscillation amplitude occurs, the cuff pressure at that time is measured as mean arterial pressure (MAP). The systolic and diastolic pressures are calculated based on analysis of the oscillation amplitude profile.

Oscillometric Method

The blood pressure values are determined by measuring the small oscillations (changes) in the cuff pressure caused by the heart's contractions as the pressure in the cuff is released. Colin's measurement technology utilizes a unique deflation technique, Dynamic Linear Deflation. This cuff deflation technique allows a Colin monitor to measure each small change in the cuff pressure oscillations that directly correspond to the measurement's systolic, mean and diastolic blood pressure values.

The cuff is first increased in pressure until it reaches a pressure above arterial occlusion. As the cuff starts to deflate, the pulse rate of the patient is determined and the deflation speed of the cuff is modified to create a patient specific deflation speed. As the pressure decreases, small cuff pressure oscillations are recorded that correspond to the applied

pressure of the blood under the cuff as the heart contracts. These oscillations increase in strength as the cuff pressure approaches the systolic blood pressure value. A sudden increase in oscillation amplitude indicates that the patient's systolic blood pressure is now able to push blood completely through beneath the cuff. The oscillation amplitude continues to increase as the pressure in the cuff is decreases until the mean blood pressure value is reached. The oscillation strength then starts to diminish and finally drop off as the diastolic blood pressure value is reached.



<< From MEASUREMENT OF BLOOD PRESSURE by L.A.GEDDES >>

The oscillometric method does not determine an instantaneous blood pressure reading like the auscultatory method employing a microphone-type auto blood pressure monitor but, as described above, determines blood pressure from an uninterrupted changing curve, which means that the oscillometric method is not easily effected by external noise and electrosurgical instruments.

Note: This equipment is suitable for use in the presence of electro-surgery.

Setup Connections

- 1. Measure the patient's limb and select a proper size cuff. As a general rule, cuff width should span approximately two-thirds of the distance between the patient's elbow and shoulder.
- 2. Connect the cuff hose to the connector on the monitor's left panel and turn to right to lock. (see Figure 3).
- 3. Connect a cuff to the cuff hose and turn the connector to right to lock the hoses together. Firm connection must be made.



- 4. Wrap the cuff around a hare arm or around an arm covered in thin clothing. Thick clothing or a rolled up sleeve will cause a major discrepancy in the blood pressure reading.
- 5. Warp the cuff around the patient's arm so that the center of the cuff's rubber bladder sits on the artery of the upper arm. The hose should be brought out from the peripheral side without bending (The Brachial artery is located on the inside of the patient's upper arm.) At this time, check that the index line on the edge of the cuff sits inside the range. Use a different sized cuff if the index line is outside of the range because this will cause a major discrepancy in blood pressure reading.



- 6. The adult cuff should be wrapped around the arm tightly enough so that only two fingers can be inserted under it, above and below the cuff.
- 7. Maintain the height of the cuff-wrapped upper arm artery to that of the heart's right ventricle during measurement.
- 8. Follow the cuff directions for use when applying the cuff to the arm.

	Model Number	Cuff width (cm)	Arm circumference (cm)
Pediatric	Cuff No.1	7	12 to 18
	Cuff No.2	9	17 to 23
Adult	Cuff No.3	12	23 to 33
	Cuff No.4	14	31 to 40
Neonate	Cuff No.10	2.5	3.5 to 6
	Cuff No.11	3	5.0 to 7.5
	Cuff No.12	4	7.5 to 10.5
	Cuff No.13	5	8.5 to 13

Table 24. Cuff Size

NIBP Measurement Modes

Blood pressure measurements can be made in three modes:

- MANUAL mode: Single measurement of systolic/diastolic/mean arterial pressure.
- Automatic (AUTO) mode: Measurements at preset intervals.
- Continuous (CONT) mode: As many measurements as possible within a 5 minutes period.

To Initiate MANUAL Measurement Mode

1. Press the NIBP Start/Stop button.

A single blood pressure measurement will be made. The measurement will be displayed for 180 minutes unless another measurement is initiated. A manual NIBP reading can be obtained in AUTO mode by pressing **NIBP start/stop button** between two AUTO measurements without the cancellation of AUTO mode.

To Initiate Automatic (AUTO) Measurement Mode

 Press NIBP interval button to select the desired automatic mode interval from NIBP interval setting menu (see Table 25). The initial measurement will start automatically in a selected interval. The automatic mode can be also activated by pressing NIBP Start/Stop button after selecting the interval.

✓ NIBP Auto Mode intervals

: Off, Cont, 1, 2, 2.5, 3, 5, 10, 15, 20, 30, 45, 60, 90, 120, 180 minutes

- 2. An NIBP reading can be cancelled by pressing *NIBP Start/Stop button* during the AUTO measurements.
- Note: When the time interval is set to 1 minute, the initial measurement will automatically start after 5 seconds, and then the measurement interval will automatically become 2.5 minutes after 12 minutes elapsed.

The NIBP numerical area will display the *NIBP auto mode interval* and *NIBP elapsed time icon*. The interval is the time from when one measurement starts to when the next measurement starts. The measurement value will be displayed until another measurement starts. When the AUTO mode is cancelled, the last measurement will be displayed for 180 minutes.

In AUTO mode, the monitor attempts to meet the requirement of SVRP (Safe Venous Return Pressure) as long as starting a new reading does not violate the requirement of being 30 seconds below SVRP between readings. A new blood pressure reading will not start until the 30 second period has elapsed. When CONT and 1 minute is selected in *NIBP interval setting menu*, this SVRP can be shorten over 2 seconds since CONT is the intensive measurement during the short term which is 5 minutes in the BP-S510.

To Initiate Continuous (CONT) Measurement Mode

You may select *Cont* to activate the continuous measurement mode. The initial measurement will automatically starts in 5 seconds. The measurement interval will automatically become 2.5 minutes after 5 minutes elapsed. Also, if the *NIBP start/stop button* is pressed during CONT mode, the measurement will be canceled and the interval will be changed to 2.5 minutes.

To Stop Blood Pressure Measurements

You may press **NIBP Start/Stop button** at any time to stop the current measurement and deflate the cuff. If an automatic measurement is underway, next measurement will start at the next interval after the current measurement stops.

Description of NIBP Menu Functions



- 1 NIIBP icon
- 2 Systolic pressure icon
- 3 Systolic alarm icon
- 4 Systolic alarm limits value
- 5 MAP icon
- 6 Systolic pressure value
- 7 MAP value
- 8 MAP alarm icon

- 9 MAP alarm limits value
- 10 Diastolic pressure value
- 11 Diastolic pressure alarm limits value
- 12 Diastolic pressure alarm icon
- 13 Diastolic pressure icon
- 14 NIBP elapsed time
- 15 NIBP auto mode interval
- 16 NIBP unit

Figure 27. NIBP display



Figure 28. NIBP menu

	Table 25. NIBP Menu
Level 1 Menu	Level 2 Menu or Response
NIBP MENU	
Initial Inflation Pressure	120, 140, 160, 180, 200, 220 (mmHg) (Adult)
	80, 100, 120, 140 (mmHg) (Neonatal)
BP On alarm	On, Off
Smart Clock	On, Off
Completion Sound	On, Off
Smart Inflation	On, Off
(Alarm Limits Adjustment)	
	Upper Alarm Limit
▼	Lower Alarm Limit
NIBP Alarm Suspend	On, Off
Return	

Note: Initial Inflation Pressures shown above are for Adult patient mode. In order to set alarm limits to Neonatal mode, change Patient mode via **Setup Menu**.

Note: The NIBP unit can only be changed by authorized personnel via Service menu.

Initial Inflation Pressure

The inflating pressure can be set from 120 to 220 mmHg for adult or from 80 to 140 mmHg for neonatal.

BP on Alarm

If the BP on Alarm is **On**, the monitor will automatically take a measurement when an alarm condition occur.

Smart Clock

If the smart clock is **On**, the start of measurements will synchronize to the time. For example, after a measurement made at 10:03 with five-minute interval and then the smart clock set to On, next measurements will start at 10:05, 10:10 and another.

Completion Sound

When the completion sound is **On**, the monitor sounds beep tones to notify the completion of the NIBP measurement.

Smart Inflation

When the smart inflation is **On**, the suitable inflation value to the patient is automatically calculated during the inflation. The inflation value is automatically calculated around the expected systolic BP value + 45 mmHg.

SpO₂ MONITORING

	WARNING: For best product performance and measurement accuracy, use only		
	accessories manufactured by Tyco Healthcare Inc. Use accessories according to		
	the manufacturer's directions for use and your facility's standards.		
	WARNING: Tissue damage can be caused by incorrect application or use of an SpO ₂ sensor. Harm can be caused, for example, by wrapping the sensor too tightly, by applying supplemental tape, or by leaving a sensor on too long in one place. Inspect the sensor site as directed in the sensor directions for use to ensure skin integrity, correct positioning, and adhesion of the sensor.		
	WARNING: Do not use damaged SpO ₂ sensors. Do not use an SpO ₂ sensor with exposed optical components. Do not immerse sensor completely in water, solvents, or cleaning solutions because the sensor and connectors are not waterproof. Do not sterilize SpO ₂ sensors by irradiation, steam or ethylene oxide. Refer to the cleaning instructions in the directions for use for reusable SpO ₂ sensors.		
	WARNING: Inaccurate measurements may be caused by:		
	 incorrect sensor application or use 		
	 significant levels of dysfunctional hemoglobin 		
	(such as carboxyhemoglobin or methemoglobin)		
	 intravascular dyes such as indocyanine green or methylene blue 		
	 exposure to excessive illumination, such as surgical lamps (especially ones with a xenon light source), bilirubin lamps, fluorescent lights, infrared heating lamps, or direct sunlight 		
	• excessive patient movement		
	 high-frequency electrosurgical interference and defibrillators 		
	 venous pulsations 		
	 placement of a sensor on an extremity with a blood pressure cuff, arterial catheter, or intravascular line 		
	• patient conditions such as hypotension, severe vasoconstriction, severe		
	anemia, hypothermia, cardiac arrest, or shock		
	 arterial occlusion proximal to the sensor 		
	 environmental conditions 		
	unspecified length of the extension cable		
	WARNING: Do not attach any cable to the sensor port connector that is intended for computer use.		
•	CAUTION: The sensor disconnect error message and associated alarm indicate the sensor is either disconnected or the wiring is faulty. Check the sensor connection and, if necessary, replace the sensor, extension cable or both.		



CAUTION: Reusable sensors may be used on the same site for a maximum of 4 hours, provided the site is inspected routinely to ensure skin integrity and correct positioning.

General

The monitor uses pulse oximetry to measure functional oxygen saturation in the blood. Because a measurement of SpO₂ is dependent upon light from the SpO₂ sensor, excessive ambient light can interfere with this measurement. SpO₂ and Pulse rate is updated every second. This monitor measures functional saturation - oxygenated hemoglobin expressed as a percentage of the hemoglobin that can transport oxygen. It does not detect significant amounts of dysfunctional hemoglobin, such as carboxyhemoglobin or mehtemoglobin.

Functional versus Fractional Saturation

This monitor measures functional saturation — oxygenated hemoglobin expressed as a percentage of the hemoglobin that can transport oxygen. It does not detect significant amounts of dysfunctional hemoglobin, such as carboxyhemoglobin or methemoglobin. In contrast, hemoximeters such as the IL482 report fractional saturation — oxygenated hemoglobin expressed as a percentage of all measured hemoglobin, including measured dysfunctional hemoglobin. To compare functional saturation measurements to those from an instrument that measures fractional saturation, fractional measurements must be converted as follows:

functional saturation =	fractional saturation	~ 100
	100 – (%carboxyhemoglobin + %methemoglobin)	~ 100

Measured versus Calculated Saturation

When saturation is calculated from a blood gas partial pressure of oxygen (PO2), the calculated value may differ from the SpO_2 measurement of the monitor. This usually occurs because the calculated saturation was not appropriately corrected for the effects of variables that shift the relationship between PO2 and pH, temperature, the partial pressure of carbon dioxide (PCO₂), 2, 3-DPG, and fetal hemoglobin.

Automatic Calibration

Because light absorption by hemoglobin is wavelength dependent and because the mean wavelength of LEDs varies, an oximeter must know the mean wavelength of the *OXIMAX* sensor's red LED to accurately measure SpO₂. The wavelength range of the light emitted are near 660 nm and 890 nm with the energy not exceeding 15 mW. During monitoring, the instrument's software selects coefficients that are appropriate for the wavelength of that individual sensor's red LED; these coefficients are then used to determine SpO₂. Additionally, to compensate for differences in tissue thickness, the light intensity of the sensor's LEDs is adjusted automatically.

Setup Connections

When selecting a sensor, consider the patient's weight and activity, adequacy of perfusion, availability of sensor sites, need for sterility, and anticipated duration of monitoring. Refer to Table 26, or contact Tyco Healthcare Inc. sales department for ordering information.

- 1. Select the proper sensor for the patient.
- 2. Connect the extension cable to the SpO₂ connector on the monitor's left panel and lock it. (see Figure 3)
- 3. Connect the sensor to the extension cable and lock it.
- 4. Carefully apply the sensor to the patient, as described in the sensor directions for use. Observe all warnings and cautions in the directions for use.



- Note: Refer to directions for use to make sure the proper placement for various types of SpO₂ sensor.
- Note: Periodically check to see that the sensor remains properly positioned on the patient and that skin integrity is acceptable. Refer to the sensor directions for use.

Sensor	Model	Patient Size
OXIMAX oxygen transducer (Sterile, single-use	MAX-N	<3 or >40 kg
only)	MAX-I	3 to 20 kg
	MAX-P	10 to 50 kg
	MAX-A	>30 kg
	MAX-AL	>30 kg
	MAX-R	>50 kg
OXIMAX Oxiband ® oxygen transducer	OXI-A/N	<3 or >40 kg
(Reusable with disposable non-sterile adhesive)	OXI-P/I	3 to 40 kg
OXIMAX Durasensor ® Oxygen transducer	DS-100A	>40 kg
(Reusable, non-sterile)		
OXIMAX OxiCliq ® oxygen transducers	Р	10 to 50 kg
(Sterile, single-use only)	N	<3 or >40 kg
	1	3 to 20 kg
	А	>30 kg
OXIMAX Dura-Y ® multisite oxygen transducer	D-YS	>1 kg
(Reusable, non-sterile)		
For use with the Dura-Y sensor:		
Ear clip (Reusable, non-sterile)	D-YSE	>30 kg
Pedi-Check TM pediatric spot-check clip		
(Reusable, non-sterile)	D-YSPD	3 to 40 kg
OXIMAX MAX-FAST adhesive reflectance	MAX-FAST	>40 kg
oxygen transducer		

Table 26. SpO₂ Sensors

Pulse Amplitude Indicator

The pulse amplitude indicator is the segmented display within the SpO₂ numerical area that shows the relative strength of the detected pulse. A stronger pulse causes a larger amplitude indicator.

Description of SpO2 Menu Functions



Level 1 Menu	Level 2 Menu or Response
SpO ₂ MENU	
C-Lock	On, Off
(Alarm Limits Adjustment)	
	Upper Alarm Limit
▼	Lower Alarm Limit
SpO ₂ Alarm Suspend	On, Off
Return	

C-Lock

When C-Lock is turned on in the SpO₂ menu, C-Lock automatically becomes operational any time a valid ECG signal is detected by the monitor. It is not necessary to turn C-Lock off if an ECG signal is not available; the monitor handles this function automatically. If the ECG signal is noisy, or of poor quality, SpO₂ performance may be improved by turning C-Lock off. C-Lock provides ECG synchronization for more reliable saturation measurements. An ECG (R-wave) signal can be used as a time reference to identify the pulse and synchronize saturation measurements. C-Lock enhances performance while maintaining rapid response time.

Description of Pleth Waveform Menu Functions



Figure 32. Pleth Waveform Menu

Table 28.	Pleth	Waveform	Menu
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Level 1 Menu	Level 2 Menu or Response	
PLETH WAVEFORM MENU		
Sweep Speed	12.5 mm/s, 25.0 mm/s, 50.0 mm/s	
Waveform Select	ECG, Pleth, Respiration, (P1 Label), (P2 Label), Capno	
Return		

Sweep Speed

The user-selectable Sweep Speed determines the speed at which pleth waveform trace moves across the screen. *Sweep Speed* can be selected from 12.5 mm/s, 25.0 mm/s and 50.0 mm/s, and Pleth waveform is synchronized with ECG waveform and IBP waveform.

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RESPIRATION MONITORING

WARNING: For best product performance and measurement accuracy, use only accessories supplied or recommended by Colin Medical Technology. Use accessories according to the manufacturer's directions for use and your facility's standards.
WARNINGS: The monitor does not detect apnea when the respiration signal is measured by trans-thoracic impedance.
WARNING: Keep patients under close surveillance when monitoring respiration. Respiration signals are relatively more sensitive to interference from radiated electromagnetic signals. Thus, it is possible, although unlikely, that radiated electromagnetic signals from sources external to the patient and monitor can cause inaccurate respiration readings. Do not rely entirely on the monitor respiration readings for patient assessment. If measured waveforms are not appropriate readings, check external condition to ensure there is no equipment to affect electromagnetic interference.

General

The impedance respiration measurement uses the impedance between ECG electrodes. Human respiration takes place by chest expansion by the respiratory muscle. As the chest expands in the inspiratory movement, the impedance between the ECG electrodes will change. The monitor detects respiration rate by putting high-frequency current between RA and LA of the ECG electrodes.

The airway respiration measurement uses gases coming into the airway adapter in case of an $EtCO_2$ option equipped. The monitor detects respiration rate by computing each breath cycle from continuous capno waveform.

Setup Connections

Refer to the **ECG Monitoring** section to acquire the respiration signal by the patient impedance using the ECG electrodes, leads and cable.

The performance of the impedance respiration can be improved by the particular placement of the Left arm (LA) and Right arm (RA) electrodes. (See **Standard ECG** *electrode placement* in Figure 21)

Refer to the **Capnography Monitoring** section to detect the respiration signal by the airway adapter in case that an EtCO2 option equipped.

Description of Respiration Menu Functions



- 1 Respiration icon 2
- 4 Respiration alarm limits value Respiration alarm icon
- 5 Respiration source icon
- 3 Respiration value





Figure 34. Respiration Menu

Table 29. Respiration Menu

Level 1 Menu	Level 2 Menu or Response
RESPIRATION MENU	
Respiration/Apnea	Off, Auto (awRR>imRR), awRR, imRR
Apnea Time Setting	Off, 20, 30, 40, 60, Step 60, Step 90
(Alarm Limits Adjustment)	
	Upper Alarm Limit
▼	Lower Alarm Limit
Respiration Alarm Suspend	On, Off
Return	

Respiration / Apnea Source

You can select either awRR or imRR for the source of the respiration rate. If you select Auto, the monitor will automatically derive the respiration rate from one of the monitoring parameters in this order of priority; awRR and imRR.

Note: You can select awRR as the source when CO_2 module is installed.

Apnea Time Setting

When the patient's breath is not detected from the airway measurement for a selected time setting, the monitor will activate an apnea alarm. When *Step 60* is selected, the monitor will sound two beeps in 20 second interval from no breath. At 60 seconds elapsed, an apnea alarm will be activated. When *Step 90* is selected, the monitor will sound three beeps in 30 second interval from no breath. At 90 seconds elapsed, an apnea alarm will be activated, the monitor does not detect an apnea alarm.

When the monitor does not detect a respiration signal from the impedance measurement for 40 seconds, the monitor will activate a loss of respiration alarm.

Check the condition of the patient, then check the connections of the patient cables.

Description of Respiration Waveform Menu Functions



1 Impedance respiration 2 Respiration waveform waveform icon





Figure 36. Respiration Waveform Menu

Level 1 Menu	Level 2 Menu or Response	
RESPIRATION WAVEFORM MENU		
Sweep Speed	6.25 mm/s, 12.5 mm/s, 25.0 mm/s	
Size	×1/4, ×1/2, ×1, ×1.5, ×2 (5 steps)	
Waveform Select	ECG, Pleth, Respiration, (P1 Label), (P2 Label) Capno	
Return		

Sweep Speed

The user-selectable sweep speed determines the speed at which the respiration waveform trace moves across the screen. *Sweep Speed* can be selected from 6.25 mm/s, 12.5 mm/s and 25.0 mm/s, and the respiration waveform is synchronized with Capno waveform.

Size

Size allows you to adjust the waveform size. (5 steps: ×1/4, ×1/2, ×1, ×1.5, ×2)
TEMPERATURE MONITORING

WARNING: For best product performance and measurement accuracy, use only YSI 400 series temperature probes recommended by Colin Medical Technology. Use accessories according to the manufacturer's directions for use and your facility's standards.

General

Measurement of patient temperature is accomplished by processing the signal from a probe containing a resistance element whose impedance is temperature dependent. These devices are called thermistors. The measuring time required to obtain accurate readings at the specific body site is about 10 seconds.

Setup Connections

The monitor is designed to accept signals from the temperature probes, YSI 400 series for skin, rectal or etc. Refer to the temperature probe directions for use for details

- 1. Insert a body temperature probe into the temperature connector on the monitor's left panel (see Figure 3).
- 2. Follow the directions for use accompanying the temperature probe.

Description of Temperature Menu Functions





Figure 38. Temperature Menu

Table	31.	Temperature	Menu
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Level 1 Menu	Level 2 Menu or Response
TEMPERATURE MENU	
(T1-Alarm Limits Adjustment)	
	Upper Alarm Limit
▼	Lower Alarm Limit
T1 Alarm Suspend	On, Off
(T2-Alarm Limits Adjustment)	
	Upper Alarm Limit
▼	Lower Alarm Limit
T2 Alarm Suspend	On, Off
Return	

Note: The temperature unit can only be changed by authorized personnel via *Service menu*.

IBP MONITORING

	WARNING: Proper measurements may not be possible,
	 if improper zero calibration was performed.
	 if air bubbles are mixed into the patient circuit.
	 if the height of the three-way tap for zero calibration and the right auricle have changed.
	WARNING: If the pressure transducer has been dropped or subjected to strong
	physical shock, check for faults before use.
	WARNING: Be sure to thoroughly read the instruction manuals for each item-such
	as the transducer, monitoring kit and transfusion set-that are used in invasive
	blood pressure measurements before using them. The cautions and warnings for
	such items are not included in this manual.
	WARNING: Use a CE certified transducer with a sensitivity of 5µV/V/mmHg at an
	excitation voltage of 5VDC, and a measurement range of -50 to 300mmHg.
	WARNING: Do not reuse disposable (single use) transducers.
	WARNING: Ensure that reusable transducers are sufficiently sterilized.
	WARNING: To ensure patient safety, do not contact any conductive parts to the
	applied part.
	WARNING: Never install or remove the IBP module while the monitor is powered
	on. It is likely to break when installing or removing it while powered on.
-	

General

The invasive blood pressure measurement measures the systolic pressure, mean pressure, diastolic pressure and pulse rate for up to 2 blood pressure line channels using blood pressure transducers, and displays the blood pressure waveform.

Installing an IBP module



Figure 39. IBP Module Installation

- 1. Turn off the monitor.
- 2. Push the IBP module into the main unit until click.

Setup Connections

- 1. Connect the interface cable for the transducer to the IBP connector on the monitor's left panel. An interface cable for the transducer has to be selected correctly as it depends on the each transducer type. (see Figure 3)
- 2. Set up the patient circuit according to the directions for use of the transducer, monitoring kit and IV set.

The drawing below shows the example.





Figure 41. IBP Menu

Table	32	IBP 1	Menu
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Level 1 Menu	Level 2 Menu or Response
(P1 Label) or (P2 Label) MENU	
P1 Label	P1, ABP
Pressure Zero Setting	No, Yes
(SYS Alarm Limits Adjustment)	
	Upper Alarm Limit
▼	Lower Alarm Limit
(MEAN Alarm Limits Adjustment)	
	Upper Alarm Limit
▼	Lower Alarm Limit
(DIA Alarm Limits Adjustment)	
	Upper Alarm Limit
▼	Lower Alarm Limit
(P1 Label) Alarm Suspend	On, Off
Return	

Table 33. IBP 2 Menu

Level 1 Menu	Level 2 Menu or Response		
(P2 Label) MENU			
P2 Label	P2, CVP, PAP, LAP		
Pressure Zero Setting	No, Yes		
(SYS Alarm Limits Adjustment)			
	Upper Alarm Limit		
▼	Lower Alarm Limit		
(MEAN Alarm Limits Adjustment)			
	Upper Alarm Limit		
▼	Lower Alarm Limit		
(DIA Alarm Limits Adjustment)			
	Upper Alarm Limit		
▼	Lower Alarm Limit		
P2 Alarm Suspend	On, Off		
Return			

Note: The IBP unit can only be changed by authorized personnel via Service menu.

Label

Name of each channel (P1 or P2) can be selected in this menu.

Pressure Zero Setting

- 1. Three way tap for zero calibration should be set at the same level as patient's heart.
 - ✓ Make sure if there is no bubble in the each part.
 - ✓ Make sure if the drop volume of the IV set is at the proper position.
- 2. The handle of the three way tap should be to the patient so that the pressure to the transducer is open to the atmosphere level.
 - \checkmark Please make sure if release part to the atmosphere in not covered by cap.
- 3. Set the pressure zero setting to Yes. "Zero CAL in progress" message will appear.
- 4. When the message "Zero CAL in progress" is disappeared and measurement will be initiated.
- 5. When Zero Calibration is done, the handle of three way tap should be turned to the atmosphere direction and the release part should be covered by the cap.
- 6. IBP measurement can start by changing the lever of three way tap so as to transfer the pressure from the catheter to transducer.
- Note: If there is bubble in the any part, you can remove the bubble by controlling the transducer.
- Note: If zero calibration can not be done correctly, the readings will not be accurate.
- Note: When zero calibration is done, make sure that release port of three way tap for Zero calibration is open, not covered by the cap of injector.

Description of IBP Waveform Menu Functions





Figure 43. IBP Waveform Menu

Table 34.	(P1 Label)) Waveform	Menu
-----------	------------	------------	------

Level 1 Menu	Level 2 Menu or Response	
(P1 Label) WAVEFORM MENU		
Sweep Speed	12.5 mm/s, 25.0 mm/s, 50.0 mm/s	
P1 Label	P1, ABP	
P1 Scale	0~50, 0~100, 0~200, 0~300 mmHg, Auto	
Pressure Zero Setting	No, Yes	
Waveform Select	ECG, Pleth, Respiration, (P1 Label), (P2 Label), Capno	
Return		

Table 35. (P2 Label) Waveform Menu

Level 1 Menu	Level 2 Menu or Response	
(P2 Label) WAVEFORM MENU		
Sweep Speed	12.5 mm/s, 25.0 mm/s, 50.0 mm/s	
P2 Label	P2, CVP, PAP, LAP	
P2 Scale	0~20, 0~50, 0~100, 0~200, 0~300 mmHg, Auto	
Pressure Zero Setting	No, Yes	
Waveform Select	ECG, Pleth, Respiration, (P1 Label), (P2 Label), Capno	
Return		

Sweep Speed

The user-selectable sweep speed determines the speed at which IBP waveform trace moves across the screen. *Sweep Speed* can be selected from 12.5 mm/s, 25.0 mm/s and 50 mm/s, and the IBP waveform is synchronized with ECG waveform and Pleth waveform.

Scale

The use can select the scale of P1 and P2 in the IBP waveform menu. When Auto is selected, the monitor automatically set the scale upon the IBP measurement value.

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CAPNOGRAPHY MONITORING

Make sure that there is no leakage in the tubes connected to the patient. If there is leakage, external air will be sucked in and the sampling gas will be diluted, causing measurement values to drop.
Make sure that selection of a volatile anesthetic is done carefully. If an improper selection is made, the measurement values will be erroneous.
Do not use consumables other than those which are specified. Different sampling tube lengths or inside diameters may have an effect on the measurement values.
If the patient's breathing becomes fast, the dead space inside the sampling circuit or device, etc. is causing the response to drop, so correct measurements cannot be taken. Use it together with close observation of the capnogram. If the sampling flow rate is 65 ml/min, the response will drop still lower, so particular caution is necessary.
For the same reason as stated above, in the case of newborn infants and other patients who have a low air exchange rate per minute, correct measurements cannot be taken.
Do not use device without rectifying a gas calibration failure, as correct measurement readings will not be obtained.
The monitor is not use for in breathing system and inhalation anaesthetic agents

General

The monitor provides the continuous end-tidal carbon dioxide (EtCO₂), inspired carbon dioxide (InCO₂), and respiration rate by using a side stream type gas module which uses a suction pump to sample the gases in the patient's inhaled and exhaled breath. CO_2 has the characteristic of absorbing infrared light, so the concentration of the gas is measured by the amount of infrared absorption. The sampled gases are analyzed using an internal sensor. The inhalation and exhalation gas concentrations are measured respectively. The respiration rate is measured from the changes of this curve.

Setup Connections

CAUTION: Connect the sampling tube to the monitor before turning power ON. Failure to do so will cause the error message to be displayed.

• CAUTION: Firmly tighten each section when connecting because external air will be sucked in at any leaking connection, which will cause sample gas dilution lowering the measurement reading.



CAUTION: Confirm the connector on the airway adaptor for the Nafion tube is facing up to prevent liquid from flowing in.

 CAUTION: The airway adaptor, Nafion tube and sampling tube are all consumables. It is recommended that they be replaced for each new patient. (Definitely change these components if the last patient having been examined had an infectious disease.) CAUTION: Be sure to dispose of discharged gas in order to prevent pollution of treatment room.

CAUTION: Do not reuse discharged gas in order to prevent the spread of infectious diseases.

1. Connect the sample line in the following manner:



2. Check the volume of water in the water trap on the monitor's left panel.

Warming Up

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The monitor requires maximum of 3 minutes to warm up after the power has been turned on. After the warm-up has finished, auto atmosphere calibration will commence.

Sample Line Checks

WARNING: Do not use the device if the "Occlusion" message has not displayed after the sample line was seal-checked, as measurement readings will be incorrect.
 Check to see that the sample gas is not diluted. If gas is diluted, measurement readings will be affected due to intake of external air from some point.
 Seal both ends of the airway adaptor (see diagram below).
 After a few seconds, check to see that the "Occlusion" message appears on screen and then unseal the ends of the airway adaptor.
 Press the jog dial to return to normal measurement.
 If the "Occlusion" message does not appear, there is probably a loose connection or leakage, so check each part of the setup.
 Key points of attention:
 Coclude the inlet port of the water trap to see which side of the patient circuit may be the problem

 \checkmark Also replace components to see if that solves the problem.

Description of CO₂ Menu Functions



Table 36. CO₂ Menu

Level 1 Menu	Level 2 Menu or Response	
CO ₂ MENU		
Capno Measurement	On, Off	
(InCO ₂ Alarm Limits Adjustment)		
	Upper Alarm Limit	
▼	Lower Alarm Limit	
InCO ₂ Alarm Suspend	On, Off	
(EtCO ₂ Alarm Limits Adjustment)		
	Upper Alarm Limit	
▼	Lower Alarm Limit	
EtCO ₂ Alarm Suspend	On, Off	
Return		

Note: The EtCO₂ and InCO₂ unit can only be changed by authorized personnel via *Service menu*.

Capno Measurement

When the capno measurement is On, EtCO₂, InCO₂ and capno waveform measurement are activated.

Description of Capno Waveform Menu Functions



- 1 Capno waveform icon 3 Capno waveform
- 2 Capno scale bar

Figure 47. Capno Waveform Display



Figure 48. Capno Waveform Menu

Table 37. C	Capno Wav	eform Menu
-------------	-----------	------------

Level 1 Menu	Level 2 Menu or Response		
CAPNO WAVEFORM MENU			
Sweep Speed	6.25 mm/s, 12.5 mm/s, 25 mm/s		
Scale	Auto, 0~40, 0~60, 0~80		
Calibration	Yes, No		
Waveform Select	ECG, Pleth, Respiration, (P1 Label), (P2 Label), Capno		
Return			

Sweep Speed

The user-selectable sweep speed determines the speed at which Capno waveform trace moves across the screen. *Sweep Speed* can be selected from 6.25 mm/s, 12.5 mm/s and 25.0 mm/s, and the respiration waveform is synchronized with Respiration waveform.

Scale

The user can select the scale of capno waveform. When Auto is selected, the monitor will automatically set the scale upon the measurement value.

Calibration

Check gas concentration once a month or when the measurement reading seems dubious to ensure reliable measurement readings.

The selections required for this check are the optional designated calibration gas and pressure adjustment regulator.

1. Warming Up

The gas monitor's internal temperature must stabilize to ensure stable performance of the monitor. Therefore, wait for minimum 3 minutes to perform a gas calibration after turn on the power of the monitor.



 \bigcirc

 \bigcirc

CAUTION: When the calibration is not performed by the instructions stated in this section, the monitor could cause inaccurate measurement readings.

CAUTION: Perform the concentration check in an environment where the room temperature is 15°C to 27 °C to enhance calibration accuracy.

CAUTION: Move on to the next procedure when the "Warming up in progress" message disappears. If this message remains displayed, the sample line check cannot be performed and the gas calibration mode cannot be selected.

2. Calibration Gas Preparation

Prepare the optional designated calibration gas and the flowmeter, pressure adjustment instrument. Thoroughly check the component ratio of the calibration gas, as there are numerous types on the market.

CAUTION:

Calibration Gas Usage Cautions

- Calibration gas must be used at between 15 °C to 27 °C (60 °F to 80 °F).
- The expiry date (EXP. DATE:) must printed on the calibration gas bottle. Do
 not use a calibration gas that has passed expiry date, as accuracy cannot be
 assured.
- When using calibration gas, be sure to use a surplus gas discharge device or thoroughly ventilate installation site.
- Do not breathe gas.
- Use with equipment rated for cylinder pressure.

Calibration Gas Storage Cautions

- Calibration gas must be stored at between 15 °C to 27 °C (60 °F to 80 °F).
- Exposure to temperature above 52 °C (125 °F) may cause contents to vent or cause bursting.
- Keep away from heat, including the sun, flame and sparks.
- Never discard container into fire or incinerator.

3. Sample Line Checks

Check to see that the sample gas is not diluted. If gas is diluted, measurement readings will be affected due to intake of external air from some point. Refer to **Sample Line Checks** paragraph.

4. Connecting Calibration Gas

Connect the calibration gas cylinder and the flowmeter instead of the airway adapter to the sampling tube connected to the water trap

5. Accessing Calibration

Set *Calibration* in the capno waveform menu to **Yes** using the jog dial. The message "Calibration in progress" will appear.

6. Commencing Calibration Gas Flow

Once zero calibration has ended successfully, the message "Turn on CO_2 (10%) calibration gas." will appear. Commence calibration gas flow by turning the knob of the flowmeter. The flow rate should be between 130~170 ml/min. The message "Calibration gas detected" will appear.

7. Ending Calibration Gas

Once sampling is completed, the message "Please turn off CO_2 (10%) calibration gas" will appear. Turn off the calibration gas by turning the knob of the flowmeter.

If Gas Calibration Fails

If gas calibration fails while in progress, a screen like the following message will be displayed.

- ✓ Zero calibration range error
- ✓ Zero calibration signal unstable error.
- ✓ High calibration range error.
- ✓ High calibration signal unstable error.

In this case, gas calibration must be performed again, but before doing so, check the following selections.

- ✓ Is the correct calibration gas being used?
- ✓ Are connections loose?
- ✓ Is the calibration gas supply flow too low?
- ✓ Is the calibration gas supply flow too high?

TRENDS

General

Trend data in either graphical or tabular format may be displayed or printed if a recorder is installed. (see **Using the monitoring** section)

- 1. Press *Trend button* to display the tabular trends.
- 2. Press *Trend button* once again to display the graphical trends.
- 3. Press Trend button once again, or press Home button to return to the main screen

The trend data is stored in a memory. When the monitor turns on and starts to measure vital signs, the monitor saves data at a selected interval. Also, the monitor saves all physiological alarm conditions, NIBP measurement and error event. After the monitor has stored 1500 trend data, the monitor begins to store the new data over the oldest data.

Tabular Trend Data

The monitor presents trend information in tabular format for all monitored parameters. The newest data appears at the bottom of tabular trends. Use the scroll function to view more data. The gray bar at the right side of the trend screen presents the memory saved. The red point indicates the location of currently scrolling.



For the NIBP trends, the data may display the latest measurement.

Figure 49. Tabular Trend Screen

To scroll or change viewing options, push the jog dial on the tabular trend screen when the tabular trend screen is highlighted. The monitor displays the *Tabular trend menu*.



Figure 50. Tabular Trend Menu

Table 38. Tabular Trend Menu

Level 1 Menu	Level 2 Menu or Response
TABULAR TREND MENU	
Scroll	The jog dial is activated for scrolling.
	Note: Press the <i>Home button</i> to exit.
Save Time Interval	Off, 0.5, 1, 2, 2.5, 5, 10, 15, 20, 30, 60, 120 minutes
Trend Clear	Yes, No
Return	Exits Tabular trend menu immediately, returns to
	Tabular Trend Screen

Scrolling Tabular Trend Data

- 1. Rotate the jog dial to highlight Scroll.
- 2. Press the jog dial to activate scrolling.
- 3. Rotate the jog dial to scroll through the trend data. Clockwise rotation moves forward to newer data. Counter clockwise rotation moves backward to older data.
- 4. After viewing the trends, press the jog dial to exit the scrolling.

Setting Save Time Interval

The monitor saves the trend data at the interval selected by user, and it displays all data stored when the tabular trend screen is activated. When **Off** is selected, the monitor stores trend data only when the NIBP measurement and the alarm condition occur.

Trend Clear

To clear trend data in the trend memory, set Trend Clear to Yes.

Graphical Trend Data

Trend information in graphical format for all monitored parameters is displayed in one graph. The user can select each parameter to display via interaction with *Graphical trend menu*.

The graphical trend data of each parameter is indicated by the symbols specified in table 3. The vertical range of a graphical trend is presented with fixed value, and the horizontal range is 90 minutes. The newest data appears at the right of graphical trend. Use the scroll function to view more data.



Figure 51. Graphical Trend Screen

To scroll or change viewing options, push the jog dial on the graphical trend screen when the graphical trend screen is highlighted. The monitor displays *Graphical trend menu*.



Figure 52. Graphical Trend Menu

Level 1 Menu	Level 2 Menu or Response
GRAPHICAL TREND MENU	
Scroll	The jog dial is activated for scrolling.
	Note: Press the <i>Home button</i> to exit.
HR/PR	On, Off
NIBP	On, Off
(PI Label)	On, Off
(P2 Label)	On, Off
SpO ₂	On, Off
RESP	On, Off
EtCO ₂	On, Off
T1	On, Off
T2	On, Off
Trend clear	Yes, No
Return	Exits Graphical Trend Menu immediately,
	returns to Graphical Trend Screen

Table 39. Graphical Trend Menu

Selecting Graphical Trend Data

- 1. Rotate the jog dial to select *HR/PR*, *NIBP*, *IBP1*, *IBP2*, *SpO*₂, *RESP*, *EtCO*₂, *T1* or *T2*, *graphical trend*.
- 2. Press the jog dial to set to *On*.
- 3. Rotate the jog dial to highlight *Return*, then press the jog dial to return to the graphical trend screen. Only parameters set to *On* will be displayed in the graphical trend screen.

Note: Setting Off will not display the trends of the selected parameter.

Scrolling Graphical Trend Data

- 1. Rotate the jog dial to highlight Scroll.
- 2. Press the jog dial to activate scrolling.
- 3. Rotate the jog dial to scroll through the trend data. Clockwise rotation moves forward to newer data. Counterclockwise rotation moves backward to older data.
- 4. After viewing the trends, press the jog dial to exit scrolling.

Trend Clear

To clear trend data in the trend memory, set Trend Clear to Yes.

MENU STRUCTURE

ECG WAVEFORM MENU

ECG Cable Select		
-	-	3 Leads
-	-	5 Leads
-	-	AUTO
-	-	Return
-	Lead Se	lect
-	-	I
-	-	II
-	-	III
-	-	aVR
-	-	aVL
-	-	aVF
-	-	V (Chest Lead)
-	-	Return
-	Sweep S	Speed
-	-	12.5 mm/s
-	-	25.0 mm/s
-	-	50 mm/s
-	-	Return
-	Size	
-	-	▲ "Up"
-	-	▼ "Down"
-	-	Return
-	Pacer D	etect
-	-	On
-	-	Off
-	Filter Mo	ode
-	-	Monitor
-	-	Low Extend
-	-	Filter
-	-	Respiration Rejection
-	-	Return
-	Wavefor	m Select
-	-	ECG
-	-	Pleth
-	-	Respiration
-	-	(P1 Label)
-	-	(P2 Label)
-	-	
-	-	Return
-	Return	

PLETH WAVEFORM MENU

-	Sweep S	Speed
-	-	12.5 mm/s
-	-	25.0 mm/s
-	-	50.0 mm/s
-	-	Return
-	Wavefor	rm Select
-	-	ECG
-	-	Pleth
-	-	Respiration
-	-	(P1 Label)
-	-	(P2 Label)
-	-	Capno
-	-	Return
-	Return	

RESPIRATION WAVEFORM MENU

-	Sweep Speed	
-	-	6.25 mm/s
-	-	12.5 mm/s
-	-	25.0 mm/s
-	-	Return
-	Size	
-	-	▲ "Up"
-	-	▼ "Down"
-	-	Return
-	Wavefor	m Select
-	-	ECG
-	-	Pleth
-	-	Respiration
-	-	(P1 Label)
-	-	(P2 Label)
-	-	Capno
-	-	Return
-	Return	

••		
-	Sweep S	Speed
-	-	12.5 mm/s
-	-	25.0 mm/s
-	-	50.0 mm/s
-	-	Return
-	P1 Labe	ł
-	-	P1
-	-	ABP
-	-	Return
-	P1 Scale	9
-	-	0~50
-	-	0~100
-	-	0~200
-	-	0~300
-	-	AUTO
-	-	Return
-	Pressur	e Zero Setting
-	-	Yes
-	-	No
-	Wavefor	rm Select
-	-	ECG
-	-	Pleth
-	-	Respiration
-	-	(P1 Label)
-	-	(P2 Label)
-	-	Capno
-	-	Return
-	Return	

(P2 ARFI)	WAVEFORM	MENII

(Г 🚄		
-	Sweep	Speed
-	-	12.5 mm/s
-	-	25.0 mm/s
-	-	50.0 mm/s
-	-	Return
-	P2 Lab	el
-	-	P2
-	-	CVP
-	-	PAP
-	-	LAP
-	-	Return
-	P2 Sca	le
-	-	0~20
-	-	0~50
-	-	0~100
-	-	0~200
-	-	0~300
-	-	Auto
-	-	Return
-	Pressu	re Zero Setting
-	-	Yes
-	-	No
-	Wavefo	orm Select
-	-	ECG
-	-	Pleth
-	-	Respiration
-	-	(P1 Label)
-	-	(P2 Label)
-	-	Capno
-	-	Return
-	Return	

CAPNO WAVEFORM MENU

CAFINO		
-	Sweep S	Speed
-	-	6.25 mm/s
-	-	12.5 mm/s
-	-	25.0 mm/s
-	-	Return
-	Scale	
-	-	0~40
-	-	0~60
-	-	0~80
-	-	AUTO
-	-	Return
-	Calibrat	ion
-	-	Yes
-	-	No
-	-	Return
-	Wavefor	m Select
-	-	ECG
-	-	Pleth
-	-	Respiration
-	-	(P1 Label)
-	-	(P2 Label)
-	-	Capno
-	-	Return
-	Return	

Note: You can select the same waveform to display in two consecutive waveform areas.

HR/PR MENU

-	HR/PR Source	9
-	- AUTO	0 "ECG> IBP>SpO ₂ >NIBP"
-	- HR	"ECG"
-	- PR	"IBP>SpO₂>NIBP"
-	- Retu	rn -
	"Alarm limits a	djustment/Alarm suspend setting"

- HR/PR
- 180
- **40**
- -
- On
- 💥
- Off
- Return

	IENU			
-	Initial Inflation Pressure			
-	-	"Adul	t"	
-	-	120 n	nmHg	
-	-	140 n	nmHg	
-	-	160 n	nmHg	
-	-	180 n	nmHg	
-	-	200 n	nmHg	
-	-	220 n	nmHg	
-	-	"Neol	natal"	
-	-	80 m	mHg	
-	-	100 n	nmHg	
-	-	120 n	nmHg	
-	-	140 n	nmHg	
-	-	Retu	rn	
-	BP On	Alarm		
-	-	On		
-	- Ours a set			
-	Smart	CIOCK		
-	-	On		
-	- Compl	UII ation S	ound	
-	Compi	On On	ouna	
-	-	Off		
-	- Smart	Un Inflatio	n	
-	-	On		
-	-	Off		
	"Alarm	limits a	diustme	ent/Alarm suspend setting (mmHa)"
_	svs	ΜΔΡ		
	010			
	200	180	160	
	70	40	30	
-		Å		
	On	On	On	
-	×	×	X	
	Off	Off	Off	
-	Return	•		

- Automatic Mode Interval

-	Autor	matic Mode
-	-	Off
-	-	Cont
-	-	1 min
-	-	2 min
-	-	2.5 min
-	-	3 min
-	-	5 min
-	-	10 min
-	-	15 min
-	-	20 min
-	-	30 min
-	-	45 min
-	-	60 min
-	-	90 min
-	-	120 min
-	-	180 min
-	-	Return

SpO₂ MENU

- C-Lock
- - On
- - Off

"Alarm limits adjustment/Alarm suspend setting"

- SpO₂
- 100
- 90
- _
- • On
- 💥
- Off
- Return

(P1 LABEL) MENU

-	P1	Label
		D4

- --P1 --ABP
- -Return
- **Pressure Zero Setting**
- Yes -
- --No

"Alarm limits adjustment/Alarm suspend setting (mmHg)"

-	SYS	MEAN	DIA
	200	180	160
	70	40	30
-	Å		Å
	On	On	On
-	×	×	×
	Off	Off	Off
	Dotur	`	

-Return

(P2 LABEL) MENU

-	P2 La	bel		
-	-	P2		
-	-	CVP		
-	-	PAP		
-	-	LAP		
-	-	Return		
-	Press	ure Zero S	Setting	
-	-	Yes		
-	-	No		
	"Alarn	n limits adj	ustment	/Alarm suspend setting (mmHg)"
-	SYS	MEAN	DIA	
	200	180	160	
	70	40	30	
-	\		Å	

	-	•	•
	On	On	On
-	×	×	×
	Off	Off	Off
-	Return	1	

RESPIRATION MENU

- Respiration/Apnea
- - Off
- - AUTO
- - awRR - - imRR
- - Return
- Apnea Time Se
 - Apnea Time Setting "40 sec preset for loss of respiration"
- - Off
- - 20 sec
- - 30 sec
- - 40 sec
- - 60 sec
- - Step 60
- - Step 90
- Return

"Alarm limits adjustment/Alarm suspend setting"





- Return

CO₂ MENU

- Capno Measurement
- - On
- - Off

"Alarm limits adjustment/Alarm suspend setting (mmHg)"

InCO₂ EtCO₂

	20	80
	0	0
-		
	On	On
-	×	×
	Off	Off

- Return

TEMPERATURE MENU

"Alarm limits adjustment/Alarm suspend setting (°C)"

	T1	Т2
	38.0	38.0
	14.5	14.5
-	Å	
	On	On
	¥ .	¥

- A A Off Off
- Off

TABULAR TREND MENU

-	Scroll	
-	Save Tir	ne Interval
-	-	Off
-	-	0.5 min
-	-	1 min
-	-	2 min
-	-	2.5 min
-	-	5 min
-	-	10 min
-	-	15 min
-	-	20 min
-	-	30 min
-	-	60 min
-	-	120 min
	-	Return
-	Trend C	lear
-	-	Yes
-	-	No
-	Return	

GRAPH		REND MENU
-	Scroll	
-	HR/PR	
-	-	On
-	-	Off
-	NIBP	
-	-	On
-	-	Off
-	(P1 Lab	el)
-	-	On
-	-	Off
-	(P2 Lab	el)
-	-	On
-	-	Off
-	SPO ₂	
-	-	On
-	-	Off
-	RESP	
-	-	On
-	-	Off
-	EtCO ₂	
-	-	On
-	-	Off
-	T1	
-	-	On
-	-	Off
-	T2	
-	-	On
-	-	Off
-	Trend C	lear
-	-	Yes
-	-	No
-	Return	

DATE/TIME MENU

-	Date		
-	-	YYY	Y
-	-	MM	
-	-	DD	
-	Time		
-	-	HH	"24 hours only"
-	-	MM	
-	Return		

SETUP	MENU	
-	Patient	Mode
-	-	Adult
-	-	Neonatal
-	-	Return
-	Record	Speed
-	-	25 mm/s
_	_	50 mm/s
_	_	Doturn
-	- Waya Dr	
-		
-	-	20 Sec
-	-	Continuous
-	-	
-	wave Re	ecord Select
-	-	ECG 1 + ECG 2
-	-	ECG 1 + PLETH
-	-	ECG 1 + RESP
-	-	ECG 1 + (P1 Label)
-	-	ECG 1 + (P2 Label)
-	-	ECG 1+ CAPNO
-	-	Return
-	Record	On Alarm
-	-	On
-	-	Off
-	Auto Lis	st Record
-	-	On
-	-	Off
-	Alarm V	olume
		=
-	-	
-	HR/PR 1	one Volume
		F
-	-	=
_	_	Off
_	- Kov Boo	on Volume
-	Ney Dee	
-	-	=
		Off
-	- Sloon M	odo
-	Sleep IM	Oue
-	-	
-	-	
-	-	20 min
-	-	30 min
-	-	Return
-	Main Sc	reen
-	-	4-ch Wave
-	-	6-ch Wave
-	-	Big Number
-	-	Return
-	Service	Menu
-	-	Pass Code
-	-	Return

ALARM	I LIMITS	MEN	U							
-	Record	On Ala	arm							
-	-	On								
-	-	Off	0.1			" — 1				
-	Audible	Alarm	1 Siler	ice Pe	riod	"The (curren	t setting	g is sho	wn only.
-	-	30 Se	C		i nis p	erioa m	ay be	set via	Servic	e menu"
-	-	60 Se	C							
-	-	30 Se 120 c								
-	- Audible	Δlarm	ec 1 Susr	end F	Period	"The	curren	t settin	a is shr	wn only
-	-	Off	000		This n	eriod m	av be	set via	Servic	e menu"
-	-	10 mi	n		rino p	onoum	ay 80	oot na		•
-	-	20 mi	n							
-	-	30 mi	n							
-	-	60 mi	n							
-	-	Indef	inite							
-	Alarm L	imits	"A	larm lir	nits adj	ustment	/Alarn	n suspe	end sett	ing"
		HR/PF	R SF	O 2	RESP	InCO;	2 Et(CO2	T1	T2
		180	10	0	30	20	8	0	38.0	38.0
		40	9	0	0	0		0	14.5	14.5
-	-		÷							≜
		On	0	n	On	On		On	On	On
-	-	×	×		×	×		×	×	×
		Off	0	ff	Off	Off	(Off	Off	Off
					/1	71 Lohal	、	,		.n
		eve			1) 9V9) ^	(eve		
		200	180	160	200		160	200		160
		70	40	30	70	40	30	70	40	30
		10	40	00	10	- I U	00	10		
-	-		-			-			-	
			On			On			On	
-	-		×			×			×	
			Off			Off			Off	
-	-	Retur	'n							
-	Alarm L	imits I	Displa	y						
-	-	On								
-	-	Off								
-	Auto Ala	arm								
-	-	On								
-	- Auto Ali		ottina		"0/ oott	ing for c	ooh n	oromot	~~"	
-	Auto Ala	HR/PF	etting Sing	0.	% Sell	InCO	acrip Ft	aramet 20.	e/ T1	Т2
		400/		02 00/	400/	400/		002 007	400/	12
		40%	5 40	J%	40%	40%	40	0%	40%	40%
		-20%	-2	0%	-20%	-20%	-2	0%	-20%	-20%
			NIBP		(F	P1 Label)	(P2 Labe	el)
		SYS	MAP	DIA	SYS	MEAN	DIA	SYS	MEAN	DIA
		40%	40%	40%	40%	40%	40%	40%	40%	40%
		-20%	-20%	-20%	-20%	-20%	-20%	-20%	-20%	-20%
-	-	Retur	'n							
-	Return									

Note: Alarm limits shown above are for Adult patient mode. In order to set alarm limits to Neonatal patient mode, change Patient mode via *Setup menu*.

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PRINTING

General

The monitor can print real-time measurement and trend data as follows.

- 1. Set Record Speed, Wave Record Time, Wave Record Select, Record on Alarm, Auto List Record via Setup menu (refer Table 11).
- 2. To start printing, press Record button.
- 3. To stop printing during print out, press *Record button* again.

Record Speed

The record speed for is user-selectable: either 25.0 or 50.0 mm/s. When 50.0 mm/s is selected, the wave record time is fixed to 20 sec.

Wave Record Time

20 sec

A 20-second print, recording real-time graphical and numeric information beginning 10 seconds before the print initiation and ending 10 seconds after that event.

Continuous

A print of real-time graphical and numeric information, beginning 10 seconds before initiating the action and continuing until stopped.

Wave Record Select

The monitor prints out the real-time waveforms selected by the user.

- ECG 1+ ECG 2
- ECG 1+ PLETH
- ECG 1+ RESP
- ECG 1+ IBP 1
- ECG 1+ IBP 2
- ECG 1 + CAPNO

Record On Alarm

If **Record on alarm** is set to **On** in **Setup menu**, the monitor will automatically print out whenever a physiological alarm condition occurs.

Auto list record

If the Auto List Record is **On**, 8 data recorded in trend memory will be automatically printed out.

Print Out Configuration

20 Sec Print Out

If *Wave record time* is set to *20 sec*, the monitor will print out numeric data and waveforms by pressing *Record button* as shown Figure 53



Figure 53. 20 Sec Printing

Continuous Print Out

If *Wave record time* is set to *Continuous*, the monitor will print out numeric data every minute and waveforms continuously by pressing the *Record button* as shown in Figure 54.



Figure 54. Continuous Printing

Tabular Trend Data Print Out

When tabular trend data is displayed on the screen, the monitor will print out the displayed data by pressing the *Record button* as shown in Figure 55.

{т	'IME	HR/PR	NIBP	ABP	CVP	SpO ₂	RESP	EtCO ₂	T1	T2
т	MM/DD	[BPM]	[mmHg]	[mmHg]	[mmHg]	[%SpO ₂]	[/min]	[mmHg]	[°C]	[°C]
05/ 8/ 05/ 8/ 05/ 8/ 05/ 8/	6 18:00 6 18:02 6 18:02 6 18:02 6 18:04	80 88 74 74		118/84(93) 118/72(96) 119/79(98) 124/72(89)	120/80(93) 128/82(96) 109/89(95) 114/72(90)	94 95 95 92	25 18 18 17	35 32 <mark>80↑</mark> 31	38.5 38 37.7 38.2	37.5 38.2 38.4 38.1
05/ 8/	6 18:04	80	/()	104/71(82)	114/51(80)	80]	19	20	38.1	38.6
05/ 8/	6 18:06	75	/()	111/74(81)	112/76(82)	91	17	25	37.8	37.8
05/ 8/	6 18:06	74	102/68(83)	108/78(85)	118/68(85)	91	16	30	38.0	38.4
05/ 8/	6 18:08	76	/()	111/62(84)	118/63(85)	95	20	31	37.9	37.6

Figure 55. Tabular Trend Printing

Graphical Trend Data Print Out

When graphical trend data is displayed on the screen, the monitor will print out the displayed data by pressing the *Record button* as shown in Figure 56.



Figure 56. Graphical Trend Printing

Setting Information Print Out

The monitor can print out all internal settings by selecting *Print value of configuration* in *Service menu* as shown in Figure 57. Refer to the service manual for the detailed instructions.

ξ	Y/M/D 2005/ 8/ 6	TIME 18:00	P1 Scale	AUTO	BO On Alarm	Off	-CVP MENU		Alarm (InCO ₂)	20 / 0
ξ	-ECG WAVE MENU-		-CVP WAVE MENU-		Smart Clock	On	Alarm (SYS)	200 / 70	Alarm (EtCO ₂)	80 / 0
ş	ECG Cable Select	AUTO	Sweep Speed	25.0mm/s	Completion Sound	On	Alarm (MEAN)	180 / 40	-TEMPERATURE M	ENU
Ş	Sweep Speed	25.0mm/s	P1 Label	CVP	Smart Inflation	On	Alarm (DIA)	160 / 30	Alarm (T1)	38.0 / 14.5
3	Pacer Detect	Off	P2 Scale	AUTO	Alarm (SYS)	200 / 70	-SpO2 MENU		Alarm (T2)	38.0 / 14.5
ζ	Filter Mode	Monitor	-CAPNO WAVE MEN	U	Alarm (MEAN)	180 / 40	C-Lock	Off	-ALARM LIMITS ME	NU (
ξ	-PLETH WAVE MEN	ل	Sweep Speed	12.5mm/s	Alarm (DIA)	160 / 30	Alarm	100 / 90	Record On Alarm	Off 2
ξ	Sweep Speed	25.0mm/s	Scale	AUTO	-NIBP INTERVAL SE	TTING MENU	-RESPIRATION MEN	U	Audible Alarm Silen	ce Period
ş	-REPIRATION WAVE	MENU	-HR/PR MENU		Interval	Off	Respiration/Apnea	AUTO		120 sec
Ş	Sweep Speed	12.5mm/s	HR/PR Source	AUTO	-ABP MENU		Apnea Time Setting	40 sec	Audible Alarm Susp	end Period
3	-ABP WAVE MENU-		Alarm	180 / 40	Alarm (SYS)	200 / 70	Alarm	30/0		Indefinite 3
ζ	Sweep Speed	25.0mm/s	-NIBP WAVE MENU-		Alarm (MEAN)	180 / 40	-CO2 MENU		Alarm Limits Display	y S
ş	P1 Label	ABP	Initial Inflate	180	Alarm (DIA)	160 / 30	Capno Measurement	On		On

Auto Alarm	Off	ABP (MEAN)	40% / -20%	RESP	On	Key Beep Volume	4	Unit of Temp	°C
Auto Alarm Setting(%	6)	ABP (DIA)	40% / -20%	EtCO ₂	On	Sleep Mode	Off	Language Setting	English
{ HR/PR	40% / -20%	CVP (SYS)	40% / -20%	T1	On	-SERVICE MENU		Date Format	Year/Month/Day
{ SpO₂	40% / -20%	CVP (MEAN)	40% / -20%	T2	On	Save Settings On Por	wer Off	Demo Mode	Off
S RESP	40% / -20%	CVP (DIA)	40% / -20%	-SETUP MENU		-	Back Up		
\$ InCO ₂	40% / -20%	-TABULAR TRENDS	MENU	Patient Mode	Adult	Audible Alarm Silence	e Period		
	40% / -20%	Save Time Interval 0	.0 min	Record Speed	25 mm/s		120 sec		
λ T1	40% / -20%	- GRAPHICAL TREN	DS MENU	Wave Record Time	Continuous	Audible Alarm Suspe	nd Period		
{ т2	40% / -20%	HR/PR	On	Wave Record Time	ECG1 + PLETH		Indefinite		
SVIBP (SYS)	40% / -20%	NIBP	On	Record On Alarm	Off	AC Line Frequency	60Hz		
S NIBP (MAP)	40% / -20%	ABP	On	Auto List Record	Off	Unit of NIBP	mmHg		
S NIBP (DIA)	40% / -20%	CVP	On	Alarm Volume	5	Unit of IBP	mmHg		
ABP (SYS)	40% / -20%	SpO ₂	On	HR/PR Tone Volume	4	Unit of CO ₂	mmHg		:

Figure 57. Setting Information Printing

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EXTERNAL INTERFACE

General

The monitor provides external connectors on the right panel to support communication with external equipment and functions such as a nurse call, software upgrades or PC connection.

WARNING: Any connections between this monitor and other devices must comply with applicable medical systems safety standards such as IEC 60601-1. Failure to do so could result in unsafe leakage current and grounding conditions.

Note: This equipment is to be used on a network and the communication wirings (RJ45 LAN and RS232) are limited to inside of the building.

Cable Connections

RS-232

The pin layouts of 15-pin RS-232 interface are illustrated below.

Figure 58. Data Port Pin Layout

Table 40. RS-232 Serial Inter	face Connections
-------------------------------	------------------

Pin #	Signal
1	Not used (open)
2	RS232 RX
3	RS232 TX
4	not used (open)
5	RS232 ground
6	not used (open)
7	Nurse call normally open
8	Nurse call normally closed
9	not used (open)
10	RS232 ground
11	not used (open)
12	not used (open)
13	not used (open)
14	not used (open)
15	Nurse call common

Nurse Call Interface

WARNING: The nurse call feature should not be used as the primary source of alarm notification. The audible alarms of the monitor, used in conjunction with clinical signs and symptoms, are the primary sources for notifying medical personnel that an alarm condition exists.



CAUTION: The nurse call feature is not functional whenever the monitor alarms are silenced.



CAUTION: The nurse call function needs to be tested after it has been set up in your facility. The nurse call feature should be tested whenever setting up the BP-S510 in a location that uses nurse call. One way to test the nurse call function is to create an alarm condition (for example, sensor disconnect) and verify that your facility's nurse call system is activated.

The nurse call feature of the monitor is operational when the monitor is powered by AC power or battery power. The nurse call feature of the monitor works in conjunction with the nurse call system of your institution when the monitor sounds an audible alarm.

The monitor provides the nurse call interface of relay closure type. The interface functions when the monitor is operating either on AC power or battery power.

The remote location is signaled anytime there is an audible alarm. If the audible alarm has been turned off or silenced, the nurse call function is also turned off.

Nurse Call Relays Normally Open/Closed

Pins 7 and 15 provide a relay that closes when an alarm is sounding on the monitor. Pins 8 and 15 provide a relay that opens when an alarm is sounding. Pin 15 is a common lead for both relays.

MAINTENANCE

WARNING: The cover should be removed only by qualified service personnel. There are no internal user-serviceable parts except for the battery.

WARNING: Do not spray, pour, or spill any liquid on the monitor, its accessories, connectors, switches or openings in the chassis.

WARNING: Unplug the power cord from the monitor before cleaning the monitor.

Recycling and Disposal

When the monitor, battery, or accessories reach the end of useful life, recycle or dispose of the equipment according to appropriate local and regional regulations.

- Note: The monitor should be disposed of separately from the municipal waste stream via designated collection facilities appointed by the government or the local authorities.
- Note: The correct disposal of your old appliance will help prevent potential negative consequences for the environment and human health.
- Note: 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 monitor.

Returning the Monitor and System Components

Pack the monitor with sensors, cable or other accessory items in its original shipping carton. If the original carton is not available, use a suitable carton with appropriate packing material to protect the monitor during shipping.

Service

The monitor requires no routine service other than cleaning, battery maintenance, and service activity which is mandated by the user's institution. For more information, refer to the monitor service manual. Qualified service personnel in the user's institution should perform periodic inspections of the monitor. If service is necessary, contact qualified service personnel or your local supplier.

Periodic Safety Checks

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

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

Cleaning

The monitor may be surface-cleaned by using a soft cloth dampened with either a commercial, nonabrasive cleaner or one of the solution listed in the below. Lightly wipe the top, bottom and front surfaces of the monitor lightly.

- 70% Isopropyl alcohol
- 10% Chlorine bleach solution

For cables, sensors, cuffs, and probes, follow the cleaning instructions in the directions for use shipped with those components.

Avoid spilling liquid on the monitor, especially in connector areas. If liquid is accidentally spilled on the monitor, clean and dry thoroughly before reuse. If in doubt about monitor safety, refer the unit to qualified service personnel for checking.

Battery Maintenance



CAUTION: Recharging the battery is strongly recommended when the battery has not been recharged for 2 or more months.



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

If the monitor has not been used 2 months, the Ni-MH battery will need charging. To charge the battery, connect the monitor to an AC power source as described in the **Battery Operation** section.

- Note: Storing the monitor for a long period without charging the battery may degrade the battery capacity. A full charge of a depleted battery takes over 12 hours
- Note: The service menu displays the number of deep discharge cycles seen by the battery. The monitor records a deep discharge cycle when the battery reaches the voltage at which a "Critically low battery" alarm is issued. Refer to the service manual for the details.
- Note: The battery should be removed from the monitor if placed in storage or will not be used for a long period.

Colin Medical Technology recommends that the monitor's Ni-MH battery be replaced if it has been stored for 2 years or more. Refer to the monitor service manual for battery replacement and general service instructions.

Loading Recorder Paper



CAUTION: Use only recorder paper specified by Colin Medical Technology.

Note: The paper roll is easier to load if it is held horizontally with your thumb on top and your forefinger and/or index finger underneath it.

Load recorder paper as follows:

- 1. Open the recorder door by pulling the latch on the recorder slightly and carefully. The door should tilt open. Gently pull the door open if necessary.
- 2. Reach in and remove the empty paper core by pulling it over gently with your thumb and index finger.
- 3. Insert a new paper roll oriented properly.
- 4. Pull the paper out towards you until approximately 2 inches (5 cm) of paper have been unrolled.
- 5. Align the paper with the pinch roller attached to the recorder door.
- 6. Close the recorder door.
- Note: To make sure that the paper is aligned in the slot and has not been pinched in the door, pull the loose edge until a few inches of paper is showing. If the paper will not move, open the door and return to step 4.



Figure 59. Recorder Paper Replacement

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TROUBLESHOOTING

WARNING: If you are uncertain about the accuracy of any measurement, check the patient's vital signs by alternate means; then make sure the monitor is functioning correctly.



WARNING: The cover should be removed only by qualified service personnel. There are no user-serviceable parts inside except for the battery.

General

If the monitor detects an error, it can display an error code. The error codes are listed in the monitor service manual. If an error code is displayed, write down the code and contact your service department. Before calling your local supplier, make sure that the battery is charged and that all power connections are in place.

Corrective Action

If you experience a problem while using the monitor and are unable to correct it, contact qualified service personnel or your local supplier. The service manual provides additional troubleshooting information for qualified personnel.

Following is a list of possible errors and suggestions for corrective action.

1. There is no response to the Power on/off button.

- A fuse may be blown. Notify service personnel to check and replace the fuse.
- If operating on battery power, the battery may be missing or discharged. If the battery is discharged, charge the battery (see **Battery Operation** section).
- 2. The monitor screen does not function properly and the power-on beep tones do not sound during the power-on self test.
 - Do not use the monitor; contact qualified service personnel or your local supplier.

3. The monitor is operating on battery power, even though it is connected to AC.

- Make sure that the power cord is properly connected to the monitor.
- Check to see if power is available to other equipment on the same AC circuit.
- The monitor operates from its internal battery if there is no AC power source.

4. When the alarm condition occurs, check the following items.

- Check the alarm message in the alarm message area or informative message area.
- Follow the check items in the below table to remove the alarm condition

Alarm Messages	Check Items	
NIBP: Check cuff (C11)	Cuff pressure did not increase enough even after activating the	
	pump for more than 30 seconds (adult). There is a possibility that a	
	cuff hose is disconnected, or a cuff may not be wrapped around an	
	arm. Check cuff and cuff hose.	
	This error possibly occurs in case of large cuffs that are wrapped	
	around loosely.	
	When the error still occurs even after checking above, there is a	
	possible air leakage from a ruptured cuff.	
	Replace it with a new one.	
NIBP: Check cuff / Patient	Blood pressure could not be measured even after cuff pressure	
(C12)	decreased. It is possibly because pulse was not strong enough for	
	measurement, or because change of pulse amplitude could not be	

Alarm Messages	Check Items		
	obtained. Check whether cuffs are not wrapped around thick clothing. After wrapping cuffs around property, measure again. When the error occurs in the initial measurement in continuous mode, the second measurement will start unless Stop button is pressed.		
NIBP: Cuff excessive	Measurement failed because of patient movement during		
artifact (C13)	measurement. Tell the patient to stay still, then, measure again. When it occurs in the initial measurement in continuous mode, the second measurement will start unless Stop button is pressed.		
NIBP: Cuff insufficient pressure (C14)	Measurement failed because of insufficient pressurizing. There is a possibility that standard cuff pressure might be detected wrongly due to noises, motion artifact or external vibration. Check whether cuffs are not wrapped around thick clothing, whether the patient is moving and whether cuffs are free from outside vibrations, then, measure again. When it occurs in the initial measurement in continuous mode, the		
	second measurement will start unless Stop button is pressed.		
NIBP: Cuff irregular pulses (C15)	Blood pressure could not be measured because oscillation graph was not normal. There is a possibility that motion artifact or vibration from outside might interrupt the measurement. Check whether the patient stays still and cuffs are free from external vibration, then, measure again. When it occurs in the initial measurement in continuous mode, the		
NIDD: Cuff motion ortifact	second measurement is continued unless Stop button is pressed.		
(C16)	pulse waveform signal. Check for motion artifacts, or external vibration and then, measure again. When it occurs in the initial measurement in continuous mode, the second measurement will start unless Stop button is pressed.		
NIBP: Cuff time-out (C17)	Measurement was preventively stopped because measurement time exceeded 160 seconds (adult), There is a possibility that blood pressure might be repeatedly measured due to insufficient pressurizing caused by calcified pseudohypertension.		
NIBP: Cuff time-out, over 160 pulses (C18)	Pulse waveform signal more than 160 beats are detected during measurement. There is possibility that noises might interrupt signal. Motion artifact or external vibrations possibly affected cuffs. Check for patient movement and if the cuff is free from stays still and cuffs are free from outside vibration, then, measure again.		
NIBP: Cuff pressure failure (C19)	Cuff pressure exceeded more than 300 mmHg (adult) during measurement. There is a possibility that the patient moved during measurement or strong pressure from outside might be added to cuffs. Considering above, measure again.		
NIBP: Cuff weak pulse (C20)	Amplitude of pulse obtained from cuffs are too weak. This error possibly occurs when cuffs are wrapped around loosely in ASO patients or when cuffs are wrapped around thick clothing. Wrap cuffs around properly, then, measure again.		
NIBP: Check cuff, hose and mode (C21)	Patient to be measured, and cuff size used, do not match. This error may occur if the blood pressure measurement mode setting is incorrect, if the cuff has been wrapped tightly in the adult mode, loosely in the neonatal mode or if the arm has been bent during measurement. Check the measurement mode setting and application of the cuff, and measure again.		

Alarm Messages	Check Items	
NIBP: Internal error (E03)	NIBP module error	
- \'/	BPM pressure sensor fault.	
	Pump operated for ten seconds, however pressure does not	
	change. Check the connection of the cuff hose.	
NIBP: Internal error (E07)	Reboot the monitor. If the problem persists, cease use immediately	
	and contact qualified service personnel or your local supplier.	
NIBP: Internal error (E08)	Reboot the monitor. If the problem persists, cease use immediately	
	and contact qualified service personnel or your local supplier.	
NIBP: Internal error (E09)	NIBP module error	
	Fault detected in accordance with safety monitoring to BPM IEC	
	standards.	
	The pressure inside the cuff reaches the standard pressure.	
	Standard pressure Adult: 320mmHg	
	Neonatal: 157mmHg	
NIBP: Internal error (ROM)	Reboot the monitor. If the problem persists, cease use immediately	
	and contact qualified service personnel or your local suppler.	
NIBP: Internal error (RAM)	Reboot the monitor. If the problem persists, cease use immediately	
	and contact qualified service personnel or your local suppler.	
NIBP: Internal error (COM)	Reboot the monitor. If the problem persists, cease use immediately	
	and contact qualified service personnel or your local suppler.	
ECG: Internal error.	Reboot the monitor. If the problem persists, cease use immediately	
	and contact qualified service personnel or your local suppler.	
RESP: Internal error.	Reboot the monitor. If the problem persists, cease use immediately	
	and contact qualified service personnel or your local suppler.	
<i>{label}</i> : Loss of pulse.	Check the condition of the patient and fitting of the transducer and	
	cable, and measure again.	
IBP: Internal error.	Verify that the IBP module is correctly installed after the BP-S510	
	is powered off. If the problem persists, reboot the monitor.	
SpO ₂ : Loss of pulse.	Signal obtained from sensor is weak. SpO ₂ could not be measured.	
	I nere may be a problem with fitting of the SpO_2 sensor, or blood	
	flow at the sensor site may be unsatisfactory. Check the condition	
SpOo: Internal error	A problem with the SpOr measurement has been detected. The	
SpO2. Internal error.	A problem with the $3pO_2$ measurement has been detected. The $3pO_2$ measurement function does not operate. If switching power	
	OFE/ON has no effect it is possible that a fault has occurred. Stop	
	using the monitor immediately and contact qualified service	
	personnel or your local suplier.	
TEMP{n}: Internal error.	An internal circuit fault has been detected. If switching power	
	OFF/ON has no effect it is possible that a fault has occurred in the	
	monitor. Cease use immediately.	
CAPNO: Internal error.	A problem with the capnography measurement function has been	
	detected. The capnography measurement function does not	
	operate. If switching power OFF/ON has no effect it is possible that	
	a fault has occurred. Cease use immediately.	
CAPNO: Sensor error.	The connector may be damaged or a fault may have occurred	
	within the gas unit or main unit. Cease use immediately.	
SYSTEM: Critically low-	Connect the AC power cord of the monitor to the AC main to	
battery condition.	recharge the battery.	
SYSTEM: Real time clock	Reboot the monitor. If the problem persists, cease use immediately	
error.	and contact qualified service personnel or your local supplier.	
SYSTEM: WDT error.	Reboot the monitor. If the problem persists, cease use immediately	
	and contact qualified service personnel or your local supplier.	
SYSTEM: RAM error.	Reboot the monitor. If the problem persists, cease use immediately	
	and contact qualified service personnel or your local supplier.	

Alarm Messages	Check Items		
SYSTEM: Failure.	Reboot the monitor. If the problem persists, cease use immediately		
	and contact qualified service personnel or your local supplier.		
ECG : Check ECG leads &	ECG error is detected, Electrodes or lead wires may not be		
electrodes.	correctly attached or a circuit is possibly saturated due to offset		
	voltage. Check whether electrodes are correctly attached and		
	electrodes are new and wet. Confirm the patient's skin is clean.		
{label}: Cable/Sensor	Cable not connected to the main unit. If connected, the cable may		
disconnected.	be damaged. Replace with a new cable. If replacing the cable has		
	no effect the problem may be within the device. In this case, cease		
	use immediately.		
SpO ₂ : Check probe.	Sensor is not in contact with patient. SpO_2 could not be measured. Fit the sensor correctly to the patient, and measure again		
TEMP{n}: Temperature	Sensor not connected to the main unit. If connected, the cable may		
probe disconnected.	be damaged. Replace with a new cable. If replacing the cable has		
p	no effect the problem may be within the device. In this case, cease		
	use immediately.		
CAPNO: Occlusion.	Check the patient condition.		
CAPNO: Water trap full.	Disconnect the sampling tubing from the water trap and replace the		
	water trap with a new one.		
ECG: Signal saturation.	Decrease the ECG size via the setup menu.		
RESP: Check Resp leads	Electrodes or lead wires may not be correctly attached or a circuit		
& electrodes.	is possibly saturated due to offset voltage.		
	Check whether electrodes are correctly attached and electrodes		
	are new and wet. Confirm the patient's skin is clean.		
{label}: Unable to zero	Could not zero calibrate pressure. Check that the transducer is		
calibration.	open to the atmosphere, and check the three-way tap. As it is also		
	possible that the measured pressure incorporates noise, check the		
<u>"</u>	measurement circuit.		
{ <i>label</i> }: Out of range.	A value outside the measurement range was obtained. As the		
	measurement circuit		
SpO2 [.] Check sensor	Sensor not connected. If connected, the cable or connector may be		
	damaged. Replace with a new cable. If replacing the cable has no		
	effect the problem may be within the device. In this case, cease		
	use immediately.		
SpO2: Sensor failure.	A problem with the SpO ₂ sensor has been detected. The SpO ₂		
	measurement function does not operate. The possible cause is a		
	connection failure of the SpO ₂ sensor and the extension cable, or a		
	failure of the sensor or cable. Reconnect the sensor and extension		
	cable or replace them with new ones. If the problem doesn't clear		
	up after carrying out the remedies above or switching power		
	OFF/ON, a grave fault can develop, Cease the use of the sensor		
	immediately.		
SpO ₂ : Module reset.	A problem with the SpO ₂ measurement has been detected. The		
	SpO ₂ measurement function does not operate. If switching power		
	OFF/ON has no effect it is possible that a fault has occurred.		
	Cease use immediately.		
TEMP{n}: Out of range.	A measure reading outside the measurement range was obtained.		
	It is possible that the temperature in the vicinity of the sensor is		
	extremely low (less than 15.0°C) or extremely high (more than		
040N0 7	45.0°C). Adjust the ambient temperature and measure again.		
CAPNO: Zero calibration	i ry again the gas calibration.		
range error			

Alarm Messages	Check Items		
CAPNO: Zero calibration	Try again the gas calibration.		
signal unstable error			
CAPNO: High calibration	Try again the gas calibration.		
range error			
CAPNO: High calibration	I ry again the gas calibration.		
signal unstable error	Diverties AC nerven could to the AC main to non-bound the bottom.		
NIRD: Potry Chock	Plug the AC power cold to the AC main to recharge the battery.		
cuff/Patient (C12)	decreased. It is possibly because pulse was not strong enough for		
	measurement, or because change of pulse amplitude could not be		
	obtained. Check whether cuffs are not wrapped around thick		
	clothing. After wrapping cuffs around properly, measure again.		
	When the error occurs in the initial measurement in continuous		
	mode, the second measurement will start unless Stop button is		
	pressed.		
NIBP: Retry, Cuff	Measurement failed because of patient movement during		
excessive artifact (C13)	Measurement. Tell the patient to stay still, then, measure again.		
	second measurement will start unless Ston button is pressed		
NIBP [.] Retry Cuff	Measurement failed because of insufficient pressurizing. There is a		
insufficient pressure (C14)	possibility that standard cuff pressure might be detected wrongly		
	due to noise, patient movement or external vibrations. Check		
	whether cuffs are not wrapped around thick clothing, whether the		
	patient is moving and whether there is external vibration. When it		
	occurs in the initial measurement in continuous mode, the second		
	measurement will start unless Stop button is pressed.		
NIBP: Retry, Cuff irregular	Blood pressure could not be measured because oscillation graph		
puises (CTS)	was not normal. There is a possibility that patient movement of		
	patient stays still and cuffs are free from outside vibrations, then		
	measure again. When it occurs in the initial measurement in		
	continuous mode, the second measurement will start unless Stop		
	button is pressed.		
NIBP: Retry, Cuff motion	Blood pressure could not be measured because noises interrupted		
artifact (C16)	pulse waveform signal. There is a possibility that patient movement		
	or external vibration interrupted the measurement. Confirm the		
	patient is not moving and the cuff is free of external vibration, then,		
	measure again. When it occurs in the initial measurement in continuous mode, the second measurement will start unless. Step		
	button is pressed		
NIBP: Retry, Cuff time-out,	Pulse waveform signal more than 160 beats are detected during		
over 160 pulses (C18)	measurement. There is a possibility that noise, patient movement		
	or external vibrations. Confirm the patient is not moving and the		
	cuff is free of external vibration, then, measure again.		
NIBP: Retry, Cuff pressure	Cuff pressure exceeded more than 300mmHg during		
failure (C19)	measurement. There is a possibility that the patient moved during		
	measurement or strong pressure from outside might be added to		
NIPD: Dotry Chook ouff	Curis, Considering above, measure again.		
hose and mode (C21)	Falleni to be measured, and cun size used, do not match. This error may occur if the blood pressure measurement mode setting is		
	incorrect, if the cuff has been applied tightly in the adult mode		
	loosely in the neonatal mode or if the arm has been bent during		
	measurement. Check the measurement mode setting and		
	application of the cuff, and measure again.		
{label}: No zero reading.	Perform the pressure zero setting.		

Alarm Messages	Check Items
SpO2: Motion artifact.	SpO2 could not be measured due to signal noise thought to be due
	to body movement. Ensure that the patient remains at rest, and
	measure again.
SYSTEM: No recorder	In case the recorder door is open, close the door.
paper.	In case the recorder paper is empty, insert new paper and close
	the door.
SYSTEM: Abnormally shut	The monitor has been abnormally shut down last time. Contact
down last time.	qualified personnel in your facility or your local supplier.
SYSTEM: No recorder	The recorder is not installed in your monitor. If required, contact
installed.	your local supplier.

EMI (Electromagnetic Interference)

WARNING: Keep patients under close surveillance when monitoring. It is possible, although unlikely, that radiated electromagnetic signals from sources external to the patient and monitor can cause inaccurate measurement readings. Do not rely entirely on the monitor readings for patient assessment.
WARNING: It is possible that any radio frequency transmitting equipment and other nearby sources of electrical noise may result in disruption in the monitor operation.
WARNING: It is possible, although unlikely, that large equipment using a switching relay for its power on/off may affect monitor operation. Do not operate the monitor in such environments.
This device has been tested and found to comply with the limits for medical devices to the IEC60601-1-2, and the Medical Device Directive 93/42/EEC. These limits are designed to provide reasonable protection against harmful interference in a typical medical installation.
However, because of the proliferation of radio-frequency transmitting equipment and other sources of electrical noise in health care environments (such as electrosurgical equipment, defibrillator, cellular phones, mobile two-way radios, electrical appliances, and high-definition television), it is possible that high levels of such interference due to close proximity or strength of a source may affect monitor operation.
WARNING: The monitor is designed for use in environments in which the signal can be obscured by electromagnetic interference. During such interference, measurements may seem inappropriate or the monitor may not seem to operate correctly.

Monitor disruption may be indicated by erratic readings, cessation of operation, or other incorrect functioning. If this occurs, survey the site to determine the source of this disruption. Try the following actions to see if they eliminate the disruption:

- Turn equipment in the vicinity off and on to isolate the offending equipment.
- Reorient or relocate the interfering equipment.
- Increase the separation between the interfering equipment and this equipment.

The monitor generates, uses, and can radiate radio frequency energy. If the monitor is not installed and used in accordance with these instructions, the monitor may cause harmful interference with other devices in the vicinity.

If assistance is required, contact your local supplier.

Obtaining Technical Assistance

For technical information and assistance, or to order a monitor service manual, call your local supplier. The service manual provides information required by qualified service personnel when servicing the monitor.

When calling your local supplier, you may be asked to provide the software version number of your monitor. The software version is displayed when monitor power is activated.

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General

The monitor is shipped with factory default settings. Authorized personnel can use the procedures described in the service manual to change default settings.

Parameter Ranges and Default Settings

Paramotor	Pangos/Soloctions	Factory Defaults	
	Kanges/Selections	Adult	Neonatal
ECG			
ECG Cable Select	3 Leads, 5 Leads, AUTO	AUTO	AUTO
ECG Lead Select	I, II, III, aVR, aVL, aVF, V(Chest Lead)	-	-
ECG Size (mm/mV)	×1/4, ×1/2, ×1, ×1.5, ×2	×1	×1
ECG Filter Mode	Monitor, Low Extend, Filter,	Monitor	Monitor
	Respiration Rejection	WORITO	wormor
ECG Pacer Detect	On, Off	Off	Off
ECG Sweep Speed	12.5, 25.0, 50.0 mm/s	25.0 mm/s	25.0 mm/s
HR/PR Source	AUTO, HR, PR	AUTO	AUTO
HR/PR Upper Alarm Limits	35 to 305 BPM (Adult/Neo) (5 BPM steps)	180 BPM	200 BPM
HR/PR Lower Alarm Limits	30 to 300 BPM (Adult/Neo) (5 BPM steps)	40 BPM	50 BPM
NIBP			
NIBP Initial Cuff Inflation	120, 140, 160, 180, 200, 220mmHg (Adult)		
	(16.0, 18.7, 21.3, 24.0, 26.7, 29.3, kPa)	180 mmHg	120 mmHg
	80, 100, 120, 140 mmHg (Neo)	24.0 kPa	16.0 kPa
	(9.3, 12.0, 14.7, 16.0, 18.7 kPa)		
BP On Alarm	On, Off	Off	Off
Smart Clock	On, Off	On	On
Smart Inflation	On, Off	On	On
Completion Sound	On, Off	On	On
NIBP Automatic Mode Interval	Off, Cont, 1, 2, 2.5, 3, 5, 10, 15, 20, 30, 45, 60,	o."	
	90,120 ,180 minutes	Off	Off
NIBP SYS Upper Alarm Limits	60 to 260 mmHg (Adult), 40 to 130 mmHg (Neo)		400 11
	8.0 to 34.7 kPa (Adult), 5.3 to 17.3 kPa (Neo)	200 mmHg	130 mmHg
	(10 mmHg / 1.3 kPa steps)	26.7 KPa	17.3 кРа
NIBP SYS Lower Alarm Limits	50 to 250 mmHg (Adult), 30 to 120 mmHg (Neo)	70	50 mml la
	6.7 to 33.3 kPa (Adult), 4.0to 16.0 kPa (Neo)		50 mmHg
	(10 mmHg / 1.3 kPa steps)	9.3 KPa	0.7 KPa
NIBP DIA Upper Alarm Limits	40 to 210 mmHg (Adult), 20 to 100 mmHg (Neo)	100 mml la	100 mml/m
	5.3 to 28.0 kPa (Adult), 2.7 to 13.3 kPa (Adult)		100 mmHg
	(10 mmHg / 1.3 kPa steps)	21.3 KPa	13.3 KPa
NIBP DIA Lower Alarm Limits	30 to 200 mmHg (Adult), 10 to 90 mmHg (Neo)	20 mmHa	10 mmHa
	4.0 to 26.7 kPa (Adult), 1.3 to 12.0 kPa (Neo)		
	(10 mmHg / 1.3 kPa steps)	4.0 KF a	1.3 KF a
NIBP MAP Upper Alarm Limits	50 to 240 mmHg (Adult), 30 to 110 mmHg (Neo)	180 mmHa	110 mmHa
	6.7 to 32.0 kPa (Adult), 4.0 to 14.7 kPa (Neo)	24.0 kPa	14.6 kPa
	(10 mmHg / 1.3 kPa steps)	27.0 N a	17.0 KI a
NIBP MAP Lower Alarm Limits	40 to 230 mmHg (Adult), 20 to 100 mmHg (Neo)	40 mmHa	20 mmHa
	5.3 to 30.7 kPa (Adult), 2.7 to 13.3 kPa (Neo)	53kPa	20 mm ly 27 kPa
	(10 mmHg / 1.3 kPa steps)	5.5 KI a	2.1 NI a

Table 41. Parameter Ranges and Factory Defaults

Parameter Ranges/Selections Aduit Neonatal IBP Aduit Neonatal IBP 25.0 mm/s 25.0 mm/s 25.0 mm/s 25.0 mm/s 25.0 mm/s Pressure Zero Setting Yes, No No No No P1 Label 0~50, 0~100, 0~200, 0~300, AUTO AUTO AUTO P1 Scale 0~50, 0~100, 0~200, 0~300, AUTO AUTO AUTO P1 Stole -50 to 280 mmHg (Aduit/Neo) (13 kPa steps) 26.7 KPa 17.3 kPa P1 SVS Lower Alarm Limits -60 to 250 mmHg (Aduit/Neo) (13 kPa steps) 21.3 kPa 10 mmHg P1 DIA Upper Alarm Limits -60 to 220 mmHg (Aduit/Neo) (10 mmHg steps) 100 mmHg 100 mmHg P1 MEAN Upper Alarm Limits -60 to 220 mmHg (Aduit/Neo) (10 mmHg steps) 100 mmHg 24.0 kPa 17.3 kPa P1 MEAN Upper Alarm Limits -60 to 220 mmHg (Aduit/Neo) (10 mmHg steps) 100 mmHg 20 mmHg 100 mmHg P2 Label P2, CVP, PAP, LAP CVP CVP 27.rkPa 27.rkPa P2 Sate 0-20, 0~50, 0-100, 0~200, 0~300, AUTO AUTO AUTO 210 mm			Factory Defaults	
IBP IBP Sweep Speed 12.5, 25.0, 50.0 mm/s 25.0 mm/s 25.0 mm/s IBP Sweep Speed 12.5, 25.0, 50.0 mm/s No No No Pressure Zero Setting Yes, No ABP ABP ABP P1 Scale 0-50, 0-100, 0-200, 0-300, AUTO AUTO AUTO AUTO P1 SyS Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 267, KPa 17.3 KPa P1 DX Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 160 mmHg 100 mmHg -8.0 to 33.3 KPa (Adult/Neo) (10 mmHg steps) 160 mmHg 100 mmHg 13.3 KPa P1 DIA Lower Alarm Limits -50 to 260 mmHg (Adult/Neo) (13 KPa steps) 24.0 KPa 13.3 KPa P1 MEAN Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (13 KPa steps) 24.0 KPa 14.7 KPa P1 MEAN Lower Alarm Limits -50 to 250 mmHg (Adult/Neo) (10 mmHg steps) 130 mmHg 27.7 KPa P2 Label P2, CVP. PAP, LAP CVP CVP 27.6 KPa 27.7 KPa P2 SYS Lower Alarm Limits -50 to 230 mmHg (Adult/Neo) (10 mmHg steps) 200 mmHg 130 mmHg 130 mmHg	Parameter	Ranges/Selections	Adult	Neonatal
IBP Sweep Speed 12.5, 25.0, 50.0 mm/s 25.0 mm/s 25.0 mm/s Presure Zero Setting Yes, No No No P1 Label P1, AbP ABP ABP P1 SS Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 200 mHg 130 mmHg P1 SYS Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (11 skPa steps) 93 kPa 5.7 kPa 17.3 kPa P1 SYS Lower Alarm Limits -60 to 260 mmHg (Adult/Neo) (13 kPa steps) 160 mmHg 100 mmHg P1 DIA Lower Alarm Limits -50 to 260 mmHg (Adult/Neo) (13 kPa steps) 13.8 kPa 13.3 kPa P1 MEAN Upper Alarm Limits -50 to 230 mmHg (Adult/Neo) (10 mmHg steps) 30 mmHg 10 mmHg -8.0 to 33.8 KPa (Adult/Neo) (10 mHg steps) 30 mMHg 10 mmHg 1.3 kPa P1 MEAN Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (13 kPa steps) 20 mmHg 20 mmHg -8.0 to 33.8 KPa (Adult/Neo) (10 mmHg steps) 30 mmHg 2.7 kPa 2.7 kPa P2 Label P2. CVP, PAP, LAP CVP CVP 2.7 kPa P2 DIA Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (13 kPa steps) 70 mmHg	IBP			
Pressure Zero Setting Yes, No No No No No P1 Label P1, ABP ABP ABP ABP P1 Scale 0~50, 0~100, 0~200, 0~300, AUTO AUTO AUTO AUTO P1 SyS Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 200 mmHg 50 mmHg P1 SYS Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 9.3 kPa 6.7 kPa P1 DIA Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (13 kPa steps) 9.3 kPa 10 mmHg P1 DIA Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (13 kPa steps) 4.0 kPa 1.3 kPa P1 MEAN Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (11 mmHg steps) 40 mmHg 10 mmHg P1 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 4.0 kPa 1.3 kPa P1 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 4.0 mmHg 20 mmHg P2 Label P2, CVP, PAP, LAP CVP CVP CVP P2 P2 SVS Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 70 mmHg 50 mmHg P2 DIA Lower Alar	IBP Sweep Speed	12.5. 25.0. 50.0 mm/s	25.0 mm/s	25.0 mm/s
P1 Label P1, ABP ABP ABP P1 P1 Scale 0~50, 0~100, 0~200, 0~300, AUTO AUTO AUTO AUTO P1 SYS Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 26, 7 KPa 17.3 KPa P1 SYS Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 9.3 kPa 6, 7 KPa P1 DIA Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 100 mmHg 100 mmHg P1 DIA Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 0.0 kPa 1.3 kPa P1 DIA Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (13 mHg steps) 0.0 mmHg 10 mmHg P1 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 0.0 mmHg 10 mmHg P1 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 5.3 kPa 2.7 kPa P2 Label P2, CVP, PAP, LAP CVP CVP CVP P2 Scale 0~20, 0~50, 0~100, 0~200, 0~300, AUTO AUTO AUTO P2 DIA Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 20 mmHg 50 mmHg P2 DIA Lower Alarm Limits -60 to 2	Pressure Zero Setting	Yes No	No	No
P1 Scale D-100. P-100. AUTO AUTO AUTO AUTO P1 Scale -50.0 = 260 mmHg (Adult/Neo) (10 mmHg steps) 200 mmHg 130 mmHg P1 SYS Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 70 mmHg 50 mmHg P1 DIA Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 9.3 kPa 6.7 kPa P1 DIA Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 100 mmHg 130 mmHg P1 DIA Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 4.0 kPa 13.3 kPa P1 MEAN Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 4.0 kPa 14.7 kPa P1 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 4.0 kPa 14.7 kPa P1 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 4.0 kPa 12.7 kPa P2 Label P2. CVP, PAP, LAP CVP CVP CVP P2 SY Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 3.0 mmHg 130 mmHg P2 DIA Lower Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 100 mmHg 100 mmHg	P1 Label	P1 ABP	ARP	ARP
1 House 1 House 1 House 1 House 1 House P1 SYS Upper Alarm Limits 50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 26.7 KPa 26.7 KPa P1 SYS Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 9.3 kPa 6.7 kPa P1 DIA Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (13 kPa steps) 100 mmHg 100 mmHg P1 DIA Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 30 mmHg 100 mmHg P1 DIA Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 4.0 kPa 1.3 kPa P1 MEAN Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 4.0 kPa 1.3 kPa P1 MEAN Lower Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 4.0 kPa 1.3 kPa P2 Label P2, CVP, PAP, LAP CVP CVP 27 kPa P2 State -7 to 34.7 kPa (Adult/Neo) (10 mmHg steps) 50 mmHg 50 mmHg P2 Sty Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 200 mmHg 130 mmHg P2 DIA Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 200 mmHg 50 mmHg P2 DIA Lower	P1 Scale	0~50 0~100 0~200 0~300 AUTO		
11 D10 Opper Namir Limits -60 for 200 mml (pcdubtReb) (1.3 kPa steps) 200 mml (steps) 200 mml (steps) P1 SYS Lower Alarm Limits -60 for 250 mml (pcdubtReb) (1.3 kPa steps) 70 mml (steps) 20.7 kPa P1 DIA Upper Alarm Limits -50 to 260 mml (pcdubtReb) (1.3 kPa steps) 160 mml (steps) 21.3 kPa P1 DIA Lower Alarm Limits -50 to 260 mml (pcdubtReb) (1.3 kPa steps) 20.1 mml (steps) 160 mml (steps) P1 DIA Lower Alarm Limits -60 to 250 mml (pcdubtReb) (1.3 kPa steps) 4.0 kPa 11.3 kPa P1 MEAN Upper Alarm Limits -60 to 250 mml (pcdubtReb) (1.3 kPa steps) 4.0 kPa 1.4 r RPa P1 MEAN Lower Alarm Limits -60 to 250 mml (pcdubtReb) (1.3 kPa steps) 4.0 kPa 1.4 r RPa P1 MEAN Lower Alarm Limits -60 to 250 mml (pcdubtReb) (1.3 kPa steps) 5.3 kPa 20 mml (steps) P2 Label P2. CVP, PAP, LAP CVP CVP CVP P2 SYS Upper Alarm Limits -60 to 250 mml (pcdubtReb) (1.3 kPa steps) 70 mml (steps) 30 mml (steps) P2 DIA Upper Alarm Limits -60 to 250 mml (pcdubtReb) (1.3 kPa steps) 70 mml (steps) 70 mml (steps) 70 mml (steps) P2 DIA Upper Alarm Limits -60 to 250 mml (pcdubtReb) (1.3 kPa steps) 70 mml (steps) <td>P1 SVS Upper Alarm Limits</td> <td>-50 to 260 mmHg (Adult/Neo) (10 mmHg steps)</td> <td>200 mmHg</td> <td>130 mmHg</td>	P1 SVS Upper Alarm Limits	-50 to 260 mmHg (Adult/Neo) (10 mmHg steps)	200 mmHg	130 mmHg
P1 SYS Lower Alarm Limits For the SY and predictive of the SYS and predictive of the SYS Lower Alarm Limits For the S		-6.7 to 34.7 kPa (Adult/Neo) (1.3 kPa steps)	200 mining 26 7 kPa	17.3 kPa
P1 Dio Lower Name Limits -80 to 23.3 kPa (AduliNeo) (1.3 kPa steps) P3 mPi 21.3 kPa 6.7 kPa P1 DIA Upper Alarm Limits -50 to 250 mmHg (AduliNeo) (1.3 kPa steps) 160 mmHg 100 mmHg P1 DIA Lower Alarm Limits -60 to 250 mmHg (AduliNeo) (1.3 kPa steps) 30 mmHg 100 mmHg P1 MEAN Upper Alarm Limits -60 to 250 mmHg (AduliNeo) (1.3 kPa steps) 30 mmHg 100 mmHg P1 MEAN Lower Alarm Limits -60 to 250 mmHg (AduliNeo) (1.3 kPa steps) 24.0 kPa 1.3 kPa P1 MEAN Lower Alarm Limits -60 to 250 mmHg (AduliNeo) (1.3 kPa steps) 24.0 kPa 1.4.7 kPa P1 MEAN Lower Alarm Limits -60 to 250 mmHg (AduliNeo) (1.3 kPa steps) 200 mmHg 20 mmHg P2 Lobel P2. CVP, PAP, LAP CVP CVP 22.7 kPa P2 SYS Upper Alarm Limits -50 to 260 mmHg (AduliNeo) (1.3 kPa steps) 200 mmHg 50 mmHg P2 DIA Lower Alarm Limits -60 to 250 mmHg (AduliNeo) (1.3 kPa steps) 200 mmHg 50 mmHg P2 DIA Lower Alarm Limits -60 to 250 mmHg (AduliNeo) (1.3 kPa steps) 30 mmHg 100 mmHg P2 MEAN Upper Alarm Limits -60 to 250 mmHg (AduliNeo) (1.3 kPa steps) 30 mmHg 100	P1 SVS Lower Alarm Limits	-60 to 250 mmHg (Adult/Neo) (10 mmHg steps)	70 mmHq	50 mmHa
P1 Di O US Di O US <thdio th="" us<=""> <thdi o="" th="" us<=""> <thdio td="" us<<=""><td></td><td>-80 to 33 3 kPa (Adult/Neo) (1 3 kPa steps)</td><td>0.3 kPa</td><td>67 kPa</td></thdio></thdi></thdio>		-80 to 33 3 kPa (Adult/Neo) (1 3 kPa steps)	0.3 kPa	67 kPa
11 Disk Opper Alarm Limits -6.7 to 34.7 KPa (Adul/Neo) (1.3 KPa steps) 100 mmHg 13.3 KPa 121 Di A Lower Alarm Limits -60 to 250 mmHg (Adul/Neo) (1.3 kPa steps) 30 mmHg 110 mmHg 121 MEAN Upper Alarm Limits -50 to 260 mmHg (Adul/Neo) (1.3 kPa steps) 30 mmHg 110 mmHg 121 MEAN Lower Alarm Limits -60 to 250 mmHg (Adul/Neo) (1.3 kPa steps) 40 mmHg 120 mmHg 121 MEAN Lower Alarm Limits -60 to 250 mmHg (Adul/Neo) (1.3 kPa steps) 40 mmHg 240 kPa 1.4.7 kPa 122 Label P2 (2 VP, PAP, LAP CVP CVP CVP 27 kPa 223 Stale 0-20, 0-50, 0-100, 0-200, 0-300, AUTO AUTO AUTO 200 mmHg 50 to 260 mmHg (Adul/Neo) (1.3 kPa steps) 200 mmHg 50 mmHg 22 Sys Lower Alarm Limits -60 to 250 mmHg (Adul/Neo) (1.3 kPa steps) 70 mmHg 50 mmHg 60 mmHg 67. kPa 22 DIA Lower Alarm Limits -60 to 250 mmHg (Adul/Neo) (1.3 kPa steps) 70 mmHg 13.3 kPa 13.3 kPa 22 MEAN Upper Alarm Limits -50 to 260 mmHg (Adul/Neo) (1.3 kPa steps) 30 mmHg 10 mmHg 40 to 3.3 kPa (Adul/Neo) (1.3 kPa steps) 30 mmHg 10 mmHg 24 MEAN Upper Alarm Limits -50 to 260 mmHg (Adul/Neo	P1 DIA LIpper Alarm Limits	-50 to 260 mmHg (Adult/Neo) (10 mmHg steps)	160 mmHg	100 mmHg
P1 DIA Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 30 mmHg 10 mmHg P1 MEAN Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (11 3 kPa steps) 40 kPa 13 kPa P1 MEAN Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 180 mmHg 110 mmHg P1 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (13 kPa steps) 24.0 kPa 147. kPa P1 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 5.3 kPa 2.7 kPa P2 Label P2 (CVP, PAP, LAP CVP CVP CVP P2 SYS Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 200 mmHg 130 mmHg P2 DIA Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 9.3 kPa 6.7 kPa P2 DIA Upper Alarm Limits -60 to 260 mmHg (Adult/Neo) (10 mmHg steps) 100 mmHg 100 mmHg P2 DIA Lower Alarm Limits -60 to 260 mmHg (Adult/Neo) (10 mmHg steps) 30 mHg 100 mmHg P2 MEAN Upper Alarm Limits -60 to 260 mmHg (Adult/Neo) (10 mmHg steps) 30 mmHg 100 mmHg P2 MEAN Upper Alarm Limits -60 to 260 mmHg (Adult/Neo) (10 mmHg steps) 100 mmHg 13. kPa		-6.7 to 34.7 kPa (Adult/Neo) (1.3 kPa steps)	21.3 kPa	13.3 kPa
P1 Dik Löwer Alam Limits -60 to 23.0 kPa (Adult/Neo) (1.3 kPa steps) 30 mHrg 10 mmHrg P1 MEAN Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (1.3 kPa steps) 120 mmHg 140 mmHg P1 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.3 kPa steps) 40 mmHg 22 mmHg P1 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.3 kPa steps) 5.3 kPa 2.7 kPa P2 Label P2, CVP, PAP, LAP CVP CVP CVP P2 Scale 0-20, 0-50, 0-100, 0-200, 0-300, AUTO AUTO AUTO AUTO P2 SYS Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) -70 mmHg 50 mmHg P2 DIA Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (1.3 kPa steps) 9.0 kPa 6.7 kPa P2 DIA Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.3 kPa steps) 100 mmHg 13.3 kPa P2 MEAN Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.1 kPa steps) 100 mmHg 13.3 kPa P2 MEAN Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.1 kPa steps) 10 mmHg 13.3 kPa P2 MEAN Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.1 kPa steps) 10 mmHg 13.3 kPa	P1 DIA Lower Alarm Limite	60 to 250 mmHg (Adult/Neo) (1.5 Ki a steps)	21.5 Ki d	10.5 Ki a
P1 MEAN Upper Alarm Limits -50 to 250 mmHg (Adult/Neo) (10 mmHg steps) 140 mmHg P1 MEAN Lower Alarm Limits -50 to 250 mmHg (Adult/Neo) (10 mmHg steps) 140 mmHg P1 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (13 kPa steps) 240 kPa 147 kPa P1 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (13 kPa steps) 5.3 kPa 27 kPa P2 Label P2, CVP, PAP, LAP CVP CVP CVP P2 State 0~20, 0~50, 0~100, 0~200, 0~300, AUTO AUTO AUTO AUTO P2 SYS Upper Alarm Limits -50 to 250 mmHg (Adult/Neo) (10 mmHg steps) -60 mHg 50 mmHg 50 mmHg P2 SYS Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 9.3 kPa 130 mmHg P2 DIA Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 160 mmHg 100 mmHg P2 DIA Lower Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mHg steps) 30 mmHg 100 mmHg P2 MEAN Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mHg steps) 30 mmHg 10 mmHg P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mHg steps) 100 mmHg 100 mHg P2	FT DIA LOWEI AIAITII LIITIIUS	-00 to 250 mining (Adult/Neo) (10 mining steps)		
P1 MEAN Opper Adam Limits -50 to 34.7 kPa (Adult/Neo) (1.3 kPa steps) 24.0 kPa P1 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.3 kPa steps) 24.0 kPa 14.7 kPa P2 Label P2, CVP, PAP, LAP CVP CVP P2 P2 Scale 0~20, 0~50, 0~100, 0~200, 0~300, AUTO AUTO AUTO AUTO P2 Sys Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.3 kPa steps) 200 mmHg 130 mmHg P2 SYS Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 70 mmHg 50 mmHg P2 DIA Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.3 kPa steps) 70 mmHg 100 mmHg P2 DIA Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.3 kPa steps) 21.3 kPa 13.3 kPa P2 MEAN Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.3 kPa steps) 24.0 kPa 1.3 kPa P2 MEAN Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.3 kPa steps) 21.3 kPa 13.3 kPa P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.3 kPa steps) 24.0 kPa 1.3 kPa P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.3 kPa steps) 24.0 kPa 1.4.7 kPa	D1 MEAN Linner Alerm Limite	-6.0 to 35.3 kPa (Adult/Neo) (1.3 kPa steps)	4.0 KFd	1.3 KFd
9.7 It 03-47. NP3 (AddIt/Neo) (1.3 NP3 steps) 24.0 NP3 14.7 NP3 P1 MEAN Lower Alarm Limits 60 to 250 mmHg (AddIt/Neo) (10 mmHg steps) 5.3 kPa 2.7 kPa P2 Label P2. CVP, PAP, LAP CVP CVP CVP P2 Soale 0-20, 0-50, 0-100, 0-200, 0-300, AUTO AUTO AUTO AUTO P2 SYS Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) -6.7 to 34.7 kPa (Adult/Neo) (1.3 kPa steps) 70 mmHg 50 mmHg P2 SYS Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.3 kPa steps) 9.3 kPa 6.7 kPa P2 DIA Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 160 mmHg 100 mmHg -8.0 to 33.3 kPa (Adult/Neo) (10 mmHg steps) 160 mmHg 100 mmHg 13.3 kPa P2 DIA Lower Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 30 mmHg 10 mmHg -8.0 to 33.3 kPa (Adult/Neo) (10 mmHg steps) 30 mmHg 10 mmHg 13.3 kPa P2 MEAN Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 40 mmHg 20 mmHg -6.7 to 34.7 kPa (Adult/Neo) (1.3 kPa steps) 24.0 kPa 1.4.7 kPa P2 MEAN Upper Alarm Limits<	PT MEAN Opper Alarm Limits	-50 to 200 mmHg (Adult/Nec) (10 mmHg steps)		
PT MEAN LOWER Atalin Limits -50 to 250 milling (Adult/Neo) (1.0 milling steps) 53 kPa 2.7 kPa P2 Label P2, CVP, PAP, LAP CVP CVP CVP P2 SyS Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (1.0 mmHg steps) -6.7 to 34.7 kPa AUTO AUTO P2 SYS Upper Alarm Limits -60 to 260 mmHg (Adult/Neo) (1.0 mmHg steps) -6.7 to 34.7 kPa AUTO MUTO AUTO P2 DIA Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 9.3 kPa 6.7 kPa P2 DIA Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 160 mmHg 100 mmHg P2 DIA Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.1 kPa steps) 21.3 kPa 13.3 kPa P2 MEAN Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (1.3 kPa steps) 4.0 kPa 1.3 kPa P2 MEAN Lower Alarm Limits -50 to 260 mmHg (Adult/Neo) (1.3 kPa steps) 4.0 mHg 100 mmHg P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.3 kPa steps) 5.3 kPa 2.7 kPa SpO2 - - - 6.7 to 34.7 kPa (Adult/Neo) (1.3 kPa steps) 9.0 % SpO2 - 0.0		-6.7 to 34.7 kPa (Adult/Neo) (1.3 kPa steps)	24.0 KPa	14.7 KPa
-8.0 ID 33.3 KPa (AdUINVEO) (1.3 KPa steps) 5.3 KPa 2.7 KPa P2 Label P2. CVP, PAP, LAP CVP CVP P2 Scale 0~20, 0~50, 0~100, 0~200, 0~300, AUTO AUTO AUTO P2 SYS Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 200 mmHg 130 mmHg P2 SYS Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (11 mmHg steps) 70 mmHg 50 mmHg P2 DIA Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (13 kPa steps) 9.3 kPa 6.7 kPa P2 DIA Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (13 kPa steps) 21.3 kPa 13.3 kPa P2 DIA Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.3 kPa steps) 30 mmHg 100 mmHg -8.0 to 33.3 kPa (Adult/Neo) (1.13 kPa steps) 24.0 kPa 1.3 kPa P2 MEAN Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.3 kPa steps) 24.0 kPa 14.7 kPa P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.1 kPa steps) 24.0 kPa 14.7 kPa P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.1 kPa steps) 20 mmHg 20 mmHg %SpO2 C-Lock On, Off Off Off <td>PT MEAN LOWER AIAIM LIMIUS</td> <td>-60 to 250 mmHg (Adult/Nec) (10 mmHg steps)</td> <td>40 mm⊓g</td> <td></td>	PT MEAN LOWER AIAIM LIMIUS	-60 to 250 mmHg (Adult/Nec) (10 mmHg steps)	40 mm⊓g	
P2 Eabel P2, CVP, PAP, LAP CVP CVP CVP P2 Scale 0~20, 0~50, 0~100, 0~200, 0~300, AUTO AUTO AUTO AUTO P2 SYS Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 200 mmHg 130 mmHg P2 SYS Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (13 kPa steps) 9.3 kPa 6.7 kPa P2 DIA Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (13 kPa steps) 9.3 kPa 6.7 kPa P2 DIA Lower Alarm Limits -50 to 250 mHg (Adult/Neo) (1.3 kPa steps) 160 mmHg 100 mmHg -8.0 to 33.3 kPa (Adult/Neo) (1.3 kPa steps) 4.0 kPa 1.3 kPa P2 MEAN Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (1.3 kPa steps) 180 mmHg 110 mmHg -8.0 to 33.3 kPa (Adult/Neo) (1.3 kPa steps) 4.0 kPa 1.3 kPa P2 MEAN Lower Alarm Limits -50 to 260 mmHg (Adult/Neo) (1.3 kPa steps) 5.3 kPa 2.7 kPa Sp0_2 -6.0 to 33.3 kPa (Adult/Neo) (1.3 kPa steps) 5.3 kPa 2.7 kPa Sp0_2 Lower Alarm Limits 70 to 100 % (Adult/Neo) (1.4 ksteps) 100 % 100 % %Sp0_2 Upper Alarm Limits 69 to 99 % (Adult/Neo) (1.4 ksteps) <t< td=""><td></td><td>-8.0 to 33.3 kPa (Aduit/Neo) (1.3 kPa steps)</td><td>5.3 KPa</td><td>2.7 KPa</td></t<>		-8.0 to 33.3 kPa (Aduit/Neo) (1.3 kPa steps)	5.3 KPa	2.7 KPa
P2 SYS Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) -200 mmHg 50 mmHg P2 SYS Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) -70 mmHg 50 mmHg P2 SYS Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (13 kPa steps) 9.3 kPa 6.7 kPa P2 DIA Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 100 mmHg 100 mmHg P2 DIA Lower Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 100 mmHg 100 mmHg P2 DIA Lower Alarm Limits -50 to 260 mmHg (Adult/Neo) (11 mmHg steps) 100 mmHg 100 mmHg -8.0 to 33.3 kPa (Adult/Neo) (10 mmHg steps) 4.0 kPa 1.3 kPa -8.0 to 260 mmHg (Adult/Neo) (10 mmHg steps) 4.0 kPa 1.3 kPa -8.0 to 260 mmHg (Adult/Neo) (10 mmHg steps) 4.0 kPa 1.3 kPa -8.0 to 260 mmHg (Adult/Neo) (11 mmHg steps) 4.0 kPa 1.3 kPa -8.0 to 260 mmHg (Adult/Neo) (11 mmHg steps) 140 mmHg 20 mmHg -60 to 250 mmHg (Adult/Neo) (11 mmHg steps) 100 mmHg 2.0 mmHg -8.0 to 33.3 kPa (Adult/Neo) (1.3 kPa steps) 5.3 kPa 2.7 kPa SpO2 Cotack On, Off				CVP
P2 SYS Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 200 mmHg 130 mmHg P2 SYS Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 70 mmHg 50 mmHg P2 DIA Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 160 mmHg 100 mmHg P2 DIA Lower Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 100 mmHg 100 mmHg P2 DIA Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 30 mmHg 10 mmHg P2 DIA Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 30 mmHg 10 mmHg P2 MEAN Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 4.0 kPa 1.3 kPa P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 40 mmHg 20 mmHg P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 40 mmHg 20 mmHg P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (13 kPa steps) 5.3 kPa 2.7 kPa SpO2		0~20, 0~50, 0~100, 0~200, 0~300, AUTO	AUTO	AUTO
-6.7 to 34.7 kFa (Adult/Neo) (1.3 kFa steps) 70 mmHg P2 SYS Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 70 mmHg 60 mMHg P2 DIA Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (13 kFa steps) 9.3 kFa 160 mmHg 100 mmHg P2 DIA Lower Alarm Limits -50 to 260 mmHg (Adult/Neo) (13 kFa steps) 160 mmHg 100 mmHg -6.7 to 34.7 kFa (Adult/Neo) (13 kFa steps) 30 mmHg 10 mmHg -8.0 to 33.3 kFa (Adult/Neo) (13 kFa steps) 4.0 kFa 1.3 kFa P2 MEAN Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (13 kFa steps) 4.0 kFa 14.7 kFa P2 MEAN Lower Alarm Limits -50 to 260 mmHg (Adult/Neo) (13 kFa steps) 4.0 kFa 14.7 kFa P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (13 kFa steps) 4.0 mmHg 20 mmHg -6.7 to 34.7 kFa (Adult/Neo) (10 mmHg steps) 5.3 kFa 2.7 kFa 26.0 mmHg P2 MEAN Lower Alarm Limits -60 to 250 mm/s 25.0 mm/s 25.0 mm/s %SpO2 - - - 0.0 ff 0ff 0.10 % (Adult/Neo) (1 % steps) 90 % 85 % 25.0 mm/s 25.0 mm/s	P2 SYS Upper Alarm Limits	-50 to 260 mmHg (Adult/Neo) (10 mmHg steps)	200 mmHg	130 mmHg
P2 SYS Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 70 mmHg 50 mmHg P2 DIA Upper Alarm Limits -50 to 250 mmHg (Adult/Neo) (11 mmHg steps) 9.3 kPa 6.7 kPa P2 DIA Lower Alarm Limits -50 to 250 mmHg (Adult/Neo) (13 kPa steps) 160 mmHg 100 mmHg P2 DIA Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 30 mmHg 100 mmHg P2 MEAN Upper Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 4.0 kPa 1.3 kPa P2 MEAN Lower Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 400 mmHg 20 mmHg P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 40 mmHg 20 mmHg P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (13 kPa steps) 40 mmHg 20 mmHg P2 MEAN Lower Alarm Limits -60 to 250 mm/s 25.0 mm/s 25.0 mm/s %SpO2 -8.0 to 33.3 kPa (Adult/Neo) (13 kPa steps) 40 mmHg 20 mmHg %SpO2 Upper Alarm Limits 70 to 100 % (Adult/Neo) (1 % steps) 100 % 100 % %SpO2 Lower Alarm Limits 69 to 99 % (Adult/Neo) (1 % steps) 90 % 85 % Respiration/A		-6.7 to 34.7 kPa (Adult/Neo) (1.3 kPa steps)		
-8.0 to 33.3 kPa (Adult/Neo) (1.3 kPa steps) 9.3 kPa 6.7 kPa P2 DIA Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 160 mmHg 100 mmHg -6.7 to 34.7 kPa (Adult/Neo) (1.3 kPa steps) 21.3 kPa 13.3 kPa 13.3 kPa P2 DIA Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.0 mmHg steps) 30 mmHg 10 mmHg -8.0 to 33.3 kPa (Adult/Neo) (1.1 kPa steps) 30 mmHg 10 mmHg 13. kPa P2 MEAN Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (1.0 mmHg steps) 4.0 kPa 1.3 kPa P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.0 mmHg steps) 24.0 kPa 14.7 kPa P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.3 kPa steps) 24.0 kPa 14.7 kPa P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.0 mmHg steps) 40 mmHg 20 mmHg P2 MEAN Lower Alarm Limits -60 to 250 mm/s (Adult/Neo) (1.3 kPa steps) 5.3 kPa 2.7 kPa SpO2 - - - 0 ff Off Off VSpO2 Upper Alarm Limits 70 to 100 % (Adult/Neo) (1 % steps) 100 % 85 % Respiration -	P2 SYS Lower Alarm Limits	-60 to 250 mmHg (Adult/Neo) (10 mmHg steps)	70 mmHg	50 mmHg
P2 DIA Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 160 mmHg P2 DIA Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.3 kPa steps) 30 mmHg P2 DIA Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 30 mmHg 110 mmHg P2 MEAN Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 180 mmHg 110 mmHg P2 MEAN Lower Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 180 mmHg 100 mmHg P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.3 kPa steps) 24.0 kPa 14.7 kPa P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.3 kPa steps) 40 mmHg 20 mmHg P2 MEAN Lower Alarm Limits -60 to 250 mm/s 25.0 mm/s 2.7 kPa SpO2 -6.7 to 33.3 kPa (Adult/Neo) (1.3 kPa steps) 100 m/Hg 20 mmHg %SpO2 Loper Alarm Limits 70 to 100 % (Adult/Neo) (1.3 kPa steps) 5.3 kPa 2.7 kPa SpD2 C-Lock On, Off Off Off 0ff VSpO2 Lower Alarm Limits 70 to 100 % (Adult/Neo) (1 % steps) 90 % 85 % Respiration -6.25, 12.5, 25.0 mm/s 12.5 mm/s 12.5 mm/s 12.5 mm/s <t< td=""><td></td><td>-8.0 to 33.3 kPa (Adult/Neo) (1.3 kPa steps)</td><td>9.3 kPa</td><td>6.7 kPa</td></t<>		-8.0 to 33.3 kPa (Adult/Neo) (1.3 kPa steps)	9.3 kPa	6.7 kPa
-6.7 to 34.7 kPa (Adult/Neo) (1.3 kPa steps) 21.3 kPa 13.3 kPa P2 DIA Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 30 mmHg 10 mmHg -8.0 to 33.3 kPa (Adult/Neo) (10 mmHg steps) 4.0 kPa 1.3 kPa P2 MEAN Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 180 mmHg 110 mmHg -6.7 to 34.7 kPa (Adult/Neo) (10 mmHg steps) 180 mmHg 110 mmHg -6.7 to 34.7 kPa (Adult/Neo) (10 mmHg steps) 4.0 kPa 14.7 kPa P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 5.3 kPa 2.7 kPa SpO2 -60 to 33.3 kPa (Adult/Neo) (1.3 kPa steps) 5.3 kPa 2.7 kPa <i>G-Lock</i> On, Off Off Off Off PLETH Sweep Speed 12.5, 25.0, 50.0 mm/s 25.0 mm/s 25.0 mm/s %SpO2 Upper Alarm Limits 70 to 100 % (Adult/Neo) (1 % steps) 100 % 100 % %SpO2 Lower Alarm Limits 69 to 99 % (Adult/Neo) (1 % steps) 90 % 85 % Respiration Apnea Time Setting Off, AUTO, awRR, imRR AUTO AUTO Respiration Size Graphic size bar (×	P2 DIA Upper Alarm Limits	-50 to 260 mmHg (Adult/Neo) (10 mmHg steps)	160 mmHg	100 mmHg
P2 DIA Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) -8.0 to 33.3 kPa (Adult/Neo) (1.3 kPa steps) 30 mmHg 4.0 kPa 11.3 kPa P2 MEAN Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (1.3 kPa steps) 180 mmHg 24.0 kPa 11.7 kPa P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.3 kPa steps) 24.0 kPa 14.7 kPa P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.3 kPa steps) 40 mmHg 20 mmHg P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.3 kPa steps) 40 mmHg 20 mmHg P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.3 kPa steps) 40 mmHg 20 mmHg P2 MEAN Lower Alarm Limits -60 to 250 mm/s (Adult/Neo) (1.3 kPa steps) 5.3 kPa 2.7 kPa SpO2		-6.7 to 34.7 kPa (Adult/Neo) (1.3 kPa steps)	21.3 kPa	13.3 kPa
-8.0 to 33.3 kPa (Adult/Neo) (1.3 kPa steps) 4.0 kPa 1.3 kPa P2 MEAN Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 180 mmHg 110 mmHg -6.7 to 34.7 kPa (Adult/Neo) (1.3 kPa steps) 24.0 kPa 14.7 kPa P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 40 mmHg 20 mmHg -8.0 to 33.3 kPa (Adult/Neo) (1.3 kPa steps) 5.3 kPa 2.7 kPa SpO2 -6.0 to 250 mmHg (Adult/Neo) (1.3 kPa steps) 5.3 kPa 2.7 kPa SpO2 -6.0 to 33.3 kPa (Adult/Neo) (1.3 kPa steps) 5.3 kPa 2.7 kPa SpO2 -6.0 to 33.3 kPa (Adult/Neo) (1.3 kPa steps) 5.3 kPa 2.7 kPa SpO2 -6.0 to 33.3 kPa (Adult/Neo) (1.3 kPa steps) 5.0 mm/s 2.5.0 mm/s %SpO2 Upper Alarm Limits 70 to 100 % (Adult/Neo) (1 % steps) 100 % 100 % %SpO2 Lower Alarm Limits 69 to 99 % (Adult/Neo) (1 % steps) 90 % 85 % Respiration -70 610.0 % 40.0 % 100 % Apnea Time Setting Off. 20, 30, 40, 50, 60 sec, Step 60, Step 90 30 sec 30 sec 30 sec Respiration Si	P2 DIA Lower Alarm Limits	-60 to 250 mmHg (Adult/Neo) (10 mmHg steps)	30 mmHg	10 mmHg
P2 MEAN Upper Alarm Limits -50 to 260 mmHg (Adult/Neo) (10 mmHg steps) 180 mmHg 110 mmHg P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (1.3 kPa steps) 24.0 kPa 14.7 kPa P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) 40 mmHg 20 mmHg BpO2 -8.0 to 33.3 kPa (Adult/Neo) (1.3 kPa steps) 5.3 kPa 2.7 kPa C-Lock On, Off Off Off Off PLETH Sweep Speed 12.5, 25.0, 50.0 mm/s 25.0 mm/s 25.0 mm/s %SpO2 Upper Alarm Limits 70 to 100 % (Adult/Neo) (1 % steps) 100 % 100 % %SpO2 Lower Alarm Limits 69 to 99 % (Adult/Neo) (1 % steps) 90 % 85 % Respiration		-8.0 to 33.3 kPa (Adult/Neo) (1.3 kPa steps)	4.0 kPa	1.3 kPa
-6.7 to 34.7 kPa (Adult/Neo) (1.3 kPa steps) 24.0 kPa 14.7 kPa P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) -8.0 to 33.3 kPa (Adult/Neo) (1.3 kPa steps) 40 mmHg 20 mmHg SpO2 -6.0 to 250 mm/s 0.13 kPa steps) 5.3 kPa 2.7 kPa C-Lock On, Off Off Off 0ff 0ff PLETH Sweep Speed 12.5, 25.0, 50.0 mm/s 25.0 mm/s 25.0 mm/s 25.0 mm/s %SpO2 Upper Alarm Limits 70 to 100 % (Adult/Neo) (1 % steps) 100 % 100 % %SpO2 Lower Alarm Limits 69 to 99 % (Adult/Neo) (1 % steps) 90 % 85 % Respiration	P2 MEAN Upper Alarm Limits	-50 to 260 mmHg (Adult/Neo) (10 mmHg steps)	180 mmHg	110 mmHg
P2 MEAN Lower Alarm Limits -60 to 250 mmHg (Adult/Neo) (10 mmHg steps) -8.0 to 33.3 kPa (Adult/Neo) (1.3 kPa steps) 40 mmHg 5.3 kPa 20 mmHg 2.7 kPa SpO2 C-Lock On, Off Off Off Off Off PLETH Sweep Speed 12.5, 25.0, 50.0 mm/s 25.0 mm/s 25.0 mm/s 25.0 mm/s 25.0 mm/s 25.0 mm/s %SpO2 Upper Alarm Limits 70 to 100 % (Adult/Neo) (1 % steps) 100 % 100 % 100 % %SpO2 Lower Alarm Limits 69 to 99 % (Adult/Neo) (1 % steps) 90 % 85 % Respiration Apnea Time Setting Off, 20, 30, 40, 50, 60 sec, Step 60, Step 90 30 sec 30 sec 30 sec Respiration/Apnea Off, AUTO, awRR, imRR AUTO AUTO AUTO Respiration Size Graphic size bar (×1/4, ×1/2, ×1, ×1.5, ×2) ×1 ×1 Respiration Size Graphic size bar (×1/4, ×1/2, ×1, ×1.5, ×2) ×1 ×1 Respiration Sweep Speed 6.25, 12.5, 25.0 mm/s 30 BPM 50 BPM RR Lower Alarm Limits 5 to 155 BPM (5 BPM steps) 0 BPM 0 BPM CAPNO Sweep Speed 6.25, 12.5, 25.0 mm/s		-6.7 to 34.7 kPa (Adult/Neo) (1.3 kPa steps)	24.0 kPa	14.7 kPa
-8.0 to 33.3 kPa (Adult/Neo) (1.3 kPa steps) 5.3 kPa 2.7 kPa SpO2	P2 MEAN Lower Alarm Limits	-60 to 250 mmHg (Adult/Neo) (10 mmHg steps)	40 mmHg	20 mmHg
SpO2 C-Lock On, Off Off Off PLETH Sweep Speed 12.5, 25.0, 50.0 mm/s 25.0 mm/s 25.0 mm/s %SpO2 Upper Alarm Limits 70 to 100 % (Adult/Neo) (1 % steps) 100 % 100 % %SpO2 Lower Alarm Limits 69 to 99 % (Adult/Neo) (1 % steps) 90 % 85 % Respiration		-8.0 to 33.3 kPa (Adult/Neo) (1.3 kPa steps)	5.3 kPa	2.7 kPa
C-Lock On, Off Off Off PLETH Sweep Speed 12.5, 25.0, 50.0 mm/s 25.0 mm/s 25.0 mm/s %SpO2 Upper Alarm Limits 70 to 100 % (Adult/Neo) (1 % steps) 100 % 100 % %SpO2 Lower Alarm Limits 69 to 99 % (Adult/Neo) (1 % steps) 90 % 85 % Respiration	SpO ₂			
PLETH Sweep Speed 12.5, 25.0, 50.0 mm/s 25.0 mm/s 100 % 101 % 110 % 110 % 110 % 110 % 110 % 110 % 110 % 110 % 110 % 110 % 110 % <th11 %<="" th=""> 110 % 110 %<td>C-Lock</td><td>On, Off</td><td>Off</td><td>Off</td></th11>	C-Lock	On, Off	Off	Off
%SpO2 Upper Alarm Limits 70 to 100 % (Adult/Neo) (1 % steps) 100 % 100 % %SpO2 Lower Alarm Limits 69 to 99 % (Adult/Neo) (1 % steps) 90 % 85 % Respiration Apnea Time Setting Off, 20, 30, 40, 50, 60 sec, Step 60, Step 90 30 sec 30 sec Respiration/Apnea Off, AUTO, awRR, imRR AUTO AUTO RUTO Respiration Size Graphic size bar (×1/4, ×1/2, ×1, ×1.5, ×2) ×1 ×1 Respiration Sweep Speed 6.25, 12.5, 25.0 mm/s 12.5 mm/s 12.5 mm/s RR Upper Alarm Limits 5 to 155 BPM (5 BPM steps) 30 BPM 50 BPM RR Lower Alarm Limits 0 to 150 BPM (5 BPM steps) 0 BPM 0 BPM CAPNO Sweep Speed 6.25, 12.5, 25.0 mm/s 12.5 mm/s 12.5 mm/s Scale 0~40, 0~60, 0~80, AUTO AUTO AUTO Capno On On On On Calibration Yes, No No No No EtCO2 Upper Alarm Limits 2 to 80 mmHg (Adult/Neo) (0.3 kPa steps) 10.7 kPa 10.7 kPa 0.3 to 10.5 %	PLETH Sweep Speed	12.5, 25.0, 50.0 mm/s	25.0 mm/s	25.0 mm/s
%SpO2 Lower Alarm Limits 69 to 99 % (Adult/Neo) (1 % steps) 90 % 85 % Respiration Apnea Time Setting Off, 20, 30, 40, 50, 60 sec, Step 60, Step 90 30 sec 30 sec 30 sec Respiration/Apnea Off, AUTO, awRR, imRR AUTO AUTO AUTO Respiration Size Graphic size bar (×1/4, ×1/2, ×1, ×1.5, ×2) ×1 ×1 Respiration Sweep Speed 6.25, 12.5, 25.0 mm/s 12.5 mm/s 12.5 mm/s 12.5 mm/s RR Upper Alarm Limits 5 to 155 BPM (5 BPM steps) 30 BPM 50 BPM RR Lower Alarm Limits 0 to 150 BPM (5 BPM steps) 0 BPM 0 BPM Capno Capno Capno No No No Scale 0~40, 0~60, 0~80, AUTO AUTO AUTO AUTO Capno Measurement On, Off On On On Calibration Yes, No No No No EtCO2 Upper Alarm Limits 2 to 80 mmHg (Adult/Neo) (0.3 kPa steps) 10.5 % 10.5 % 10.5 % EtCO2 Lower Alarm Limits 0 to 78 mmHg (Adult/Neo) (2 mmHg steps) <t< td=""><td>%SpO₂ Upper Alarm Limits</td><td>70 to 100 % (Adult/Neo) (1 % steps)</td><td>100 %</td><td>100 %</td></t<>	%SpO ₂ Upper Alarm Limits	70 to 100 % (Adult/Neo) (1 % steps)	100 %	100 %
Respiration Apnea Time Setting Off, 20, 30, 40, 50, 60 sec, Step 60, Step 90 30 sec 30 sec Respiration/Apnea Off, AUTO, awRR, imRR AUTO AUTO Respiration Size Graphic size bar (×1/4, ×1/2, ×1, ×1.5, ×2) ×1 ×1 Respiration Size Graphic size bar (×1/4, ×1/2, ×1, ×1.5, ×2) ×1 ×1 Respiration Sweep Speed 6.25, 12.5, 25.0 mm/s 12.5 mm/s 12.5 mm/s RR Upper Alarm Limits 5 to 155 BPM (5 BPM steps) 30 BPM 50 BPM RR Lower Alarm Limits 0 to 150 BPM (5 BPM steps) 0 BPM 0 BPM CAPNO Sweep Speed 6.25, 12.5, 25.0 mm/s 12.5 mm/s 12.5 mm/s Scale 0~40, 0~60, 0~80, AUTO AUTO AUTO Capno V VITO AUTO AUTO Capno Measurement On, Off On On On Capno Measurement On, Off On No No EtCO2 Upper Alarm Limits 2 to 80 mmHg (Adult/Neo) (2 mmHg steps) 80 mmHg 10.7 kPa 0.3 to 10.5 % (Adult/Neo) (0.3 kPa steps) 10.5 %<	%SpO ₂ Lower Alarm Limits	69 to 99 % (Adult/Neo) (1 % steps)	90 %	85 %
Apnea Time Setting Off, 20, 30, 40, 50, 60 sec, Step 60, Step 90 30 sec 30 sec Respiration/Apnea Off, AUTO, awRR, imRR AUTO AUTO Respiration Size Graphic size bar (×1/4, ×1/2, ×1, ×1.5, ×2) ×1 ×1 Respiration Sweep Speed 6.25, 12.5, 25.0 mm/s 12.5 mm/s 12.5 mm/s 12.5 mm/s RR Upper Alarm Limits 5 to 155 BPM (5 BPM steps) 30 BPM 50 BPM RR Lower Alarm Limits 0 to 150 BPM (5 BPM steps) 0 BPM 0 BPM CAPNO Sweep Speed 6.25, 12.5, 25.0 mm/s 12.5 mm/s 12.5 mm/s Scale 0~40, 0~60, 0~80, AUTO AUTO AUTO Capno 0 0, 0/60, 0~80, AUTO AUTO AUTO Capno Measurement 0n, Off On On On Capno Measurement 0, 0/76 0n No No EtCO2 Upper Alarm Limits 2 to 80 mmHg (Adult/Neo) (2 mmHg steps) 80 mmHg 80 mmHg 0.3 to 10.5 % (Adult/Neo) (0.3 kPa steps) 10.5 % 10.5 % 10.5 % EtCO2 Lower Alarm Limits 0 to 78 mmHg (Adult/Neo) (0.3 kPa s	Respiration			
Respiration/ApneaOff, AUTO, awRR, imRRAUTOAUTORespiration SizeGraphic size bar (×1/4, ×1/2, ×1, ×1.5, ×2)×1×1Respiration Sweep Speed6.25, 12.5, 25.0 mm/s12.5 mm/s12.5 mm/sRR Upper Alarm Limits5 to 155 BPM (5 BPM steps)30 BPM50 BPMRR Lower Alarm Limits0 to 150 BPM (5 BPM steps)0 BPM0 BPMCAPNO Sweep Speed6.25, 12.5, 25.0 mm/s12.5 mm/s12.5 mm/sScale0~40, 0~60, 0~80, AUTOAUTOAUTOCapno MeasurementOn, OffOnOnCalibrationYes, NoNoNoEtCO2 Upper Alarm Limits2 to 80 mmHg (Adult/Neo) (2 mmHg steps)80 mmHg0.3 to 10.5 % (Adult/Neo) (0.3 % steps)10.5 %10.5 %EtCO2 Lower Alarm Limits0 to 78 mmHg (Adult/Neo) (2 mmHg steps)0 mmHg0.4 kPa0.4 kPa0.4 kPa0 kPa0.4 kPa0.4 kPa0.4 kPa0.4 kPa0.4 kPa0.4 kPa0.4 kPa0.4 kPa	Apnea Time Setting	Off, 20, 30, 40, 50, 60 sec, Step 60, Step 90	30 sec	30 sec
Respiration SizeGraphic size bar (×1/4, ×1/2, ×1, ×1.5, ×2)×1×1Respiration Sweep Speed $6.25, 12.5, 25.0 \text{ mm/s}$ 12.5 mm/s 12.5 mm/s 12.5 mm/s RR Upper Alarm Limits $5 \text{ to } 155 \text{ BPM (5 BPM steps)}$ 30 BPM 50 BPM RR Lower Alarm Limits $0 \text{ to } 150 \text{ BPM (5 BPM steps)}$ 0 BPM 0 BPM CAPNOCapno 0 Capno 0 Capno 12.5 mm/s CAPNO Sweep Speed $6.25, 12.5, 25.0 \text{ mm/s}$ 12.5 mm/s 12.5 mm/s Scale $0^{\sim}40, 0^{\sim}60, 0^{\sim}80, \text{ AUTO}$ AUTOAUTOCapno MeasurementOn, OffOnOnCalibrationYes, NoNoNoEtCO2 Upper Alarm Limits $2 \text{ to } 80 \text{ mmHg}$ (Adult/Neo) ($2 \text{ mmHg steps$) 80 mmHg $0.3 \text{ to } 10.7 \text{ kPa}$ 0.3 steps) 10.5% 10.7 kPa 0.5% $0 \text{ to } 78 \text{ mmHg}$ (Adult/Neo) ($2 \text{ mmHg steps$) 0 mmHg 0 mmHg $0 \text{ to } 78 \text{ mmHg}$ (Adult/Neo) (0.3 kPa steps) 0 mmHg 0 mmHg	Respiration/Apnea	Off, AUTO, awRR, imRR	AUTO	AUTO
Respiration Sweep Speed 6.25, 12.5, 25.0 mm/s 12.5 mm/s 12.5 mm/s 12.5 mm/s RR Upper Alarm Limits 5 to 155 BPM (5 BPM steps) 30 BPM 50 BPM RR Lower Alarm Limits 0 to 150 BPM (5 BPM steps) 0 BPM 0 BPM Capno 0 Capno 12.5 mm/s 12.5 mm/s CAPNO Sweep Speed 6.25, 12.5, 25.0 mm/s 12.5 mm/s 12.5 mm/s Scale 0~40, 0~60, 0~80, AUTO AUTO AUTO Capno Measurement On, Off On On Calibration Yes, No No No No EtCO2 Upper Alarm Limits 2 to 80 mmHg (Adult/Neo) (2 mmHg steps) 80 mmHg 10.7 kPa 10.7 kPa 0.3 to 10.5 % (Adult/Neo) (0.3 % steps) 10.5 % 10.5 % 0.5 % 0.5 %	Respiration Size	Graphic size bar (×1/4, ×1/2, ×1, ×1.5, ×2)	×1	×1
RR Upper Alarm Limits 5 to 155 BPM (5 BPM steps) 30 BPM 50 BPM RR Lower Alarm Limits 0 to 150 BPM (5 BPM steps) 0 BPM 0 BPM 0 BPM Capno Capno 12.5 mm/s 12.5 mm/s 12.5 mm/s 12.5 mm/s Scale 0~40, 0~60, 0~80, AUTO AUTO AUTO AUTO AUTO Capno Measurement On, Off On On On On Calibration Yes, No No No No No EtCO2 Upper Alarm Limits 2 to 80 mmHg (Adult/Neo) (2 mmHg steps) 80 mmHg 10.7 kPa 10.7 kPa 0.3 to 10.5 % (Adult/Neo) (0.3 % steps) 10.5 % 10.5 % 0.5 % 0.5 %	Respiration Sweep Speed	6.25, 12.5, 25.0 mm/s	12.5 mm/s	12.5 mm/s
RR Lower Alarm Limits 0 to 150 BPM (5 BPM steps) 0 BPM 0 BPM 0 BPM Capno CAPNO Sweep Speed 6.25, 12.5, 25.0 mm/s 12.5 mm/s 12.5 mm/s 12.5 mm/s Scale 0~40, 0~60, 0~80, AUTO AUTO AUTO AUTO On On Capno Measurement On, Off On On No No No No No EtCO2 Upper Alarm Limits 2 to 80 mmHg (Adult/Neo) (2 mmHg steps) 0.3 to 10.7 kPa (Adult/Neo) (0.3 kPa steps) 10.7 kPa 10.7 kPa 10.7 kPa 10.5 % 10.5 % 0 mmHg 0 to 78 mmHg (Adult/Neo) (2 mmHg steps) 0 mmHg	RR Upper Alarm Limits	5 to 155 BPM (5 BPM steps)	30 BPM	50 BPM
Capno CAPNO Sweep Speed 6.25, 12.5, 25.0 mm/s 12.5 mm/s 12.5 mm/s Scale 0~40, 0~60, 0~80, AUTO AUTO AUTO Capno Measurement On, Off On On Calibration Yes, No No No EtCO2 Upper Alarm Limits 2 to 80 mmHg (Adult/Neo) (2 mmHg steps) 80 mmHg 80 mmHg 0.3 to 10.7 kPa (Adult/Neo) (0.3 kPa steps) 10.7 kPa 10.7 kPa 0.3 to 10.5 % (Adult/Neo) (0.3 % steps) 10.5 % 10.5 % 0.5 % EtCO2 Lower Alarm Limits 0 to 78 mmHg (Adult/Neo) (2 mmHg steps) 0 mmHg 0 mmHg	RR Lower Alarm Limits	0 to 150 BPM (5 BPM steps)	0 BPM	0 BPM
CAPNO Sweep Speed 6.25, 12.5, 25.0 mm/s 12.5 mm/s 12.5 mm/s Scale 0~40, 0~60, 0~80, AUTO AUTO AUTO Capno Measurement On, Off On On Calibration Yes, No No No EtCO2 Upper Alarm Limits 2 to 80 mmHg (Adult/Neo) (2 mmHg steps) 80 mmHg 80 mmHg 0.3 to 10.7 kPa (Adult/Neo) (0.3 kPa steps) 10.7 kPa 10.7 kPa 0.3 to 10.5 % (Adult/Neo) (0.3 % steps) 10.5 % 10.5 % EtCO2 Lower Alarm Limits 0 to 78 mmHg (Adult/Neo) (2 mmHg steps) 0 mmHg 0 to 10.4 kPa (Adult/Neo) (0 3 kPa steps) 0 mmHg 0 mmHg	Capno			-
Scale 0~40, 0~60, 0~80, AUTO AUTO AUTO Capno Measurement On, Off On On On Calibration Yes, No No No No EtCO2 Upper Alarm Limits 2 to 80 mmHg (Adult/Neo) (2 mmHg steps) 80 mmHg 80 mmHg 0.3 to 10.7 kPa (Adult/Neo) (0.3 kPa steps) 10.7 kPa 10.7 kPa 0.3 to 10.5 % (Adult/Neo) (0.3 % steps) 10.5 % 10.5 % 0 mmHg EtCO2 Lower Alarm Limits 0 to 78 mmHg (Adult/Neo) (2 mmHg steps) 0 mmHg 0 mmHg	CAPNO Sweep Speed	6.25, 12.5, 25.0 mm/s	12.5 mm/s	12.5 mm/s
Capno MeasurementOn, OffOnOnCalibrationYes, NoNoNoEtCO2 Upper Alarm Limits2 to 80 mmHg (Adult/Neo) (2 mmHg steps)80 mmHg0.3 to 10.7 kPa (Adult/Neo) (0.3 kPa steps)10.7 kPa0.3 to 10.5 % (Adult/Neo) (0.3 % steps)10.5 %EtCO2 Lower Alarm Limits0 to 78 mmHg (Adult/Neo) (2 mmHg steps)0 mmHg0 to 78 mmHg (Adult/Neo) (2 mmHg steps)0 mmHg0 to 10.4 kPa (Adult/Neo) (0.3 kPa steps)0 mmHg0 to 78 mmHg (Adult/Neo) (0.3 kPa steps)0 mmHg0 to 10.4 kPa (Adult/Neo) (0.3 kPa steps)0 kPa	Scale	0~40 0~60 0~80 AUTO	AUTO	AUTO
Calibration Yes, No No No EtCO2 Upper Alarm Limits 2 to 80 mmHg (Adult/Neo) (2 mmHg steps) 80 mmHg 80 mmHg 0.3 to 10.7 kPa (Adult/Neo) (0.3 kPa steps) 10.7 kPa 10.7 kPa 0.3 to 10.5 % (Adult/Neo) (0.3 % steps) 10.5 % 10.5 % EtCO2 Lower Alarm Limits 0 to 78 mmHg (Adult/Neo) (2 mmHg steps) 0 mmHg 0 to 10.4 kPa (Adult/Neo) (2 mmHg steps) 0 mmHg 0 mmHg	Cappo Measurement	On Off	On	On
EtCO2 Upper Alarm Limits 2 to 80 mmHg (Adult/Neo) (2 mmHg steps) 80 mmHg 80 mmHg 0.3 to 10.7 kPa (Adult/Neo) (0.3 kPa steps) 10.7 kPa 10.7 kPa 0.3 to 10.5 % (Adult/Neo) (0.3 % steps) 10.5 % 10.5 % 10.5 % EtCO2 Lower Alarm Limits 0 to 78 mmHg (Adult/Neo) (2 mmHg steps) 0 mmHg 0 mmHg	Calibration	Yes No	No	No
Etco2 Copper Alarm Limits 2 to comming (Adult/Neo) (2 mming steps) oo mming (Adult/Neo) 0.3 to 10.7 kPa (Adult/Neo) (0.3 kPa steps) 10.7 kPa 10.7 kPa 0.3 to 10.5 % (Adult/Neo) (0.3 % steps) 10.5 % 10.5 % 10.5 % EtcO2 Lower Alarm Limits 0 to 78 mmHg (Adult/Neo) (2 mmHg steps) 0 mmHg 0 mmHg 0 to 10.4 kPa (Adult/Neo) (0 3 kPa steps) 0 mmHg 0 kPa	EtCO ₂ Upper Alarm Limite	2 to 80 mmHg (Adult/Neo) (2 mmHg stops)	80 mm ^L a	
0.3 to 10.7 kFa 10.7 kFa 10.7 kFa 0.3 to 10.5 % (Adult/Neo) (0.3 % steps) 10.5 % 10.5 % EtCO ₂ Lower Alarm Limits 0 to 78 mmHg (Adult/Neo) (2 mmHg steps) 0 mmHg 0 to 10.4 kPa (Adult/Neo) (0.3 kPa steps) 0 mmHg 0 kPa		2 to 00 mining (Adult/Neo) (2 mining steps) 0.3 to 10.7 kPa (Adult/Neo) (0.3 kPa steps)		
0.5 to 10.5 % (Addit/Neo) (0.5 % steps) 10.5 % EtCO2 Lower Alarm Limits 0 to 78 mmHg (Addit/Neo) (2 mmHg steps) 0 mmHg 0 to 10.4 kPa (Addit/Neo) (0.3 kPa steps) 0 kPa 0 kPa		$0.3 \text{ to } 10.5 \% (\Delta \text{dult/Nec}) (0.3 \% \text{steps})$	10.7 KFa	10.7 KFa
$\begin{bmatrix} 1002 \text{ Lower Aranne Linning} & 0 to 70 mining (Addit/Neo) (2 mining steps) & 0 mining & 0 mining \\ 0 to 10.4 kPa (Addit/Neo) (0.3 kPa steps) & 0 kPa & 0 kPa \\ \end{bmatrix}$	EtCO ₂ Lower Alarm Limits	0 to 78 mmHg (Adult/Neo) (2 mmHg stops)		
TO TO TO A KEALADO TO TA KEA MADELE TE TERES TE TERES		0 to 10.4 kPa (Adult/Neo) (0.3 kPa steps)	0 kPa	0 kPa

Parameter	Ranges/Selections	Factory Defaults	
	Rangeo, concento	Adult	Neonatal
	0 to 10.3 % (Adult/Neo) (0.3 % steps)	0 %	0 %
InCO ₂ Upper Alarm Limits	2 to 20 mmHg (Adult/Neo) (2 mmHg steps)	20 mmHg	20 mmHg
	0.3 to 2.7 mmHg (Adult/Neo) (0.3 kPa steps)	2.7 kPa	2.7 kPa
	0.3 to 2.6 % (Adult/Neo) (0.3 % steps)	26%	26%
InCO. Lower Alarm Limits	0 to 18 mmHg (Adult/Neo) (2 mmHg steps)	0 mmHg	0 mmHg
		0 KPa	0 кРа
-	0 to 2.3 % (Adult/Neo) (0.3 % steps)	0 %	0 %
Temperature			
T1, T2 Upper Alarm Limits	15.0 to 45.5 °C (Adult/Neo) (0.5° C steps)	38.0 °C	39.0 °C
	59.0 to 113.9 °F (Adult/Neo) (0.1°F steps)	(100.4 ° F)	(102.2 ° F)
T1, T2 Lower Alarm Limits	14.5 to 45.0 °C (Adult/Neo) (0.5° C steps)	14.5 °F	14.5 °F
	58.1 to 113.0 °F (Adult/Neo) (0.1°F steps)	(58.1 °F)	(58.1 °F)
Others		, , , , , , , , , , , , , , , , , , ,	
Patient Mode	Adult, Neonatal	Adı	,lt
Record Speed**	25.0 mm/s 50.0 mm/s	25.0 n	nm/s
Wave Record Time**	20 sec. Continuous (10 sec delay)	20 s	ec.
	ECG1 + ECG2, PLETH, RESP, IBP1, IBP2 or		
Wave Record Select**	CAPNO	ECG1 +	PLETH
Record on Alarm**	On,Off	Of	f
Auto List Record**	On,Off	Ot	ff
Alarm Volume	1, 2, 3, 4, 5, 6, 7, 8 (45to85dB)	5	
HR/PR tone Volume	Off, 1, 2, 3, 4, 5, 6, 7	4	
Key Beep Volume	Off, 1, 2, 3, 4, 5, 6, 7	4	
Sleep Mode	Off, 10, 20, 30 min	Ot	f
Main Screen	4ch-wave, 6ch-wave, Big Number	-	
Alarm Limits Display	On, Off	On	
Auto Alarm	On, Off	Ot	ff
Auto Alarm Setting (Upper)	+10 to +50%	+40%	
Auto Alarm Setting (Lower)	-50 to -10%	-20	%
Save Time Interval	Off. 0.5.1.2.2.5.5.10.15.20.30.60.120 min	Ot	f
Graphical Display On/Off	On/Off for each parameter	O	า
Save Setting on Power Off*	Custom, Back up, Default	Back	
Audible Alarm Silence Period*	30, 60, 90, 120 sec	120	sec
Audible Alarm Suspend	Off, 10, 20, 30, 60 min, Indefinite (Alarm	1	
Period*	Inhibition)	Indef	inite
AC Line Frequency*	50, 60 Hz	60 I	Ηz
NIBP Unit*	mmHg, kPa	mm	Hg
IBP Unit*	mmHg, kPa	mm	Hg
CO ₂ Unit*`	mmHg, %, kPa	mm	Hg
Temperature Unit*	°C , °F	°C	
HR/PR Tone Set*	High, Med, Low, SpO ₂	High	
Date Format*	year/month/day, month/day/year, day/month/year	year/month/day	
Jog Dial Speed	Fast, Normal	Norr	nal
	(Korean), 中文 (Chinese), Dansk (Danish), Nederlands (Dutch), English, Français		
Language*	(French), Deutsch (German), Italiano (Italian), 日本語 (Japanese), Português (Portuguese)	English	
	Español (Spanish), Svenska (Swedish)		
Note: An asterisk (*) by a parameter in the above table indicates that the parameter can only be changed by authorized personnel as described in the service manual			

Note: Asterisks (**) by a parameter in the above table indicate the settings only when an optional recorder is installed in the monitor.

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SPECIFICATION

Display

Screen Size	10.4" measured diagonally across the TFT-LCD screen
Screen Type/Color	Liquid Crystal Display (LCD) Color,
	Cold Cathode Fluorescent Backlit
Resolution	800 × 600 pixel
Number of Traces	4 or 6 waveforms

Controls

	Jog dial control;
Standard	/ soft buttons (NIBP start/stop, Main screen, NIBP interval,
	Alarm silence, menu, Record, Power on/oir)

Alarms

Categories	Patient Status and System Status	
Priorities	Low, Medium and High Priorities	
Notification	Audible and Visual	
Setting	Default and Individual	

Physical Characteristics and Recorder

Instrument		
Dimensions	360 × 280 × 215 (mm) (W×H×D)	
	including a handle and excluding options and accessories	
Weight	5.5 kg excluding optional configurations and accessories	
Degree of Protection	ECG:	Type CF with defibrillator protection
against Electric	IBP(P1-P2):	Type CF with defibrillator protection
Shock	NIBP:	Type CF with defibrillator protection
	SpO2:	Type CF with defibrillator protection
	Temperature	Type CF with defibrillator protection
	(T1-T2):	
	Gas(CO2):	Type CF with defibrillator protection
Mode of Operation	Continuous	
Classification	Class IIb (MDD An	nex IX Rule10:MEDDEV 2.4/1 Rev.8)
Recorder (Optional)		
Туре	Thermal	
Weight	150 g	
Resolution	8 dot/mm	
Number of Channels	1 to 2 channels	
Paper Type	Thermal	
Paper Width	50 mm	
Paper Speeds	25.0 mm/s and 50.0) mm/s

Electrical

Instrument		
Power Requirements	ower Requirements AC Mains	
	100-240V~50/60 Hz, 0.5-1.3A	
Fuses	q'ty 2, T6.3 A, 250 volts, (time-lag), IEC (5×20 mm)	
	Battery	
A battery typically provides operating time of 1 hour when fully charged with no		
printing, no external communication, no audible alarm sound and one NIBP		
measurement per 5 minu	ites at 25°C.	
Туре	Ni-MH	
Voltage/Capacity	12 V/ 3.8 Ampere-Hours	
Recharge	Over 12 hours with monitor turned on/off	
Shelf Life	2 years, new battery fully-charged	
Compliance	91/157/EEC	

Environmental Conditions

Operation		
Temperature	10°C to 40°C (50°F to 104°F)	
Humidity	15 % RH to 90% RH, non-condensing	
Altitude	500hPa to 1060hPa	
Transport and Storage (in shipping container)		
Temperature -20°C to 60°C (-4°F to 140°F)		
Humidity	15 % RH to 95% RH, non-condensing	
Altitude	500hPa to 1060hPa	
Note: The system may not meet its performance specifications if stored or used		
outside the specified temperature and humidity range.		

Measurement Parameters

ECG

Heart Rate		
Measurement Range	20 BPM ~ 250 BPM	
Accuracy	±3 BPM or ±5% whichever is greater	
ECG (Electrocardiograph)		
Leads	3 / 5 Lead	
	Lead I, II, III, aVR, aVL, aVF, V (Chest Lead)	
Lead Off Detection	Detected and displayed	
Input		
Input Dynamic Range	±5 mV AC, ±300 mV DC	
Voltage Range	±0.5 mV ~ ±5 mV	
Signal Width	40 ms ~ 120 ms (Q to S)	
Output		
Frequency Response (Ban	dwidth)	
Low Extend	0.05 Hz ~ 40 Hz	
Filter	0.5 Hz ~ 30 Hz	
Monitor	0.5 Hz ~ 40 Hz	
Respiration Rejection	1 Hz ~ 40 Hz	
ECG Size	×1/4, ×1/2, ×1, ×1.5, ×2	
Display Sweep Speeds	12.5 mm/sec, 25.0 mm/sec, and 50.0 mm/sec	
Defibrillator Discharge Recovery	<5 sec per IEC60601-2-27>	

Respiration

im RR		
Technique	Trans-thoracic impedance	
Range	0, 3 to 120 breaths/min	
Accuracy	±3 breaths/min	
Leads	RA to LA	
Display Sweep Speeds	6.25 mm/s, 12.5 mm/s, 25.0 mm/s	
Lead Off Condition	Detected and displayed	
aw RR (Option)		

Technique	Nondispersive Infrared Spectroscopy
Range	0 to 120 breaths/min
Accuracy	±1 breath/min
Display Sweep Speeds	6.25 mm/s, 12.5 mm/s, 25.0 mm/s

NIBP

Pulse Rate		
Pulse Rate Range	Adult 40 BPM ~ 200 BPM	
	Neonatal 40 BPM ~ 240 BPM	
Pulse Rate Accuracy	±2 BPM or ±2%, whichever is greater	
NIB	P (Non-Invasive Blood Pressure)	
Technique	Oscillometric Measurement	
Measurement Modes	MANUAL, AUTO and CONT	
MANUAL Mode	Single measurement initiated by NIBP Start/Stop button	
AUTO Mode	Automatic BP measurements at intervals of 1, 2, 2.5, 3, 5, 10, 15, 20, 30, 45, 60, 90, 120, or 180 minutes	
CONT Mode	Series of consecutive measurements for 5 minutes	
NIBP Pressure Measuring R	ange	
Systolic Pressure Range	Adult 60 mmHg ~ 250 mmHg	
	Neonatal 40 mmHg ~ 120 mmHg	
Diastolic Pressure Range	Adult 40 mmHg ~ 200 mmHg	
	Neonatal 20 mmHg ~ 90 mmHg	
MAP Pressure Range	Adult 45 mmHg ~ 235 mmHg	
	Neonatal 30 mmHg ~ 100 mmHg	
NIBP Accuracy	Mean error and standard deviation per ANSI/AAMI SP10:2002	
Pressure Display Range	10 mmHg ~ 300 mmHg	
Pressure Display Accuracy	±3mmHg	
Initial Cuff Inflation	Adult 120, 140, 160, 180, 200, 220, mmHg	
	(16.0, 18.7, 21.3, 24.0, 26.7, 29.3, kPa)	
	Neonatal 80, 100, 120, 140 mmHg	
	(9.3, 12.0, 14.7, 16.0, 18.7 kPa)	
Automatic Cuff Deflation	Measurement time exceeding 180s in adult/pediatric (90s	
	in neonatal) or maximum pressure value exceeding 300	
	mmHg in adult (150 mmHg in neonatal).	
Overpressure Protector	300 ±10 mmHg for Adult	
	150 ±5 mmHg for neonatal	

SpO₂

Pulse Rate		
Range	20 BPM ~ 250 BPM	
Accuracy	Without Interference	20 BPM ~ 250 BPM ±3 digits
	Low Perfusion	20 BPM ~ 250 BPM ±3 digits
	SpO ₂	
Range	1% to 100%	
Low Perfusion	0.03% to 20%	
Accuracy	Without Interference-Ad	ult
		70% ~ 100% ±2 digits
		1% ~ 69% unspecified
	Without Interference-Ne	onate
		70% ~ 100% ±3 digits
		1% ~ 69% unspecified
	Low Perfusion	70% ~ 100% ±2 digits
		1% ~ 69% unspecified
Display Sweep Speeds	12.5 mm/sec, 25.0 mm/s	sec and 50.0 mm/sec
Neonatal specifications ar	e shown for neonatal sen	sors with the monitor. Saturation
accuracy will vary by sensor type as specified by the manufacturer.		

Note: The wavelength range of the light emitted are near 660 nm and 890 nm with the energy not exceeding 15mW.

Temperature

Probe Type	Thermistor probe YSI 400 series	
Parameter Displayed	T1, T2	
Range	15°C ~ 45°C (59°F to 113°F)	
Display Accuracy	±0.1°C (25°C to 45°C) or ±0.2°F (77°F to 113°F)	
	±0.2°C (15°C to less than 25°C)	
	or ±0.4°F (59°F to less than 77°F)	
Probe Accuracy	±0.1°C (±0.2°F)	

IBP

Pulse Rate			
Range	20 BPM ~ 250 BPM		
Accuracy	±1% or ± 1 beat		
	IBP		
Parameter Displayed	P1, ABP		
	P2 , CVP, PAP, LAP		
Pressure Measuring Range	-50 mmHg ~ 300 mmHg		
Input Impedance	More than 1 M ohm		
Transducer Driving Voltage	DC 5V		
Transducer Input Sensitivity	5uV/V/mmHg		
Zero Calibration Range	± 100mmHg		
Zero Calibration Accuracy	Less than ±1mmHg		
Frequency Characteristics	dc to 25Hz		
Pressure Display Accuracy	Monitor: Less than ±3mmHg		
Scale	P1 0~50, 0~100, 0~200, 0~300, AUTO		
	P2 0~20, 0~50, 0~100, 0~200, 0~300, AUTO		
Display Sweep Speeds	12.5 mm/sec, 25.0 mm/sec, and 50.0 mm/sec		

Capnography

Parameter Displayed	EtCO ₂ , InCO ₂	
Range	0 mmHg ~ 100 mmHg; 0 kPa ~ 13.3 kPa; 0 % ~10 % CO ₂	
	STPD (standard temperature and pressure dry)	
Accuracy	±2mmHg or ± 4%, whichever is greater	
Stability	< 0.3 % (vol) CO ₂ /24hrs	
Rise Time	400 ms (average)	
Delay Time	1.8 (average)	
System Response	2.1 (average)	
Warm Up Time	3 minutes average	
Display Sweep Speeds	6.25mm/sec, 12.5 mm/sec and 25.0 mm/sec	
Sound Noise Level	Less than 41dB when ambient sound pressure level is 22dB	
Rise Time of Flow Rate	1.0 sec at normal condition	
at flow rate of 10 l/min	2.0 sec after sustained pressure	
Fall Time of Flow Rate	1.2 sec at normal condition	
at flow rate of 10 l/min	2.2 sec after sustained pressure	

Trends

Types	Graphical and Tabular	
Memory	saves total 1500 data	
	saves at selected time interval	
	saves alarm condition & error events	
	saves NIBP Measuremets	
Graphical Format	Total 2 graphs	
	a graph for NIBP, P1/P2, SpO ₂ , T1/T2 parameters	
	a graph for HR/PR, Resp, EtCO ₂ parameters	
	User-selectable each parameter to be desired	
Tabular Format	One table for all parameters	
Display	8 lists	
Save Time Interval	30sec or 1, 2, 2.5, 5, 10, 15, 20, 30, 60 or 120 minutes	

Compliance

ltem	Compliant with		
Classification	Class I (on AC power) Internally powered (on battery power)		
Type of protection	Type CF – Applied part		
Mode of operation	Continuous		
Degree of protection	IP21 (provided by enclosures)		
General	93/42/EEC Directives for medical devices		
	21CFR820 Code of Federal Regulations		
	2002/96/EC Waste electrical and electronic equipment Directive(WEEE)		
	91/157/EEC Battery Declaration Directive		
	93/86/EEC Battery Disposal Directive		
	ISO9001:2000 Quality Management Systems - Requirements		
	ISO13485:1996 Quality Systems– Medical Devices –Particular requirements		
	for the application of ISO9001		
	ISO14971:2000 Risk analysis managements – medical devices		
	IEC60601-1:1988+A1:1991+A2:1995		
	General requirements for Safety and Essential Performance		
	IEC60529:2001 Degree of Protection Provided by Enclosures (IP21)		
	EN ISO14155-1:2003 Clinical investigation of medical devices for human		
	subjects – part 1: General requirements		
	AAMI HE48:1993 Human factors engineering guidelines and preferred		
	practices for the design of medical devices		
	IEC60601-1-1:2000 Safety requirements for medical electrical systems		
	IEC60601-1-4:2000		
	Particular requirements for programmable medical systems		
	IEC60601-1-6:2004 Medical electrical equipment Part 1-6: General		
	requirements for safety Collateral Standard: Usability		
	ISO10993-1:2003 Biological evaluation of medical devices – Part 1:		
	Evaluation and testing		
	IEC60601-2-49:2001		
	Particular requirements for multifunction patient monitoring equipment		
Alarms	IEC60601-1-8:2003 Alarm systems requirements, tests and guidances in		
	medical electrical equipments systems		
Electrocardiograph	IEC60601-2-27:1994 Particular requirements for the safety of		
	Electrocardiographic monitoring equipment		
	AAMI EC13:2002 Cardiac monitors, heart rate meters and alarms		
	AAMI EC53:1995+A1:1998 ECG cable and leads		
Non-invasive blood	AAMI SP10:2002+A1:2003 Electronic or Automated Sphygmomanometers		
pressure	EN1060-1:1995+A1:2002 Non-invasive sphygmomanometers		
	EN1060-3:1997 Supplementary requirements for electrical-mechanical blood		
	pressure measuring systems		
	essential performance, of automatic cycling indirect blood pressure monitoring		
	equipment		
Oxvgen saturation	EN865:1997 Pulse oximeters. Particular requirements		
	ISO9919:2005 Basic safety & essential performance of Pulse oximeter for		
	medical use		
Temperature	EN12470-4:2000		
monitorina	Performance of Electrical Thermometers for continuous Measurement		

Item	Compliant with		
Invasive blood	IEC60601-2-34:2000		
pressure	Particular requirements for the safety, including essential performance, of		
	invasive blood pressure monitoring equipment		
Capnography	EN864:1996 Capnometers for use with humans-Particular requirements		
	ISO9918:1993 Capnometers for use with humans-Requirements		
	ISO21647:2004		
	Particular requirements for the basic safety and essential performance of		
	respiratory gas monitors		
Electromagnetic	IEC60601-1, sub clause 36, IEC/		
compatibility	IEC60601-1-2:2001+A1:2004		
	Electromagnetic compatibility-requirements & test		
	IEC61000-3-2:2005 Harmonic Emission Ed 3.0		
	IEC61000-3-3:2005 Voltage Fluctuations/Flicker Emission Ed 1.2		
	IEC61000-4-2:2001 Electrostatic Discharge Ed 1.2		
	IEC61000-4-3:2002 Radiated RF electromagnetic field Ed 2.1		
	IEC61000-4-4:2004 Electrical fast transient/burst Ed 2.0		
	IEC61000-4-5:2005 Surge current Ed 2.0		
	IEC61000-4-6:2004 Conducted disturbances, induced by RF field Ed 2.1		
	IEC61000-4-8:2001 Power frequency (50/60Hz) magnetic field Ed 1.1		
	IEC61000-4-11:2004 Voltage dips, short interruption and voltage variation on		
	power supply input lines Ed 2.0		
	CISPR 11:1997 (EN55011:1998) Limits and methods of measurement of radio		
	disturbance characteristics of industrial scientific and medical (ISM) radio-		
	frequency equipment RF Emissions Group 1, Class B		
Package	ISTA (Procedure 1A, 1994 Rev.)		
	Pre-Shipment Test Procedures (Package)		
Reliability	IEC60068-2-27:1987 Environmental testing – Shock		
	IEC60068-2-6:1995 Environmental testing –Vibration		
	IEC60068-2-64:1993 Environmental testing: vibration, broad-band random		
	(digital control) and guidance		
Labeling	EN1041:1998		
	Information supplied by the manufacturer with medical devices		
Marking	IEC /TR60878:2003		
	Graphical symbols for electrical equipment in medical practice		
	EN980:2003 Graphical symbols for use in the labeling of medical devices		
	ISO7000:2004 Graphical symbols for use on equipment-index and synopsis		
	EN60417-1:1999		
	Graphical symbols for use on equipment-overview and application		
	EN60417-2:1999 Graphical symbols for use on equipment-symbol originals		
	EN50419:2005 Marking of electrical and electronic equipment in accordance		

Manufacturer's Declaration

WARNING: For best product performance and measurement accuracy, use only accessories supplied or recommended by Colin Medical Technology. Use accessories according to the manufacturer's directions for use and your facility's standards. The use of accessories, transducers, and cables other than those specified may result in increased emission and/or decreased immunity of the BP-S510.

The BP-S510 is suitable for use in the specified electromagnetic environment. The customer and/or user of the BP-S510 should assure that it is used in an electromagnetic environment as described below;

Emission Test	Compliance	Electromagnetic Environment
RF emission CISPR 11	Group 1	The BP-S510 must emit electromagnetic energy in order to perform its intended function. Nearby electronic equipment may be affected.
RF emissions CISPR 11	Class B	The BP-S510 is suitable for use in all establishments.
Harmonic emissions IEC 61000-3-2	Class A	
Voltage fluctuations/flicker emission IEC 61000-3-3	Complies	

Table 42. Electromagnetic Emissions (IEC60601-1-2)

Table 43. Electromagnetic Immunit	ty ((IEC60601-1-2)	
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Immunity Test	IEC60601-1-2	Compliance	Electromagnetic
	Test Level	Level	Environment Guidance
Electrostatic discharge (ESD) IEC 61000-4-2	±6 kV contact ±8 kV air	±6 kV contact ±8 kV air	Floor should be wood, concrete, or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30 %.
Electric fast transient/burst IEC 61000-4-4	±2 kV for power supply lines ±1 kV for input/output lines	±2 kV for power supply lines ±1 kV for input/output lines	Mains power quality should be that of a typical commercial and/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 commercial and/or hospital environment
Voltage dips, short interruptions and voltage variations on power supply	<5 % U T (>95 % dip in UT) for 0.5 cycle	<5 % U T (>95 % dip in U T) for 0.5 cycle	Mains power quality should be that of a typical commercial and/or hospital environment. If the user of the BP-S510 requires continued operation during power mains interruption, it is recommended that the BP-S510 be powered
IEC 61000-4-11	40 % U T (60 % dip in UT) for 5 cycles 70 % U T (30 % dip in UT)	40 % U T (60 % dip in U T) for 5 cycles 70 % U T (30 % din in UT)	from an uninterruptible power supply or battery.
	for 25 cycles	for 25 cycles	

Immunity Test	IEC60601-1-2 Test Level	Compliance Level	Electromagnetic Environment Guidance
	<5 % U T (95 % dip in UT) for 5 sec.	<5 % U T (95 % dip in UT) for 5 sec.	
Power frequency (50/ 60 Hz) magnetic field IEC 61000-4-8	3 A/m	3 A/m	It may be necessary to position the BP-S510 further from the sources of power frequency magnetic fields or to install magnetic shielding. The power frequency magnetic field should be measured in the intended installation location to assure that it is sufficiently low.
Note: UT is the AC mains voltage prior to application of the test level.			

Table 44.	Electromagnetic	Immunity	(IEC60601-1-2	۱
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Immunity Test	IEC60601	Compliance	Electromagnetic environment
	test level	level	guidance
The BP-S510 is intended for use in the electromagnetic environment specified below. The customer or the user of the BP-S510 should assure that it is used in such an environment.			
			Potable and mobile RF communications equipment should be used no closer to any part of the BP-S510 including cables, than the recommended separation distance calculated from the equation appropriate to the frequency of the transmitter.
			Recommend separation distance
Conducted RF IEC 61000-4-6	3 Vrms 150 kHz to 80 MHz	3 Vrms	$d = 1.2 \sqrt{p}$
Radiated RF IEC 61000-4-3	3 V/m 80 MHz to 800 MHz	3 V/m	$d = 1.2 \sqrt{p}$ 80 MHz to 800 MHz
	3 V/m 800 MHz to 2 5	3 V/m	$d = 2.3 \sqrt{p}$ 800 MHz to 2.5 GHz
	GHz		where P is the maximum output power rating of the transmitter in watts (W) according to he transmitter manufacturer and d is the recommended separation distance in metres (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 he following symbol:
			((r;-))
Note: At 80 MHz Note: These guid affected by abso	and 800 MHz, the high delines may not apply in rption and reflection fro	ner frequency ra n all situations. om structures, o	ange applies. Electromagnetic propagation is bjects, and people.

Immunity Test	IEC60601	Compliance	Electromagnetic environment
	test level	level	quidance
^a Field strength telephones and I predicted theoret RF transmitters, strength in the lo level above, the performance is o relocating the BF	is from fixed transmitte and mobile radio, AM a ically with accuracy. To an electromagnetic sit ication in which the BP BP-S510 should be observed, additional m P-S510.	ers, such as bas and FM radio b assess the ele e survey should -S510 is used to baserved to heasures may l	se stations for radio (cellular/ cordless) roadcast, and TV broadcast cannot be ectromagnetic environment due to fixed d be considered. If the measured field exceeds the applicable RF compliance verify normal operation. If abnormal be necessary, such as re-orienting or
^b Over the frequ	iency range 150 kHz to	80MHz, field s	trengths should be less than 3 V/m

^b Over the frequency range 150 kHz to 80MHz, field strengths should be less than 3 V/m

Recommended Separation Distances between Portable and Mobile RF Communications Equipment and the BP-S510. (IEC60601-1-2)

Table 45. Recommended Separation Distances

Recommended separation distance between Portable and mobile RF communications equipment and the BP-S510

The BP-S510 is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of he BP-S510 can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the BP-S510 as recommended below, according to the maximum output power of the communications equipment.

Rated Maximum Output Power of	Separation distance according to frequency of transmitter in meter		
Transmitter in	150 kHz to MHz	80 MHz to 800 MHz	800 MHz to 2.5GHz
watt	$d = 1.2\sqrt{p}$	$d = 1.2\sqrt{p}$	$d = 2.3 \sqrt{p}$
0.01	0.12	0.12	0.23
0.1	0.38	0.38	0.73
1	1.2	1.2	2.3
10	3.8	3.8	7.3
100	12	12	23

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

Note: At 80MHz and 800MHz, the separation distance for the higher frequency range applies Note: These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects, and people.

Table 46. Cables (IEC60601-1-2)

Cables and Sensors	Maximum Length	Complies with	
Pulse oximetry extension cable	10 ft. (3 m)	-RF emissions, CISPR 11, Class B/ Group 1 -Harmonic emissions, IEC 61000-3-2 -Voltage fluctuations/flicker emission, IEC	
RS-232 serial communication	10 ft. (3 m)		
cable, 9 pin "D"			
Non-terminated cable, RS-232,	10 ft. (3 m)	61000-3-3	
9 pin "D"		-Electrostatic discharge (ESD), IEC 61000-4-	
Reusable SpO2 sensor	3 ft (0.91 m)	2	
AC power cord	10 ft. (3 m)	-Electric fast transient/burst, IEC61000-4-4	
	× /	-Surge, IE 61000-4-5	
		-Conducted RF IEC 61000-4-6	
		-Radiated RF, IEC 61000-4-3	