

SERVICE MANUAL

YM6000 Patient Monitor

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YM6000 Service Manual

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Introduction

Warnings



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



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



WARNING: Do not immerse the monitor or its accessories in liquid or clean with caustic or abrasive cleaners.



WARNING: Ensure that conductive portions of the electrodes, leads, and cable do not come into contact with any other conductive parts,



WARNING: Before attempting to open or disassemble the monitor, disconnect the power cord from the monitor.



WARNING: Chemicals from a broken LCD display panel are toxic when ingested. Use caution when handling a monitor with a broken display panel.



WARNING: The use of accessories, transducers, and cables other than those specified may result in increased emission and/or decreased immunity of the monitor.



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



WARNING: During the safety test, AC main voltage will be present on the applied part terminals, Exercise caution to avoid electrical shock hazard.



WARNING: Do not place the monitor into operation after repair or maintenance has been performed, until all Performance Tests and Safety Tests listed in Performance Verification section of this service manual have been performed. Failure to perform all tests could result in erroneous monitor readings.



WARNING: High voltage is generated by the LCD backlight driver. Exercise caution when operating monitor with covers open.

Cautions



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.

-  **CAUTION:** Observe ESD (electrostatic discharge) precautions when working within the unit and/or when disassembling and reassembling the monitor and when handling any of the components of the monitor.
-  **CAUTION:** When reassembling the monitor, over-tightening could strip out the screw holes in the cases, rendering it unusable.
-  **CAUTION:** If any problem with the monitor built in an optional printer, check a printer's door is closed well. Operating error may be caused if the cover is not closed correctly.
-  **CAUTION:** If internal battery cable has been disconnected, pay particular attention to polarity of the cable before reattaching. If battery cable polarity is reversed, it is likely that circuit damage will occur.
-  **CAUTION:** For continued protection against risk of fire, replace only with same type and rating of fuse.
-  **CAUTION:** Ferrite Cores are used for electromagnetic compatibility. Please do not remove Ferrite Cores while disassembling or reassembling, otherwise the monitor can be affected by electromagnetic interference and measure inaccurate data to be displayed or stored.

Manual Overview

This manual contains information for servicing the YM6000 monitor. The monitor subsequently referred to as the monitor throughout this manual. Only qualified service personnel should service this product. Before servicing the monitor, read the operator's manual carefully for a thorough understanding of safe operation.

Read and understand all safety warnings and service notes printed in this service manual.
and the operator's manual part number A7076.

Related Documents

To perform test and troubleshooting procedures and to understand the principles of operation and circuit analysis sections of this manual, you must know how to operate the monitor. Refer to the monitor operator's manual part number A7076.

To understand the various SpO₂ sensors, ECG leads, blood pressure cuffs, and temperature probes that work with the monitor, refer to the individual directions for use that accompany these accessories.

Intended Use for the YM6000 Monitor

The YM6000 monitor is intended to be used to monitor electrocardiography, heart rate, pulse rate, noninvasive blood pressure (systolic, diastolic and mean arterial pressures), functional arterial oxygen saturation, invasive blood pressure, respiration, capnography (EtCO₂ and InCO₂) and temperature for adult, pediatric and neonate 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, surgicenters, and sub-acute centers.

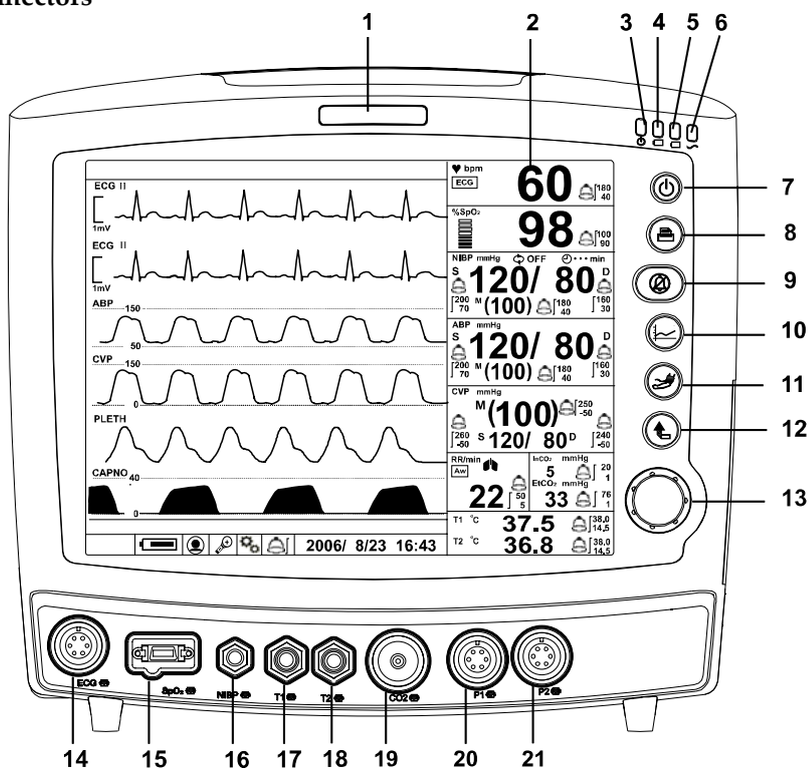
Identifying the YM6000 monitor Configurations

The following table identifies YM6000 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.

Model-option	Description
YM6000-0	Basic (ECG, NIBP, SpO ₂ , 2-channel Temperature, Respiration)
YM6000-P0	Basic + Recorder
YM6000-0B	Basic + IBP
YM6000-0E	Basic + Capnography
YM6000-P0B	Basic + Recorder +IBP
YM6000-P0E	Basic + Recorder + Capnography
YM6000-0BE	Basic +IBP + Capnography
YM6000-P0BE	Basic + Recorder + IBP + Capnography

Monitor Controls

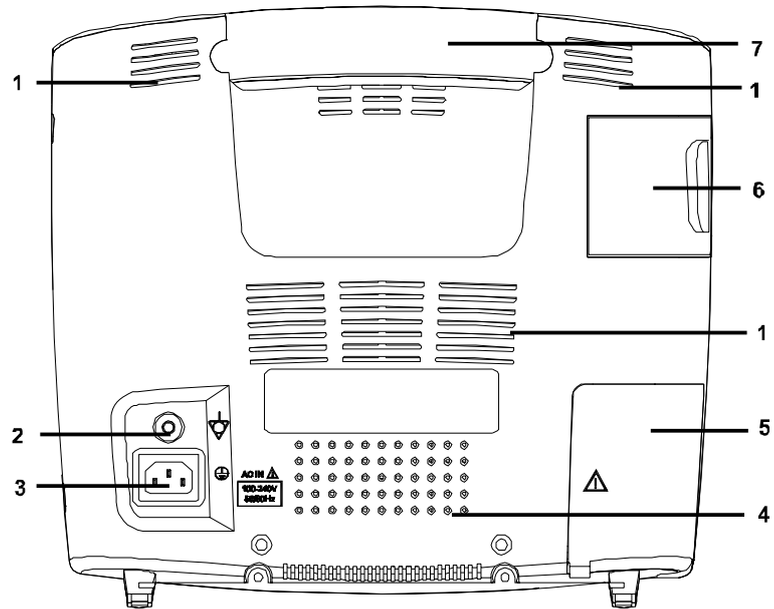
Front Panel Controls and Connectors



- 1 Visual alarm indicator
- 2 LCD (Liquid crystal display)
- 3 Power on indicator
- 4 Battery charging status indicator A
- 5 Battery charging status indicator B
- 6 AC indicator
- 7 Power button
- 8 Print button
- 9 Alarm silence/suspension button
- 10 Trend button
- 11 NIBP start/stop button
- 12 Return button
- 13 Trim knob
- 14 ECG connector
- 15 SpO₂ connector
- 16 NIBP connector
- 17 Temperature channel 1
- 18 Temperature channel 2
- 19 CO₂ connector (option)
- 20 IBP 1 connector (option)
- 21 IBP 2 connector (option)

Figure 1. Front Panel Controls and Connectors

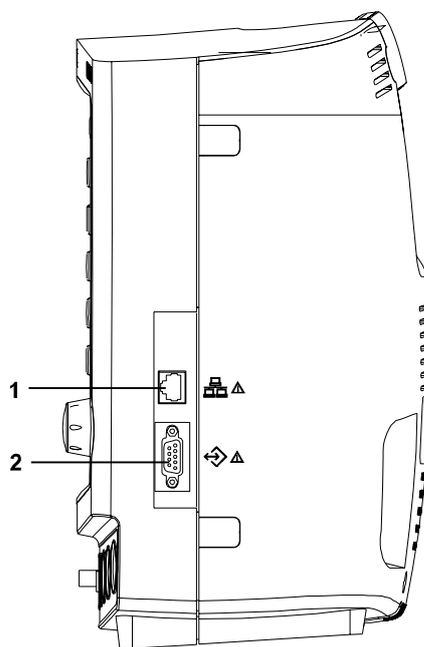
Rear Panel Components



- | | | | |
|---|------------------------|---|--------------------------|
| 1 | Air ventilation | 5 | Battery cover |
| 2 | Equipotential terminal | 6 | Thermal Printer (option) |
| 3 | Ac power connector | 7 | Handle |
| 4 | Speaker | | |

Figure 2. Rear Panel Components and Symbols

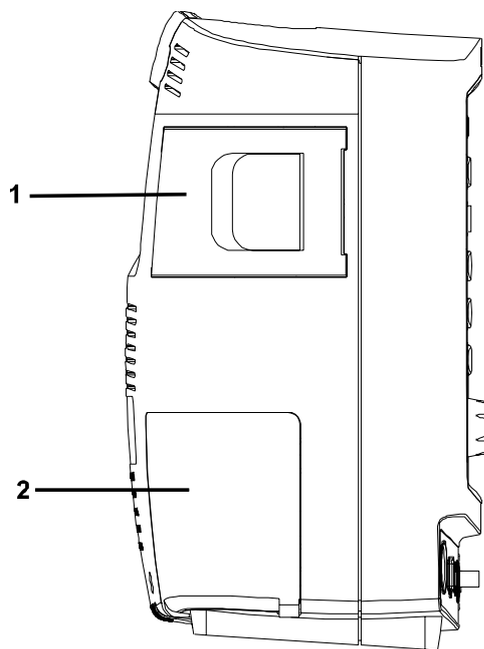
Right Panel Components



- 1 LAN connector
- 2 RS-232 interface connector

Figure 3. Right Panel Components and Symbols

Left Panel Components



- 1 Thermal printer (option)
- 2 Battery cover

Figure 4. Left Panel Components and Symbols

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Routine Maintenance



WARNING: Do not spray or pour any liquid on the monitor or its accessories. Do not immerse the monitor or its accessories in liquid or clean with caustic or abrasive cleaners.

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.

- Quaternary Ammonium
- Alcohol-70% Isopropyl
- 10% Chlorine bleach solution
- PDI Sani-System

For cables, sensors, cuffs, and probes, follow 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.

Periodic Safety and Functional Checks

The monitor requires no routine service or calibration other than cleaning and battery maintenance. The following performance verification tests may be used following repair or during routine maintenance (if required by your local institution). The following checks should be performed at least every 2 years by qualified service personnel.

1. Inspect the exterior of the monitor for damage.
2. Inspect labels for legibility. If the labels are not legible, contact Mediana Technical Service Department.
3. If the monitor has been visibly damaged or subjected to mechanical shock (for example, if dropped), perform the performance tests as described in **Performance Verification** section. If the unit fails these performance tests, refer to Troubleshooting section.
4. Perform the electrical safety tests detailed in **Performance Verification** section. If the unit fails these electrical safety tests, do not attempt to repair. Contact Mediana Technical Service Department.
5. Inspect the fuses for proper value and rating.
qty 2, 6.3 A, 250 volts for AC mains

Batteries

If the monitor has not been used for a long period of time, more than 2 months, the battery will need charging. To charge the battery, connect the monitor to an AC outlet as described in **Battery Charge** paragraph in this service manual or the **Battery Operation** section of the operator's manual.

Note: Storing the monitor for a long period without charging the battery may degrade the battery capacity. The battery may require a full charge/discharge cycle to restore normal capacity. Mediana recommends that the monitor's sealed, Ni-MH batteries be replaced at 2 year intervals. Refer to **Disassembly Guide** section.



CAUTION: If the monitor is to be stored for a period of 2 months or longer, it is recommended to notify service personnel to remove the battery from the monitor prior to storage. To replace or remove the battery, refer to Disassembly Guide section. Recharging the battery is strongly recommended when the battery has not been recharged for 2 or more months.



CAUTION: If the battery shows any signs of damage, leakage, or cracking, it must be replaced immediately.



CAUTION: Discarded battery may explode during incineration. Recycle used batteries properly. Do not dispose of batteries in refuse containers.

Environmental Protection

Follow local governing ordinances and recycling plans regarding disposal or recycling batteries and other device components.

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.

Performance Verification

General

This section discusses the tests used to verify performance following repairs or during routine maintenance. All tests can be performed without removing the monitor covers. All tests except the battery charge and battery discharge tests must be performed as the last operation before the monitor is returned to the user.

If the monitor fails to perform as specified in any test, repairs must be made to correct the problem before the monitor is returned to the user.

Required Equipment

Table 1 lists the equipment required for performance verifications.

Table 1. Required Equipment

Equipment	Description
Digital multimeter (DMM)	Fluke Model 87 or equivalent
ECG cable for 3 leads	Mediana ECG cable for 3 lead
ECG cable for 5 leads (option)	Mediana ECG cable for 5 lead
ECG leads	Mediana ECG 3 lead
ECG leads (option)	Mediana ECG 5 lead
NIBP hose	Mediana cuff hose
NIBP cuff	Mediana cuff, Durable
NIBP rigid PVC vessel	9cm diameter
SpO ₂ extension cable	Nellcor DOC-10
SpO ₂ sensor (durable)	Nellcor DS-100A
Temperature probes	YSI-400 series
SpO ₂ simulator	Nellcor SRC-MAX simulator
ECG simulator	METRON PS-420 or equivalent
NIBP simulator	Bio-Tek BP Pump 2 or equivalent
Respiration simulator	METRON PS-420 or equivalent
Temperature simulator	Medsim 300 or equivalent
IBP simulator	METRON PS-420 or equivalent
IBP test cable	Mediana IBP test cable
Calibration gas cylinder	8-12% CO ₂ , ±0.02% balance air or nitrogen
Gas valve and non-silicon tubing	Gas valve and non-silicon tubing
Sample elbow or tee	Sample elbow or tee
CO ₂ gas flow meter	BCI catalogue No. 8133 recommended
Sample line	Sample line
Water trap	Water trap
CO ₂ scrubber (or 0% CO ₂ medical-grade gas source)	CO ₂ scrubber (or 0% CO ₂ medical-grade gas source)
Safety analyzer	METRON QA-90 or equivalent
Data interface cable	9-pin RS-232 cable
PC	Common PC
LAN cable	Common used LAN cable
Stopwatch	Manual or electronic

Note: Contact Mediana Technical Service Department for pricing and ordering information.

System Tests

The monitor must be placed in the service menu. For a detailed explanation to access the service menu, refer to **Service Menu and Factory Default Settings** section.

1. Rotate the trim knob to select *System Test* in the service menu, and then press the trim knob.

LCD Test

This tests the LCD display.

1. Select *LCD Test* in the system test.
2. The screen color will change over the following sequence:
Red → Green → Blue → White → Black ... every two seconds
3. After testing the test, press trim knob twice to exit.

Pass/Fail Results

When the color of the test screen changes in the order from Red, Green, Blue, White to Black, the LCD display is in a normal state.

Alarm Audible Test

This tests the alarm tones by displaying the level of the alarm tone on the screen.

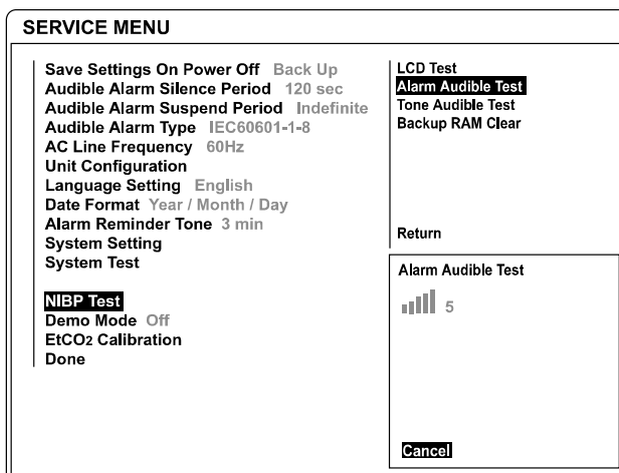


Figure 5. Alarm Audible Test

1. Select *Alarm Audible Test* in the system test.
2. The level of the alarm tone will appear on the screen as the alarm tone sounds. Then, the level goes up gradually. When the tone reaches the maximum level 8, it returns to the minimum level 1.
3. After finishing the test, press the trim knob to select *Cancel*. The menu box will disappear.

Pass/Fail Results

When the alarm tone changes to eight steps, the alarm tone is in a normal state.

Tone Audible Test

This tests HR/PR tones, key beeps and completion sounds.

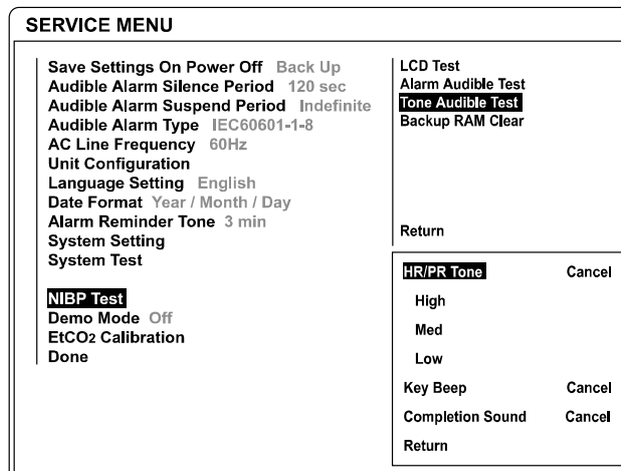


Figure 6. Tone Audible Test

1. Select *Tone Audible Test* in the system test.
2. Rotate the trim knob to select *HR/PR Tone*.
The HR/PR tone sounds intermittently as the level goes up gradually. When the tone reaches the maximum level 7, it returns to the minimum 1.
When the tone returns to the minimum, the pitch will change automatically.
There are three pitches – High, Med and Low.
3. Press *Cancel* to finish the test.
4. Rotate the trim knob to select *Key Beep*.
The key beep sounds intermittently as the level goes up gradually. When the tone reaches the maximum level 7, it returns to the minimum 1.
5. After finishing the test, press the trim knob to select *Cancel*. The menu box will disappear.

Pass/Fail Results

When the HR/PR tone changes to seven steps and the pitch changes to three steps, the HR/PR tone is normally set.

When the key beep changes to seven steps, the key beep is normally set.

When the completion sound is heard, it is normally set.

Backup RAM Clear

When *Backup RAM Clear* is set to *Yes*, all settings of the monitor including the service settings return to the factory defaults from the next cycle.

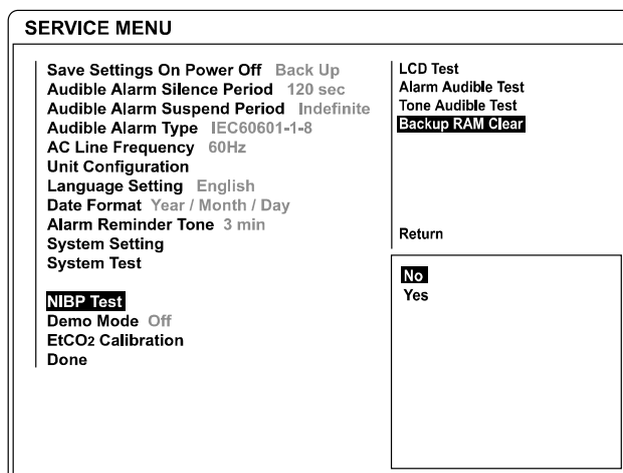


Figure 7. Backup RAM Clear

Note: Set the monitor to appropriate AC line frequency (50Hz or 60Hz) again after Back Up RAM clearance.

Performance Tests

The battery charge and battery discharge test should be performed before monitor repairs whenever the battery is suspected as being a source of the problems. All other tests may be used following repairs or during routine maintenance (if required by your local institution). Before performing the battery discharge test, ensure that the battery is fully charged. This section is written using **factory defaults set** as power-up. If your institution has preconfigured custom defaults, those values will be displayed.

Power

1. Connect the monitor to AC power source using proper power cord.
2. Verify that *AC indicator* is lit.
3. Press *Power button* over 1 second to turn on the monitor.
4. Verify that the monitor is turned on and that *Power on indicator* is lit
5. After the monitor operates in normal mode, disconnect the power cord.
6. Verify that *Battery status icon* appears on the screen instead of lighting AC indicator.
7. Press *Power button* over 1 second, and then verify that the monitor is turned off.

Battery Charge

1. Connect the monitor to AC power source using a proper power cord.
2. Verify *AC indicator* is lit.
3. Charge the battery fully until *Battery charging status indicator A and B* is no more flashing. It takes for at least 12 hours.
4. To check for a full charge, perform the procedure in paragraph "**Battery Discharge**".

Note: The battery may require a complete charge/discharge cycle to restore its normal capacity, depending on its previous usage.

Battery Discharge

1. Disconnect the power cord from the monitor with fully charge battery.
2. Turn on the monitor by pressing *Power button* over 1 second.
3. Verify *Battery status icon* appears at bottom of the screen after power-on self-test is completed. The bar in battery status icon should be filled, indicating battery is charged.
4. Connect SpO₂ simulator to monitor via the SpO₂ extension cable.
5. Connect NIBP simulator to the monitor via Mediana hose.
6. Set SpO₂ simulator as follows: SpO₂ of 90% and pulse rate of 60bpm.
7. Set NIBP simulator to simulate pressure setting of 120/80 mmHg and heart rate of 80bpm.
8. Set NIBP Automatic mode interval of the monitor to 15 minutes.
9. The monitor must operate for 1 hour with one fully charged battery. The monitor must operate for at least 15 minutes before the monitor powers down due to low battery condition.
10. Verify that low priority audible alarm occurs and a warning message "*low battery*" is displayed about 15 minutes before battery fully discharges.

11. Allow monitor to operate until it automatically powers down due to low battery condition. Verify that high priority audible alarm occurs and a warning message "*Critically low battery*" is displayed about 5 minutes before monitor automatically shuts down.
12. If monitor passes this test, immediately recharge battery. (paragraph "**Battery Charge**")

Power-On Self-Test

1. Connect the monitor to AC power source and verify that *AC indicator* is lit.
2. Observe the monitor's front panel. With monitor off, press *Power button*. The monitor must perform the following sequence.
 - a. The monitor progresses the check sum for the flash memory and displays the bar while the check sum is proceeding
 - b. When the check sum completed, the bar is filled, and then Mediana logo appears for a few seconds, with version numbers of the software and the copyright in lower left corner of display.
 - c. The monitor emits the high-pitched beep.
 - d. *Visual alarm Indicator* (red) on the top of the front panel, *AC indicator* (green) and *battery charging status indicators A and B* (green) on the right top of the front panel are illuminated.
 - e. Upon successful completion of power-on self-test, display will be in normal monitoring screen configuration.

Note: Power-on self-test takes approximately 3 seconds to complete. No vital signs numeric values or waveforms will be displayed.

Note: During the monitor's power-on-self-test, the integrity of all programming is checked first. If software testing is successful, hardware tests are initiated. If all testing is successful, the monitor is ready for use. If an error code is displayed, please refer to **Troubleshooting** section.

General Operation Tests

Alarms and Alarm Silence

1. Connect the monitor to an AC power source.
2. Press **Power button** to turn on the monitor.
3. Connect SpO₂ simulator to sensor input cable and connect cable to monitor.
4. Set SpO₂ simulator as follows: SpO₂ of 75% and pulse rate of 60bpm.
5. Verify following the monitor reaction:
 - a. Pulse bar begins to track artificial pulse signal from SpO₂ simulator.
 - b. After about 10 to 20 seconds, the monitor displays saturation and pulse rate as specified by simulator. Verify values are within following tolerances:
 - Tolerance of Oxygen Saturation : ± 2 %
 - Tolerance of Pulse Rate : ± 3 bpm
 - c. Audible alarm sounds and "*SpO₂: Lower limit violated.*" message will be displayed and % SpO₂ numerical area will flash, indicating the parameter has violated default alarm limits.
6. Press **Alarm silence/suspension button** on the monitor front panel. Audible alarm is temporarily silenced.
7. Verify the following:
 - a. An audible alarm remains silenced.
 - b. **Alarm silence icon** appears in %SpO₂ numerical area on display.
 - c. %SpO₂ display continues flashing.
 - d. Audible alarm returns in approximately 120 seconds.

HR/PR Tone Volume Control

1. Press **Power button** to turn on the monitor.
2. Connect SpO₂ simulator to sensor input cable and connect cable to monitor.
3. Set SpO₂ simulator as follows: SpO₂ of 75% and pulse rate of 60bpm.
4. Verify SpO₂ and pulse rate values are correctly displayed.
5. Press **Alarm silence/suspension button** on front panel of the monitor to temporarily silence audible alarm.
6. Verify **HR/PR source** on HR/PR numerical area is set to "*SpO₂*".
7. Select **Setup icon** on the screen to display **Setup menu**.
8. Rotate the trim knob to highlight HR/PR tone volume on **Setup menu** and press the trim knob to adjust HR/PR tone volume.
9. Set HR/PR tone volume level 1 to level 7 and return to the monitoring screen. Verify beeping pulse rate tone increases.
10. Set HR/PR tone volume level 7 to level 1 and return to the monitoring screen. Verify beeping pulse rate tone decreases.

11. Set HR/PR tone volume to “Off” and return to the monitoring screen. Verify beeping pulse rate tone is no longer audible.
12. Return HR/PR tone volume to a comfortable level.

Sensor LED Excitation Test

This procedure uses normal system components to test circuit operation. A *DS-100A*, used to examine LED intensity control. The red LED is used to verify intensity modulation caused by the LED intensity control circuit.

1. Connect the monitor to an AC power source.
2. Press the *Power button* to turn on the monitor.
3. Connect the SpO₂ extension cable to the monitor.
4. Connect SpO₂ sensor to the SpO₂ extension cable.
5. Leave the sensor open with the LEDs and photo detector visible.
6. After monitor completes its normal power-up sequence, verify that the sensor LED is brightly lit.
7. Slowly move sensor LED in proximity of photodetector element of the sensor (close the sensor slowly). Verify; as LED approaches the optical sensor, that the LED intensity decreases.
8. Open the sensor and notice that the LED intensity increases.
9. Repeat step 7 and intensity will again decrease. This variation is an indication that the microprocessor is in proper control of LED intensity.
10. Press *Power button* to turn off the monitor.

Restoring Power-On Default Settings

The following test procedures will verify that alarms are activated at the level of factory default alarm limits and that any changed settings are saved and in effect when the user changes alarm limit settings and saves the current settings as a power default.

Table 2. Parameter Alarm Limit Factory Defaults

Factory Defaults	Adult	Pediatric	Neonatal
HR/PR High Alarm Limits	120 BPM	160 BPM	200 BPM
HR/PR Low Alarm Limits	50 BPM	70 BPM	100 BPM
NIBP SYS High Alarm Limits	160 mmHg	120 mmHg	90 mmHg
NIBP SYS Low Alarm Limits	90 mmHg	70 mmHg	40 mmHg
NIBP DIA High Alarm Limits	90 mmHg	70 mmHg	60 mmHg
NIBP DIA Low Alarm Limits	50 mmHg	40 mmHg	20 mmHg
NIBP MAP High Alarm Limits	110 mmHg	90 mmHg	70 mmHg
NIBP MAP Low Alarm Limits	60 mmHg	50 mmHg	30 mmHg
% SpO ₂ High Alarm Limits	100 %	100 %	95 %
% SpO ₂ Low Alarm Limits	90 %	90 %	80 %
IBP1, 2 SYS High Alarm Limits	160 mmHg	120 mmHg	90 mmHg
IBP1, 2 SYS Low Alarm Limits	90 mmHg	70 mmHg	40 mmHg
IBP1, 2 DIA High Alarm Limits	90 mmHg	70 mmHg	60 mmHg
IBP1, 2 DIA Low Alarm Limits	50 mmHg	40 mmHg	20 mmHg
IBP1, 2 MEAN High Alarm Limits	110 mmHg	90 mmHg	70 mmHg
IBP1, 2 MEAN Low Alarm Limits	60 mmHg	50 mmHg	30 mmHg
RESP High Alarm Limits	30 BPM	30 BPM	100 BPM

Factory Defaults	Adult	Pediatric	Neonatal
RESP Low Alarm Limits	8 BPM	8 BPM	30 BPM
EtCO ₂ High Alarm Limits	80 mmHg	80 mmHg	80 mmHg
EtCO ₂ Low Alarm Limits	0 mmHg	0 mmHg	0 mmHg
InCO ₂ High Alarm Limits	20 mmHg	20 mmHg	20 mmHg
InCO ₂ Low Alarm Limits	0 mmHg	0 mmHg	0 mmHg
TEMP1, 2 High Alarm Limits	39.0°C (102.2°F)	39.0°C (102.2°F)	39.0°C (102.2°F)
TEMP1, 2 Low Alarm Limits	36.0°C (96.8°F)	36.0°C (96.8°F)	36.0°C (96.8°F)

1. Turn on the monitor at the factory default settings.
2. Select *Alarm limits icon* to display *Alarm limits menu*.
3. Verify that alarm limits are set to as shown in Table 2.
4. Change *Patient mode* from Adult to Pediatric or Neonatal, then verify that alarm limits are set to as shown in Table 2.
5. Change alarm limit value via *Alarm limits menu*.
6. Save the current changed alarm limit values as a power on default setting via the service menu (see **Service Menu and Factory Defaults** section). Turn off the monitor.
7. Press *Power button* to turn on the monitor.
8. Verify alarm limits are set to the current changed alarm limit values.

Printer Test (Option)

If Printer option is installed in the monitor, the following test procedures will verify the printer performance.

1. Connect the monitor to an AC power source.
2. Press *Power button* to turn on the monitor.
3. Connect all necessary simulators to the monitor.
4. Select *Setup icon* on the screen to display *Setup menu*.
5. Test #1: 20 sec printing
 - a. Set Wave record time to *20 sec* via *Setup menu*.
 - b. Press *Print button* when all the parameter signals display normally.
 - c. Verify that the parameter values and waveforms are printed out for 20 seconds.
6. Test #2: Continuous printing
 - a. Set Wave record time to *Continuous* via *Setup menu*.
 - b. Press *Print button* when all the parameter signals display normally.
 - c. Verify the parameter values and waveforms are printed out continuously.
 - d. Verify printing stops with pressing *Print button* again.

7. Test #3: Record speed

- a. Set *Record speed* to 25 mm/s.
- b. Press *Print button* when all the parameter signals display normally.
- c. Verify the parameter values and waveforms are printed out with 25 mm/s.
- d. Set *Record speed* to 50 mm/s.
- e. Press *Print button* when all the parameter signals display normally.
- f. Verify that the parameter values and waveforms are printed out with 50 mm/s.

8. Test #4: Wave Record Select

- a. Set Wave Record Select to *ECG1+ECG2* via *Setup menu*.
- b. Press *Print Button* when all the parameter signals display normally.
- c. Verify the two ECG waveforms are printed out.
- d. Repeat this test for other selections.

9. Test #5: Record on Alarm

- a. Set *Record on Alarm* to *On* via *Setup menu*.
- b. Set Heart rate of ECG simulator to 30 bpm.
- c. Verify "HR Lower limit violated" alarm is activated and the parameter values and waveforms are printed out.

10. Test #6: Auto List Record

- a. Set *Auto List Record* to *On* via *Setup menu*.
- b. Set *Save Time Interval* to *0.5 minutes* via Tabular trend menu.
- c. Verify that the monitor automatically prints out the data after stored 8 data in the trend memory.

Note: If there is no printer paper left or printer paper places improperly, the monitor will display an informative message.

Network Test

Perform the following procedure to test the Network. The Network connector is located on the monitor's right panel, identified with the Network symbol.



CAUTION: Do not change any other settings of the test programs while performing the Network test.

Note: Network Test is only available when YM6000 monitor is equipped with TCP/IP module for Central Monitoring System (CMS). Otherwise this test does not need to perform.

Note: Contact Mediana Technical Service Department for the software required.

1. Connect the network line to the monitor, then turn on the monitor.
2. Run "YM6000_Comm_Test.exe" on PC connected the network line using the same gateway as the monitor.

3. Press **PROBE button** on PC.
4. Verify that the number of the monitor connections to PC found is correct
5. Press **EXIT button** on PC to close "YM6000_Comm_Test.exe".

Nurse Call Test

Perform the following procedure to test the Nurse Call. The nurse call connector is Dsub-9, located on the monitor's right panel, identified with the data interface symbol (RS-232).

1. Connect the negative lead of a voltmeter to pin 5 and positive lead to pin 1 of the data port connector on the right panel of the monitor. Ensure that the audible alarm is not silence or turned off.
2. Connect the SRC-MAX simulator to the DOC-10 sensor cable.
3. Connect temperature probe to temperature input port on YM6000.
4. Turn on the monitor and wait for the monitor to complete POST.

Note: The monitor should indicate a %SpO₂ alarm of 75.

5. Verify an output voltage at pins 5 and 1 between +5 to +12 DC.
6. Press Alarm silence/suspension button. With no active audible alarm, the output voltage at pins 5 and 1 must be between -5 to -12 VDC. This verifies the RS-232 Nurse Call function.
7. With the instrument in an alarm condition, use a digital multimeter (DMM) to verify that there is no continuity (1 mega ohms or greater) between pins 8 and 9 and that there is continuity (60 ohms or less) between pins 7 and 8.
8. Press the SRC-MAX simulator %SpO₂ button to change the %SpO₂ to 90.
9. Use a DMM to verify that there is continuity between pins 8 and 9 and that there is no continuity between pins 7 and 8. This verifies the solid state Nurse Call function.

Note: The pin layouts and signal descriptions are included in the operator's manual. For the detailed information regarding the Nurse Call, refer to **Nurse Call Interface** section of the operator's manual.

Measurement Parameter Operation Tests

ECG Operation

1. Press **Power button** to turn on the monitor.
2. Connect 3 ECG leads to appropriate jacks on ECG simulator.
3. Connect leads to Mediana ECG cable.
4. Connect Mediana ECG cable to ECG connector on the monitor's front panel.
5. Set ECG simulator as follows:
 - Heart rate: 30 bpm
 - Amplitude: 1 millivolt
 - Lead select: II
 - Normal sinus rhythm
 - Adult mode
6. After normal power-up sequence, verify that the following monitor reactions:
 - a. After about 15 seconds, the monitor displays a heart rate of 30 \pm 3 bpm.
 - b. Verify that audible alarm sounds and that "**HR: Lower limit violated**" message displays on the message area.
 - c. Verify that the HR/PR numerical area flashes and that the heart rate is below default low alarm limit (medium priority alarm).
7. Increase the heart rate setting on ECG simulator to 240 bpm.
 - a. After about 15 seconds, verify the monitor displays heart rate of 240 \pm 3 bpm.
 - b. Verify that audible alarm sounds and that "**HR: Upper limit violated**" message displays
 - c. Verify that the HR/PR numerical area flashes and that the heart rate is above default high alarm limit (medium priority alarm).
8. Decrease the heart rate setting on ECG simulator to 120 bpm.
 - a. After about 15 seconds, verify the monitor displays heart rate of 120 \pm 3 bpm.
9. Disconnect the LL lead from ECG simulator.
 - a. Verify that the "**ECG: Check ECG leads & electrodes**" message appears, that three dashes are displayed in the HR/PR numerical area, and that low priority audible alarm sounds.
 - b. Reconnect the LL lead to ECG simulator. Verify that "**ECG: Check ECG leads & electrodes**" message no longer appears and that the audible alarm is stopped.
 - c. Repeat this test for the LA and the RA leads.
10. Connect all the leads to the monitor.
 - a. Select **ECG waveform menu** and set **Lead select** to Lead I.
 - b. Verify that the lead selection is displayed on the ECG waveform area.
 - c. Repeat step 10-a for all the ECG Lead selections.
11. Set ECG Lead selection to Lead II.
12. Change ECG waveform size to all the selectable sizes and verify that an appropriate size is displayed on the ECG waveform area.

13. Set *ECG Cable Select* to *Auto* via ECG waveform menu.
14. Verify that Lead Select in ECG waveform menu displays I, II, III.
15. Disconnect 3 ECG leads and connect 5 ECG leads.
16. Verify that Lead Select in ECG waveform menu displays I, II, III, aVR, aVL, aVF, V (Chest Lead).
17. Repeat step 9 to 12.
18. Turn off the monitor.

Note: The accuracy of the monitor ECG measurements is ± 3 bpm. In the procedure, add the tolerance of the simulator to the acceptable range of readings.

NIBP Operation



WARNING: A blood pressure cuff, connected to the monitor, should never be applied to a human subject while the monitor is in the pressure test mode, as injury could result.

These tests verify the functionality of the monitor's pneumatic system. The monitor must be placed in *Service menu*. For a detailed explanation to access the service menu, refer to **Service Menu and Factory Default Settings** section.

1. Rotate the trim knob to select NIBP Test in the service menu, and then press the trim knob.

Note: Before accessing the NIBP Test mode, ensure that current patient mode is proper for the pneumatic system to be test. You can set Patient mode: Adult, Pediatric or Neonatal via the setup menu.

Note: In the NIBP Test mode, no function button will have no effect except the trim knob. All the tests will start to be performed by pressing or rotating the trim knob. If you would like to stop the test during test progressing, press the trim knob.

Note: Inflation Speed Test and Deflation Speed Test are intended for factory use only.

Pressure Sensor Accuracy Test

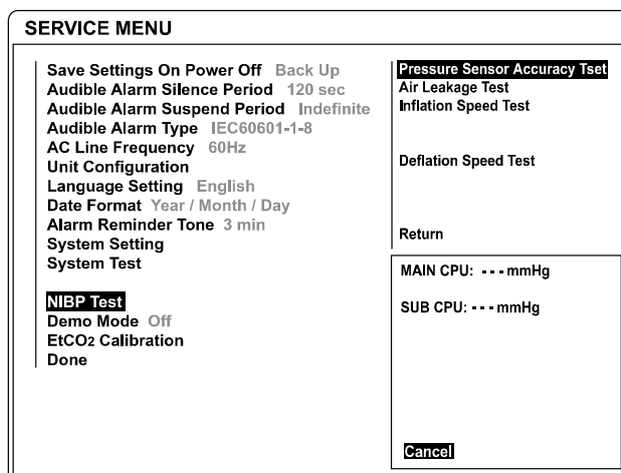


Figure 8. Pressure Sensor Accuracy Test

1. Connect the NIBP cuff hose to the NIBP connector on the monitor’s front panel.
2. Connect the other end of the NIBP cuff hose to the NIBP simulator.
3. Rotate the trim knob to select *NIBP Test* in the service menu, and then press the trim knob.
4. Rotate the trim knob to select *Pressure Sensor Accuracy Test*, and then press the trim knob.
5. Press Select button on the simulator until simulator displays “Pressure Source Set Test Pressure”. Adjust pressure on the simulator for 250, 150, 50 and 0 mmHg.
6. Press *Start Pump button* on simulator. The simulator will begin to pressurize. Allow 15~20 seconds for pressure to stabilize.
7. Compare the pressure of the simulator with the pressure displayed by the monitor.
If air leakage disables accurate comparison of the above pressures, eliminate the cause of air leakage.
8. After finishing the test, press the trim knob to select *Cancel*. The menu box will disappear. If *Cancel* is selected during the test progressing, the test will stop and the menu box will disappear.

Pass/Fail Results

The difference between the simulator’s and the monitor’s readings are as follows:

Sphygmomanometer	Monitor’s Readings
0 mmHg	±0 mmHg
50 mmHg	50±6 mmHg
100 mmHg	100±6 mmHg
200 mmHg	200±6 mmHg

Air Leakage Test

The air leakage test verifies the integrity of the pneumatic system.

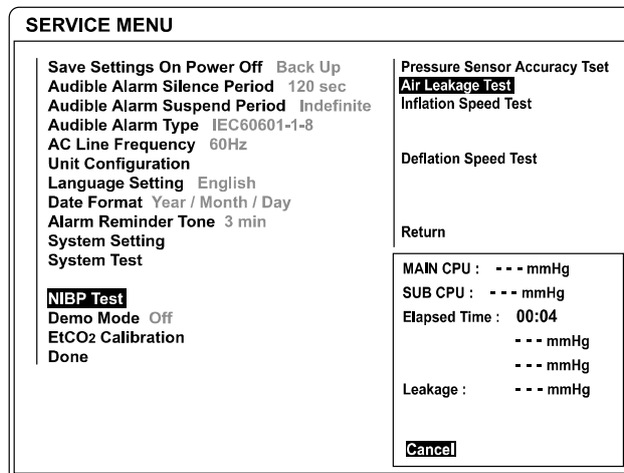


Figure 9. Air Leakage Test

1. Place the adult cuff with a rigid PVC vessel (5 cm diameter).
2. Connect the cuff to the NIBP connector on the monitor's front panel via NIBP cuff hose.
3. Rotate the trim knob to select *NIBP Test* in the service menu, and then press the trim knob.
4. Rotate the trim knob to select *Air Leakage Test*, and then press the trim knob.
5. The result will be displayed on the monitor's screen in four minutes.
6. After finishing the test, press the trim knob to select *Cancel*. The menu box will disappear. If *Cancel* is selected during the test progressing, the test will stop and the menu box will disappear.

Pass/Fail Results

It passes if the leakage value is less than 12mmHg/3minutes.

Pulse Oximetry Operation

1. Connect the monitor to an AC power source.
2. Turn on the monitor by pressing the *Power button*.
3. Connect the SpO₂ extension cable to SpO₂ connector on the monitor's front panel after the monitor completes POST.
4. Connect the SpO₂ simulator to the other end of the SpO₂ extension cable.
5. The monitor will:
 - be in SpO₂ alarm
 - display an SpO₂ of 75 (Test pass criteria is 73 to 77 % SpO₂)
 - display a pulse rate of 60 (Test pass criteria is 57 to 63 bpm)
 - display low level modulation
6. Test #1: SpO₂
 - a. Press % *SpO₂ selection button* on the SpO₂ simulator. The % SpO₂ 90 LED on the SpO₂ simulator will light.
 - b. The monitor will display three dashes until the SpO₂ simulator stabilizes at 90 % SpO₂. The test pass criteria are 88 to 92 % SpO₂.
 - c. The monitor will display:
 - 90 % SpO₂
 - 60 bpm
 - no alarm
7. Test #2: Pulse rate (bpm)
 - a. Press *PULSE RATE selection button* on the SpO₂ simulator. The PULSE RATE 200 LED on the SpO₂ simulator will light:
 - b. The monitor will increase to 200 bpm. The test pass criteria is 197 to 203 BPM.
 - c. The monitor will display:
 - 90 % SpO₂
 - 200 bpm
 - alarm: "*PR(SpO₂): Upper limit violated*" message will display and the HR/PR area will flash, indicating pulse rate is above default high alarm limit (medium priority alarm).
 - d. Press *PULSE RATE selection button* on the SpO₂ simulator. The PULSE RATE 60 LED will light.
 - e. The monitor will decrease to 60 and stabilize at 60bpm. The test pass criteria is 57 to 63 bpm.
 - f. The monitor will display:
 - 90 % SpO₂
 - 60 bpm
 - no alarm
 - low level modulation
8. Test #3: Modulation Level
 - a. Press %*MODULATION selection button* on the SpO₂ simulator. The %MODULATION LED on the SpO₂ simulator will light.
 - b. The monitor's waveform area will spike and stabilizes at a higher modulation level.
 - c. The monitor will display:
 - 90 % SpO₂

- 60 bpm
 - no alarm
- d. Disconnect all equipments and turn off the monitor.

Respiration Operation

1. Press **Power button** to turn on the monitor.
2. Connect ECG leads to an appropriate jack on Respiration simulator.
3. Connect ECG leads to Mediana ECG cable.
4. Connect Mediana ECG cable to ECG connector on the monitor's front panel.
5. Set ECG simulator for the respiration rate of 120 breaths per minute.
6. After the normal power-up sequence, verify the following reactions:
 - a. The monitor displays respiration rate of 120 ± 3 breaths per minute.
 - b. Audible alarm will sound, "**Resp: Upper limit violated**" message will display and the Respiration numerical area will flash, indicating a respiration rate is above default high alarm limits. (medium priority alarm)
7. Decrease the respiration rate setting on Respiration simulator to 20 breaths per minute.
 - a. Verify that the monitor displays the respiration rate of 20 ± 3 breaths per minute.

Note: The accuracy of Respiration rate measurements is ± 3 breaths per minute. In the procedure below, add the tolerance of the simulator to the acceptable range of readings.

Temperature Operation

1. Press **Power button** to turn on the monitor.
2. Connect the temperature probe (supplied with the temperature simulator) to an appropriate jack on temperature simulator.
3. Connect the temperature probe to the temperature connector on the monitor's front panel.
4. Set Temperature simulator as follows:
 - Temperature: 37°C (98.0°F)
 - Probe type: YSI-400 series Temperature Probes (Probe accuracy: $\pm 0.1^{\circ}\text{C}$)
5. After the normal power-up sequence, verify that the temperature reads $37^{\circ}\text{C} \pm 0.1^{\circ}\text{C}$ ($98.6^{\circ}\text{F} \pm 0.2^{\circ}\text{F}$ if Fahrenheit is selected for the temperature units).
6. Turn off the monitor.

Note: The accuracy of temperature measurements is $\pm 0.1^{\circ}\text{C}$ ($\pm 0.2^{\circ}\text{F}$) in the range of 25°C to 45°C and $\pm 0.2^{\circ}\text{C}$ in the range of 15°C to less than 25°C . In the procedure above, add the tolerance of the simulator and the probe to the acceptable range of readings.

IBP Operation

1. Connect the monitor to an AC power source.
2. Turn on the monitor by pressing the *Power button*.
3. Connect the IBP test cable to the IBP simulator, then turn on the IBP simulator.
4. Press *6, zero button* on the IBP simulator, then press *enter* button on the IBP simulator.
5. Connect the other end of IBP test cables to IBP channel 1 and 2 connectors on the monitor's front panel after completed the POST.
6. The monitor will display IBP waveform's baseline on the screen.
7. Select IBP waveform menu or IBP menu, then set *Pressure Zero Setting*, to *Yes* to perform the zero calibration.
8. If zero calibration is successful, the monitor will display the value "0" on the IBP numerical area.
9. Press the *5, wavef button* on the IBP simulator, then press *enter* button on the IBP simulator.
10. The monitor will:
 - display IBP waveform on the screen.
 - display systolic, mean, diastolic measurement values on the IBP numerical area.

Note: Initial values of IBP simulator are systolic 120mmHg, diastolic 80mmHg at channel 1, systolic 120mmHg, diastolic 0mmHg at channel 2.

CO₂ Operation

1. Connect the monitor to an AC power source. Turn on the monitor.
2. Rotate the trim knob to highlight CO₂ numerical area and then press the trim knob to display the CO₂ menu.
3. Verifying that *Capno Measurement* is set to *On* in CO₂ menu.
4. Test #1: Display Accuracy
 - a. Connect the 10% calibration gas cylinder with the valve to the sampling tube connected to the water trap of the monitor.
 - b. After passed the warm up time (about 3 minutes), repeat 3 or 4 times turning on and off the valve of the gas cylinder with the interval of 1 to 2 seconds.
 - c. Verifying that the monitor displays 76mmHg \pm 2mmHg in CO₂ numerical area.
5. Test #2: Flow Rate
 - a. Connect the flow meter to the sampling tube connected to the monitor.
 - b. Connect a sampling tube to the other side of the flow meter and make the other side of the sampling tube open.
 - c. Verify that the flow rate is between 150 \pm 2mmHg.
6. Test #4: Occlusion
 - a. Block the sample input. The module should enter the "occlusion" state. Verify that it does correctly enter this occlusion state.
 - b. Unblock the input. Verify that after the modules occlusion sequence of high flow rates and valve actuations the occlusion message disappears and the module restores normal function.
7. Test #5: EtCO₂ Calibration

To perform a Zero Calibration:

The user is required to apply a 0% CO₂ gas, or "zero" gas, during the steps of a Zero Calibration.

- a. Select a well ventilated room to perform the calibration.
- b. Make sure the monitor has been operating for at least 5 minutes prior to the Zero Calibration.
- c. Attach the CO₂ scrubber to the water trap inlet according to the CO₂ scrubber "Directions for Use" (or attach a 0% medical-grade gas source). Now the monitor operates for one minute.
- d. After approximately one minute, observe the CO₂ reading. The CO₂ reading should be between 0.0%~0.3% with well-ventilated room air.
- e. Proceed with the zero gas phase of the calibration as defined by the host system.
- f. Disconnect the CO₂ scrubber (or gas source) from the water trap after the zero gas calibration.

To perform the Two Points User Calibration:

- a. Select a well-ventilated room to perform the calibration.
- b. Let the monitor run for at least 5 minutes prior to calibration.
- c. For the "Sample Zero Reference Gas" step, attach the CO₂ scrubber to the water trap inlet according to the CO₂ scrubber "Directions for Use" (or use a 0% medical-grade gas source).
- d. Proceed with the zero gas phase of the calibration as defined by the host system. Disconnect CO₂ scrubber (or 0% gas source) from the water trap after the zero gas phase and prior to the next calibration step.
- e. For the "Sample Span Reference Gas" phase of the calibration, provide a medical-grade gas source with a known CO₂ concentration between 8% and 12%, regulated to a pressure between 5 and 7 psi.
- f. Introduce a steady stream of CO₂ gas, Set and monitor the flow rate to approximately 1 liter/minute $\pm 10\%$). To verify flow, place the bleed line of the calibration sample line in a glass of water. If bubbles emerge, gas supply is sufficient. If no bubbles emerge, there is an insufficient supply of gas.
- g. Connect a sample line from the sample elbow, or Tee, as shown above, to the monitor's water trap inlet. Make sure the bleed line is directed away from the monitor. Allow the reference gas to flow for at least one minute.
- h. Sample the Span Reference Gas.
- i. After the Two-Point User Calibration steps have been followed, disconnect the monitor from the gas supply and test setup.
- j. A verification may be performed using the same gas delivery set up. Verify the observed CO₂ gas reading is within 3 mmHg or 10%, whichever is greater, of the CO₂ value of the test gas supplied. (If not, refer the user to the appropriate troubleshooting information.)

Safety Tests

The monitor safety tests meet the standards of, and are performed in accordance with, IEC 60601-1, Clause 19 (Second Edition, 1988; Amendment 1, 1991-11, Amendment 2, 1995-03), EN60601-1 for instruments classified as Class I and Type CF.

Protective Earth Continuity

This test checks the integrity of the power cord ground wire from the AC plug to the instrument chassis ground. The current used for this test is less than or equal to 4 Volts RMS, 50 to 60 Hz, and 25 Amperes.

1. Connect the monitor AC mains plug to the analyzer recommended by the analyzer operating instructions.
2. Connect the analyzer resistance input lead to the equipotential terminal (ground lug) on the rear of the instrument. Verify that the analyzer indicates 100 milliohms or less.

Electrical Leakage

Earth Leakage Current

This test is in compliance with IEC60601-1 earth leakage current. The applied voltage for IEC60601-1 the voltage is 264 Volts AC, 50 to 60 Hz. All measurements shall be made with the power switch in both "On" and "Off" positions.

1. Connect the monitor AC plug to the electrical safety analyzer as recommended by the analyzer operating instructions.
2. Perform the test as recommended by analyzer operating instructions.

Table 3. Earth Leakage Current Values

Test Condition	Allowable Leakage Current (microamps)
Normal Condition (NC)	500
SFC Open Supply (SFC OS)	1000
Normal Condition RM (NCRM)	500
SFC Open Supply RM (SFC OSRM)	1000

SFC: Single Fault Condition / RM: Reverse Mains/Lines Voltage

Note: Earth leakage current is measured under various conditions of the AC mains and protective earth conductor. For each condition, the measured leakage current must not exceed that indicated in Table 3.

Enclosure Leakage Current

This test is in compliance with IEC60601-1 enclosure leakage current. This test is for ungrounded enclosure current, measured between enclosure parts and earth. The applied voltage for IEC60601-1 the applied voltage is 264 Volts AC at 50 to 60 Hz.

1. Connect the monitor AC plug to the electrical safety analyzer as recommended by the analyzer operating instructions.
2. Place a 200cm² foil in contact with the instrument case making sure the foil is not in contact with any metal parts of the enclosure that may be grounded.
3. Measure the leakage current between the foil and earth.

Note: The analyzer leakage current indication must not exceed the values listed in Table 4.

Table 4. Enclosure Leakage Current

Test Condition	Allowable Leakage Current (microamps)
Normal Condition (NC)	100
SFC Open Supply (OS)	500
SFC Open Earth (SFC OE)	500
Normal Condition RM (NCRM)	100
SFC Open Supply RM (SFC OSRM)	500
SFC Open Earth RM (SFC OERM)	500

Patient Leakage Current

This test measures patient leakage current in accordance with IEC60601-1, clause 19, for Class I, Type CF equipment. Patient leakage current in this test is measured from any individual patient connection to earth (power ground).

1. Configure the electrical safety analyzer as recommended by the analyzer operating instructions.
2. Connect the monitor's AC mains power cord to the analyzer as recommended by the analyzer operating instructions.
3. Connect the ECG test cable between the ECG connector on the monitor and the appropriate input connector on the analyzer.
4. Turn on the monitor.
5. Perform the patient leakage current test as recommended by the analyzer operating instructions.
6. Repeat the patient leakage current test for the SpO₂ and temperature patient connections, using the appropriate test cables.

Note: Patient leakage current is measured under various conditions of the AC mains and protective earth conductor. For each condition, the measured leakage current must not exceed that indicated in Table 5.

Note: This test requires a test cable for each patient connector. For example, the ECG test cable consists of the ECG cable connector, with all conductors shorted together, connected to a test lead from the electrical safety analyzer. Test cables for SpO₂ and temperature can be configured in a similar manner, by wrapping each sensor end individually with aluminum foil filled with conductive gel (only enough gel to ensure conductivity). Attach a wire to the foil that is connected to a test lead from the electrical safety analyzer.

Table 5. Patient Leakage Current Values

Test Condition	Allowable Leakage Current (microamps)
Normal Condition (NC)	10
SFC Open Supply (OS)	50
SFC Open Earth (SFC OE)	50
Normal Condition RM (NCRM)	10
SFC Open Supply RM (SFC OSRM)	50
SRC Open Earth RM (SFC OERM)	50

Patient Leakage Current - Mains Voltage on the Applied Part

WARNING: AC mains voltage will be present on the applied part terminals during this test. Exercise caution to avoid electrical shock hazard.



WARNING: Do not touch the patient leads clips or the simulator parts connected to patient leads during this test as an electrical shock will occur.

This test measures patient leakage current in accordance with IEC60601-1, clause 19, for Class I, type CF equipment. In this test, 110% of mains voltage is applied between each patient connection and earth (power ground). Patient leakage current is then measured from any individual patient connection to earth.

Note: Keep the patient test cable length as short as possible during the leakage test.

Note: This test requires the same test cables for each patient connector as described in paragraph "Patient Leakage Current".

1. Configure electrical safety analyzer as recommended by analyzer operating instructions.
2. Connect monitor's AC mains power cord to analyzer as recommended by analyzer operating instructions.
3. Connect ECG test cable between ECG connector on the monitor and appropriate input connector on analyzer.
4. Turn on the monitor.
5. Perform the test as recommended by analyzer operating instructions.
6. Repeat the test for SpO₂ and temperature patient connections, using appropriate test cables.

Note: Patient leakage current is measured with normal and reverse mains polarity. For each condition, the measured leakage current must not exceed that indicated in Table 6.

Table 6. Patient Leakage Current Values - Mains Voltage on Applied Part

Test Condition	Allowable Leakage Current (microamps)
Normal polarity (SFC)	50
Reverse polarity (SFCRM)	50

Patient Auxiliary Current

This test measures patient auxiliary current in accordance with IEC60601-1, clause 19, for Class I, type CF equipment. The applied voltage for IEC60601-1 the voltage is 264 volts, 50 to 60 Hz. Patient auxiliary current is measured between each ECG test lead and between each sensor connection for all possible connections.

Note: Keep the patient test cable length as short as possible during the leakage test.

Note: This test requires the same test cables for each patient connector as described in paragraph “**Patient Leakage Current**”.

1. Configure the electrical safety analyzer as recommended by the electrical analyzer’s operating instructions.
2. Connect the monitor’s AC mains power cord to the electrical analyzer as recommended by the electrical analyzer’s operating instructions.
3. Connect the patient test lead combination in Table 7 to the appropriate input connector on the electrical analyzer.
4. Turn on the monitor.
5. Perform patient auxiliary current test per Table 8 as recommended by electrical analyzer’s operating instructions.
6. Repeat the patient auxiliary current test for each test lead combination as listed in Table 7 and measure each patient auxiliary current.

Table 7. Test Lead Combinations

First Test Lead	Second Test Lead
ECG #1 (LA)	ECG #2 (LL)
ECG #1 (LA)	ECG #3 (RA)
ECG #2 (LL)	ECG #3 (RA)
ECG #1 (LA)	Temperature 1/2
ECG #2 (LL)	Temperature 1/2
ECG #3 (RA)	Temperature 1/2
ECG #1 (LA)	SpO ₂
ECG #2 (LL)	SpO ₂
ECG #3 (RA)	SpO ₂
Temperature 1/2	SpO ₂

Table 8. Allowable Leakage Current

Test Condition	Allowable Leakage Current (microamps)
Normal Condition (NC)	10
SFC Open Supply (OS)	50
SFC Open Earth (SFC OE)	50
Normal Condition RM (NCRM)	10
SFC Open Supply RM (SFC OSRM)	50
SRC Open Earth RM (SFC OERM)	50

Service Menu and Factory Default Settings

General

This section discusses use of the Service menu to configure 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, Alarm Reminder Tone, System Setting System Test, NIBP Test, Demo Mode and EtCO₂ Calibration. Also this section explains briefly the factory default settings.

Service Menu

The purpose of the Service menu (Figure 10, Table 9) is to allow the authorized user to create a Power-on default for the settings in effect each time the monitor is powered on. Once the Service menu is entered, physiological monitoring is terminated. The screen layouts do not display any information associated with normal monitoring operation. Use the following procedure to configure the Service Menu for the monitor (also see **Using the Monitor** section, of the operator's manual):

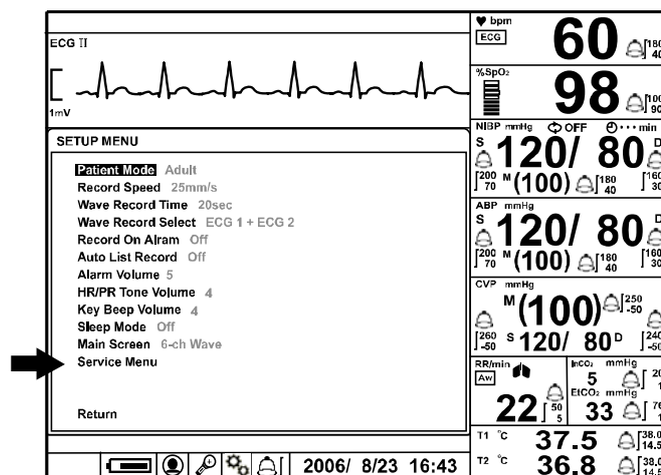


Figure 10. The access of Service Menu via Set-up menu

1. Set the monitor to normal monitoring mode.
2. Rotate the trim knob to highlight the *Setup icon* located on the bottom of the screen, and then press the trim knob. *Setup menu* displays.
3. Rotate the trim knob to highlight *Service Menu* in *Setup menu*, and then press the trim knob to access the *Service Menu*.
4. Three digits are displayed in the Level 2 Menu as shown in Figure 4.

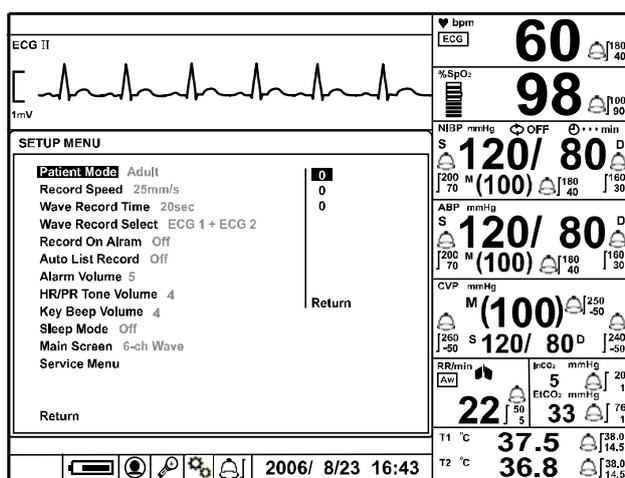


Figure 11. The access of Service Menu via Set-up menu

Note: The access code is **6, 0, 6**. It is set at the factory and may not be changed.

5. Rotate the trim knob to highlight the top of the digits. Press the trim knob to enter *Pass code*.
6. Rotate the trim knob until “6” appears, then press the trim knob.
7. Repeat step 5-6 to enter all the access code “6” “0” “6”.

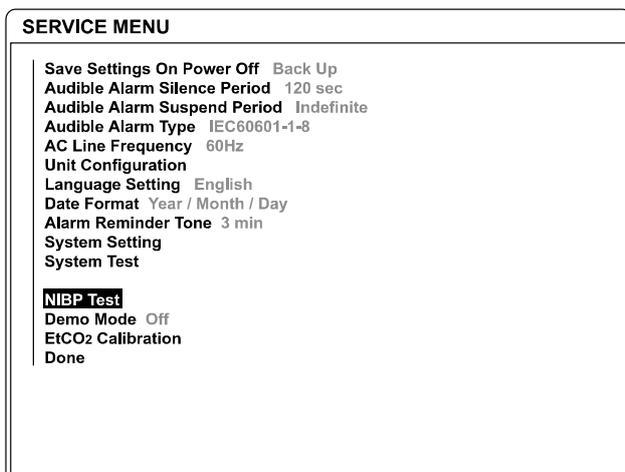


Figure 12. Service Menu

8. The Service Menu will now be present. The available Service Menu items are explained in Figure 6 and Table 9. Make changes to these menu items as desired by rotating and pressing the trim knob.
9. Select “Done”. The monitor will present the message “All changes made to the power-up defaults will be in effect the next time the monitor is turned on.” Before turned off.
10. Turn off the monitor, and then turn on the monitor again.

Note: The monitor must be powered off upon selecting “Done” to save any changes into the monitor, and then the changes made to the Power on defaults will be in effect next time the monitor is powered up.

Table 9. Service Menu

Level 1 Menu	Level 2 Menu	Level 3 Menu
Saved setting on Power off	Custom	
	Back Up	
	Default	
Audible Alarm Silence Period	60, 60, 90, 120 sec	
Audible Alarm Suspend Period	Off, 10, 20, 30, 60 min, Indefinite	
Audible Alarm Type	GN924, IEC60601-1-8	
AC Line Frequency	50Hz, 60Hz	
Unit Configuration	NIBP	mmHg, kPa
	IBP	mmHg, kPa
	CO ₂	mmHg, %, kPa
	Temperature	°C, °F
Language Setting	한국어 (Korean), 中文 (Chinese), English, Français (French), Deutsch (German), Italiano (Italian), 日本語 (Japanese), Português (Portuguese), Español (Spanish)	
Date Format	Year/Month/Day, Month/Day/Year, Day/Month/Year	
Alarm Reminder Tone	Off, 3, 10 min	
System Setting	Monitor On Time	
	Recorder On Time	
	Battery Deep Discharges	
	System Software Version	
	Module Version	
	Print value of configuration	
	Tone Set (Detail)	
System Test	LCD Test	
	Alarm Audible Test	
	Tone Audible Test	
	Backup RAM Clear	
NIBP Test	Pressure Sensor Accuracy Test	
	Air Leakage Test	
	Inflation Speed Test	
	Deflations Speed Test	
Demo Mode	On, Off	
EtCO ₂ Calibration	Zero Calibration, Calibration	
Done	The monitor will be powered off upon selecting "Done", then any changes in the service menu will be in effect next time the unit is powered up.	

Saved setting on Power off

If the save settings on power off is set to *Custom*, the monitor does not save the settings when the monitor is power off. Therefore, the first settings when the monitor is powered up become the power-up defaults in the next power-on cycling.

If the save settings on power off is set to *Back Up*, the monitor saves the current settings when the monitor is powered off. The saved settings become the power-on defaults in the next power-on cycling.

If the save settings on power off is set to *Default*, the factory default settings become the power-up defaults in the next power-on cycling.

Audible Alarm Silence Period

Pressing the *Alarm silence/suspension button* temporarily silences alarms for the period selected in the service menu. The factory default of alarm silence period is 120 seconds..

Audible Alarm Suspension Period

If *Alarm suspension period* is set to other than *Off* or *Indefinite*, the audible alarm is not activated for the time interval by pressing and holding the *Alarm silence/suspension button* for 2 seconds. If *Off* is selected, the audible alarm suspension is not allowed to activate. If *Indefinite* is selected, the audible alarm suspension continues until canceled.

Audible Alarm Type

The audible alarm has different tone pitch and on-off beep patterns for each alarm priority. The monitor has two different audible alarm types, called GN924 and IEC60601-1-8.

Table 10. Audible Alarm Characteristics

Alarm Category	Tone Pitch		Beep Rate	
	GN924	IEC60601-1-8	GN924	IEC60601-1-8
High priority	~976 Hz	~976 Hz	7 beeps in 2 sec	10 beeps in 15 sec
Medium priority	~697 Hz	~697 Hz	2 beeps in 1 sec	3 beeps in 15 sec
Low priority	~488 Hz	~488 Hz	1 beeps in 5 sec	1 beeps in 30 sec

AC Line Frequency

The monitor supports *AC line frequency* both 50 Hz and 60 Hz. Select either *50Hz* or *60 Hz* for an appropriate AC line.

Unit Configuration

The unit configuration of each parameter can be set in the service menu.

Language Setting

The selected language will be used for all the text shown on the display; and it will be effective the next time the monitor is powered up.

Date Format

The date format is selectable; Year/Month/Day, Month/Day/Year, Day/Month/Year

Alarm Reminder Tone

The monitor generates the alarm reminder tone at the preset interval to remind the user that the audible alarm is suspended. The interval can be set to *Off*, *3* or *10 minutes* via the service menu. If *Off* is selected, the reminder tone will be disabled.

System Setting

This menu displays the information for system settings.

Monitor On Time

This menu displays the number of hours, minutes and seconds, rounded to the nearest hour that the monitor has been operational. The term that the monitor is in the service mode is not included.

Recorder On Time

This menu displays the number of hours, minutes and seconds, rounded to the nearest hour that the Recorder has been operational.

Battery Deep Discharges

This menu displays the number of deep-discharge cycles seen by the battery. The monitor records a deep discharge cycle when the monitor is automatically turned off after a "Critically Low Battery" alarm is issued.

System Software Version

This menu displays the revision level of the system software. The revision level is also shown on the LCD as part of the Copyright screen.

Module Version

This menu displays information for each module or board version.

Print Value of Configuration

This menu is used to print out the set values. This function is useful when collecting a wrong value.

Tone Set (Detail)

This menu used to set the HR/PR tone with High, Medium, Low or SpO₂.

Note: The values of Monitor On Time, Recorder On Time and Battery Deep Discharges may not be reset, but they will be reset to zero when a new CPU module is installed.

System Test

These menus provide to facilitate performing verification testing for the overall system.

For a detailed procedure, refer to **Performance Verification** section.

Table 11. System Test

Tests	Description
LCD Test	tests the LCD display.
Alarm Audible Test	tests the alarm tones.
Tone audible Test	tests the HR/PR tones, the key beeps and the completion sounds.
Backup RAM Clear	is used to set the factory defaults.

NIBP Test

These menus provide to facilitate performing verification testing for the NIBP subsystem.

For a detailed procedure, refer to **Performance Verification** section.

Table 12. NIBP Test

Tests	Description
Pressure Sensor Accuracy Test	verifies that the pneumatic pressure sensor accuracy is within the specification.
Air Leakage Test	verifies that the pneumatic pressure air leakage is within a pressure drop of 3 mmHg/min.
Inflation Speed Test	is intended for factory use only.
Deflation Speed Test	is intended for factory use only.
Return	returns to Service Menu after the test completed.

Demo Mode

The purpose of Demo Mode is to show a visual presentation demonstrating how the monitor works.

EtCO₂ Calibration

The EtCO₂ calibration may be performed periodically to ensure product performance and measurement accuracy. This menu provides to facilitate performing EtCO₂ calibration. For a detailed procedure, refer to *Performance Verification* section.

Factory Default Settings

Factory default settings are divided into three groups, adult, pediatric and neonatal as described in Table 13.

The patient mode is preset to “Adult” mode. Alarm limits settings will be automatically changed to the default settings for each patient mode as the mode is changed to Adult, Pediatric or Neonatal mode.

Table 13. Factory Default Settings

Parameters	Ranges/Selections	Factory Defaults		
		Adult	Pediatric	Neonatal
NIBP				
NIBP Initial Inflation Pressure	adult/pediatric 120, 140, 160, 180, 200, 220mmHg (16.0, 18.7, 21.3, 24.0, 26.7, 29.3 kPa) neonatal 80, 100, 120, 140mmHg (13.3, 16.0, 17.3, 18.7 kPa)	180 mmHg (24.0 kPa)	180 mmHg (24.0 kPa)	120 mmHg (16.0 kPa)
BP On Alarm	On, Off	Off	Off	Off
NIBP Interval	Off, Cont, 1, 2, 2.5, 3, 5, 10, 15, 20, 30, 45, 60, 90, 120, 180 minutes	Off	Off	Off
NIBP SYS High Alarm Limits	adult/pediatric 35 to 270 mmHg (4.7 to 36.0 kPa) neonatal 45 to 130 mmHg (6.0 to 17.3 kPa) (5 mmHg, 0.7kPa steps)	160 mmHg (21.3 kPa)	120 mmHg (16.0 kPa)	90 mmHg (12.0 kPa)
NIBP SYS Low Alarm Limits	adult/pediatric 30 to 265 mmHg (4.0 to 36.3 kPa) neonatal 40 to 125 mmHg (5.3 to 16.7 kPa)	90 mmHg (12.0 kPa)	70 mmHg (9.3 kPa)	40 mmHg (5.3 kPa)

Parameters	Ranges/Selections	Factory Defaults		
		Adult	Pediatric	Neonatal
	(5 mmHg, 0.7kPa steps)			
NIBP DIA High Alarm Limits	adult/pediatric 15 to 250 mmHg (2.0 to 33.3 kPa) neonatal 25 to 90 mmHg (3.3 to 12.0 kPa) (5 mmHg, 0.7kPa steps)	90 mmHg (12.0 kPa)	70 mmHg (9.3 kPa)	60 mmHg (8.0 kPa)
NIBP DIA Low Alarm Limits	adult/pediatric 10 to 245 mmHg (1.3 to 32.7 kPa) neonatal 20 to 85 mmHg (2.7 to 11.3 kPa) (5 mmHg, 0.7kPa steps)	50 mmHg (6.7 kPa)	40 mmHg (5.3 kPa)	20 mmHg (2.7 kPa)
NIBP MAP High Alarm Limits	adult/pediatric 25 to 260 mmHg (3.3 to 34.7 kPa) neonatal 35 to 110 mmHg (4.7 to 14.7 kPa) (5 mmHg, 0.7kPa steps)	110 mmHg (14.7 kPa)	90 mmHg (12.0 kPa)	70 mmHg (9.3 kPa)
NIBP MAP Low Alarm Limits	adult/pediatric 20 to 255 mmHg (2.7 to 34.0 kPa) neonatal 30 to 105 mmHg (4.0 to 14.0 kPa) (5 mmHg, 0.7kPa steps)	60 mmHg (8.0 kPa)	50 mmHg (6.7 kPa)	30 mmHg (4.0 kPa)
IBP				
IBP Sweep Speed	12.5, 25.0, 50.0 mm/s	25.0 mm/s	25.0 mm/s	25.0 mm/s
Pressure Zero Setting	Yes, No	No	No	No
P1 Label	P1, ABP	ABP	ABP	ABP
P1 Scale	0~50, 0~100, 0~200, 0~300, AUTO	AUTO	AUTO	AUTO
IBP 1 SYS High Alarm Limits	adult/pediatric/neonatal -45 to 300 mmHg (-6 to 40 kPa) (5 mmHg, 0.7kPa steps)	160 mmHg (21.3 kPa)	120 mmHg (16.0 kPa)	90 mmHg (12.0 kPa)
IBP 1 SYS Low Alarm Limits	adult/pediatric/neonatal -50 to 295mmHg (-6.7 to 39.3 kPa) (5 mmHg, 0.7kPa steps)	90 mmHg (12.0 kPa)	70 mmHg (9.3 kPa)	40 mmHg (5.3 kPa)
IBP 1 DIA High Alarm Limits	adult/pediatric/neonatal -45 to 300 mmHg (-6 to 40 kPa) (5 mmHg, 0.7kPa steps)	90 mmHg (12.0 kPa)	70 mmHg (9.3 kPa)	60 mmHg (8.0 kPa)
IBP 1 DIA Low Alarm Limits	adult/pediatric/neonatal -50 to 295 mmHg (-6.7 to 39.3 kPa) (5 mmHg, 0.7kPa steps)	50 mmHg (6.7 kPa)	40 mmHg (5.3 kPa)	20 mmHg (2.7 kPa)
IBP 1 MEAN High Alarm Limits	adult/pediatric/neonatal -45 to 300 mmHg (-6 to 40 kPa) (5 mmHg, 0.7kPa steps)	110 mmHg (14.7 kPa)	90 mmHg (12.0 kPa)	70 mmHg (9.3 kPa)
IBP 1 MEAN Low Alarm Limits	adult/pediatric/neonatal -50 to 295 mmHg (-6.7 to 39.3 kPa) (5 mmHg, 0.7kPa steps)	60 mmHg (8.0 kPa)	50 mmHg (6.7 kPa)	30 mmHg (4.0 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
IBP 2 SYS High Alarm Limits	adult/pediatric/neonatal -45 to 300 mmHg (-6 to 40 kPa) (5 mmHg, 0.7kPa steps)	160 mmHg (21.3 kPa)	120 mmHg (16.0 kPa)	90 mmHg (12.0 kPa)
IBP 2 SYS Low Alarm Limits	adult/pediatric/neonatal -50 to 295mmHg (-6.7 to 39.3 kPa) (5 mmHg, 0.7kPa steps)	90 mmHg (12.0 kPa)	70 mmHg (9.3 kPa)	40 mmHg (5.3 kPa)
IBP 2 DIA High Alarm Limits	adult/pediatric/neonatal -45 to 300 mmHg (-6 to 40 kPa) (5 mmHg, 0.7kPa steps)	90 mmHg (12.0 kPa)	70 mmHg (9.3 kPa)	60 mmHg (8.0 kPa)
IBP 2 DIA Low Alarm Limits	adult/pediatric/neonatal -50 to 295mmHg (-6.7 to 39.3 kPa) (5 mmHg, 0.7kPa steps)	50 mmHg (6.7 kPa)	40 mmHg (5.3 kPa)	20 mmHg (2.7 kPa)
IBP 2 MEAN High Alarm Limits	adult/pediatric/neonatal -45 to 300 mmHg (-6 to 40 kPa) (5 mmHg, 0.7kPa steps)	110 mmHg (14.7 kPa)	90 mmHg (12.0 kPa)	70 mmHg (9.3 kPa)
IBP 2 MEAN Low Alarm	adult/pediatric/neonatal	60 mmHg	50 mmHg	30 mmHg

42 Service Menu and Factory Default Settings

Parameters	Ranges/Selections	Factory Defaults		
		Adult	Pediatric	Neonatal
Limits	-50 to 295mmHg (-6.7 to 39.3 kPa) (5 mmHg, 0.7kPa steps)	(8.0 kPa)	(6.7 kPa)	(4.0 kPa)
ECG				
ECG Cable Select	3 Leads, 5 Leads, AUTO	AUTO	AUTO	AUTO
ECG Lead Select	I, II, III, aVR, aVL, aVF, V(Chest Lead)	-	-	-
ECG Size (mm/mV)	Auto, 1.25, 1.7, 2.5, 5.0, 7.5, 10.0, 15.0, 20.0	10.0	10.0	10.0
ECG Filter Mode	Monitor, Low Extend, Filter	Monitor	Monitor	Monitor
ECG Pacer Detect	On, Off	Off	Off	Off
ECG Sweep Speed	12.5, 25.0, 50.0 mm/s	25.0 mm/s	25.0 mm/s	25.0 mm/s
HR/PR Source	Auto, HR, PR	Auto	Auto	Auto
HR/PR High Alarm Limits	25 to 250 BPM (5 BPM steps)	120 BPM	160 BPM	200 BPM
HR/PR Low Alarm Limits	20 to 245 BPM (5 BPM steps)	50 BPM	75 BPM	100 BPM
SpO₂				
C-Lock	On, Off	Off	Off	Off
SpO ₂ Sweep Speed	12.5, 25.0, 50.0 mm/s	25.0 mm/s	25.0 mm/s	25.0 mm/s
% SpO ₂ High Alarm Limits	21 to 100 % (1 % steps)	100 %	100 %	95 %
% SpO ₂ Low Alarm Limits	20 to 99 % (1 % steps)	90 %	90 %	80 %
Respiration				
Apnea Time Setting	Off, 20, 30, 40, 60, Step 60, Step 90	30 sec	30 sec	30 sec
Respiration/Apnea	Off, AUTO, awRR, imRR	AUTO	AUTO	AUTO
Respiration Size (mm/mV)	Level 1, 2, 3, 4, 5, 6, 7, 8	Level 6	Level 6	Level 6
Respiration Sweep Speed	6.25, 12.5, 25.0 mm/s	12.5 mm/s	12.5 mm/s	12.5 mm/s
RESP High Alarm Limits	4 to 150 BPM (1 BPM steps)	30 BPM	30 BPM	100 BPM
RESP Low Alarm Limits	3 to 149 BPM (1 BPM steps)	8 BPM	8 BPM	30 BPM
Capnography				
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
Capno Measurement	On, Off	On	On	On
O ₂ Gas	On, Off	Off	Off	Off
N ₂ O Gas	On, Off	Off	Off	Off
EtCO ₂ High Alarm Limits	2 to 80 mmHg (Adult/Neo) (2 mmHg steps) 0.3 to 10.7 kPa (Adult/Neo) (0.3 kPa steps) 0.3 to 10.5 % (Adult/Neo) (0.3 % steps)	80 mmHg 10.7 kPa 10.5 %	80 mmHg 10.7 kPa 10.5 %	80 mmHg 10.7 kPa 10.5 %
EtCO ₂ Low Alarm Limits	0 to 78 mmHg (Adult/Neo) (2 mmHg steps) 0 to 10.4 kPa (Adult/Neo) (0.3 kPa steps) 0 to 10.3 % (Adult/Neo) (0.3 % steps)	0 mmHg 0 kPa 0 %	0 mmHg 0 kPa 0 %	0 mmHg 0 kPa 0 %
InCO ₂ High Alarm Limits	2 to 20 mmHg (Adult/Neo) (2 mmHg steps) 0.3 to 2.7 mmHg (Adult/Neo) (0.3 kPa steps) 0.3 to 2.6 % (Adult/Neo) (0.3 % steps)	20 mmHg 2.7 kPa 2.6 %	20 mmHg 2.7 kPa 2.6 %	20 mmHg 2.7 kPa 2.6 %
InCO ₂ Low Alarm Limits	0 to 18 mmHg (Adult/Neo) (2 mmHg steps) 0 to 2.4 kPa (Adult/Neo) (0.3 kPa steps) 0 to 2.3 % (Adult/Neo) (0.3 % steps)	0 mmHg 0 kPa 0 %	0 mmHg 0 kPa 0 %	0 mmHg 0 kPa 0 %
Temperature				
TEMP1 High Alarm Limits	15.1 to 45.0 °C (0.1°C steps) 59.1 to 113.0 °F (0.1°F steps)	39.0 °C (102.2 °F)	39.0 °C (102.2 °F)	39.0 °C (102.2 °F)
TEMP1 Low Alarm Limits	15.0 to 44.9 °C (0.1°C steps) 59.0 to 112.8 °F (0.1°F steps)	36.0 °C (96.8 °F)	36.0 °C (96.8 °F)	36.0 °C (96.8 °F)
TEMP2 High Alarm Limits	15.1 to 45.0 °C (0.1°C steps) 59.1 to 113.0 °F (0.1°F steps)	39.0 °C (102.2 °F)	39.0 °C (102.2 °F)	39.0 °C (102.2 °F)

Parameters	Ranges/Selections	Factory Defaults		
		Adult	Pediatric	Neonatal
TEMP2 Low Alarm Limits	15.0 to 44.9 °C (0.1°C steps) 59.0 to 112.8 °F (0.1°F steps)	36.0 °C (96.8 °F)	36.0 °C (96.8 °F)	36.0 °C (96.8 °F)
Others				
Patient Mode	Adult, Pediatric, Neonatal	Adult		
Record Speed**	25.0 mm/s, 50.0 mm/s	25.0 mm/s		
Wave Record Time**	20 sec, Continuous (10sec delay)	20 sec		
Wave Record Select**	ECG1 + ECG2, PLETH, RESP, IBP1, IBP2 or CAPNO	ECG1 + PLETH		
Record on Alarm**	On,Off	Off		
Auto List Record**	On,Off	Off		
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	Off		
Main Screen	4ch-wave, 6ch-wave, Big Number	-		
Alarm Limits Display	On, Off	On		
Auto Alarm	On, Off	Off		
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	1 min		
Graphical Display On/Off	On/Off for each parameter	On		
Save Setting on Power Off*	Custom, Back up, Default	Back up		
Audible Alarm Silence Period*	30, 60, 90, 120 sec	120 sec		
Audible Alarm Suspend Period*	Off, 10, 20, 30, 60 min, Indefinite (Alarm Inhibition)	Indefinite		
Audible Alarm Type*	GN924, IEC60601-1-8	GN924		
AC Line Frequency*	50, 60 Hz	60 Hz		
NIBP Unit*	mmHg, kPa	mmHg		
IBP Unit*	mmHg, kPa	mmHg		
CO ₂ Unit*	mmHg, %, kPa	mmHg		
Temperature Unit*	°C, °F	°C		
Date Format*	year/month/day, month/day/year, day/month/year	year/month/day		
Alarm Reminder Tone*	Off, 3, 10 min	3 min		
EtCO ₂ Calibration*	On, Off	Off		
Language*	한국어 (Korean), 中文 (Chinese), English, Français (French), Deutsch (German), Italiano (Italian), 日本語 (Japanese), Português (Portuguese), Español (Spanish)	English		

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 monitor service manual

Note: Asterisks (**) by a parameter in the above table indicate the print settings when the optional printer is installed in the monitor

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Firmware Download

General

This section is for the purpose for reloading Firmware into the monitor when the possibility of corrupted Firmware exists, or updating Firmware with a new system revision (system/device version). Call Mediana Technical Service Department for the latest version of Firmware utility required.

Equipment Needed

Table 14 lists the equipment required for Firmware download.

Table 14. Required Equipments for Firmware download

Equipment	Description
Firmware Downloading Cable	LAN Cable
Firmware Downloading Software	YM6000 Field Utility
Personal Computer (PC)	With TCP/IP Port

How to Download

1. Turn the monitor off.
2. Turn the monitor on by pressing the *Power button*.
3. The monitor displays the bar while the checksum for the flash memory is in progress.
4. Press *NIBP start/stop button* and *Return button* simultaneously before filled the bar completely.
5. The monitor will display the firmware download screen as shown below figure 13.

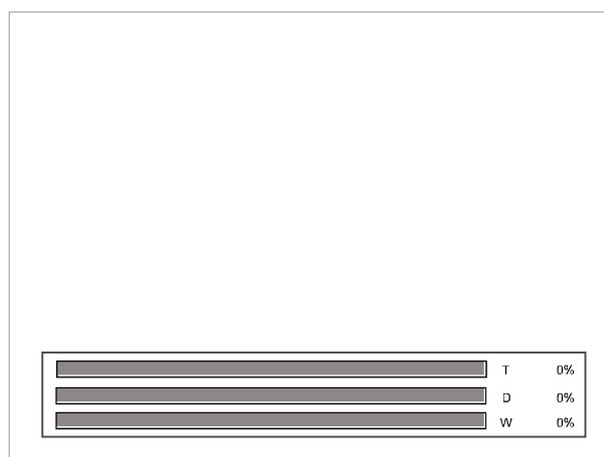


Figure 13. Firmware downloading display

6. Connect LAN cable to the TCP/IP port of the monitor and the PC.
7. Run **YM6000 Field Utility.exe** on the PC.
8. Three bars on the screen indicate the progress status for the downloading.
 - “T” displays the transferring status the PC to monitor.
 - “D” displays the deleting status of the flash memory.
 - “W” displays the writing status of the flash memory..
9. When three bars are filled, the completion code will be displayed in the box located on the center of the screen as shown Figure 14. Refer to Table 15.

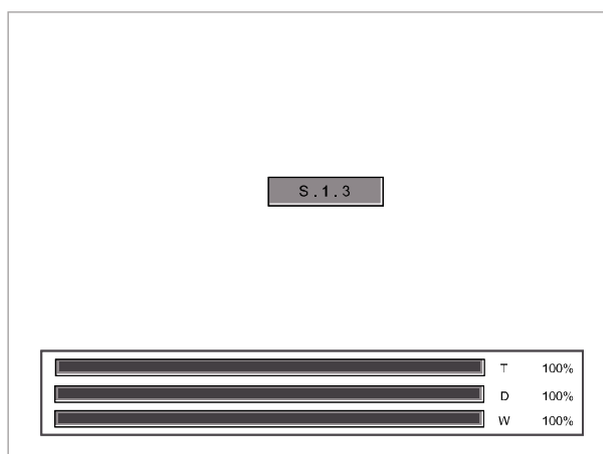


Figure 14. Firmware downloading display

Table 15. Completion codes

Code	Description
S. 1. 3	Boot download completed.
S. 2. 3	Main download completed.

Note: If any problem during Firmware downloading, the error code will be displayed in the box located on the center of the screen. Refer to Firmware Download in **Troubleshooting** section.

10. After completion of downloading, turn the monitor off.
11. Disconnect the LAN cable from the monitor.
12. After a few seconds, turn the monitor on again.
13. Perform the tests specified in **Performance Verification** section.

Note: When a new firmware downloading is completed, the monitor is set to the factory default.

Troubleshooting

General

This section provides information that can be helpful in troubleshooting the YM6000 monitor.

How to Use This Section

Use this section in conjunction with **Performance Verification** section and **Spare Parts** section. To remove and replace a part suspected a trouble, follow the instructions in **Disassembly Guide** section.

Who Should Perform Repairs

Only qualified service personnel should open the monitor housing, remove and replace components, or make adjustments in accordance with this service manual. If your medical facility does not have qualified service personnel, contact Mediana Technical Service Department.

Replacement Level Supported

The replacement level supported for this product is to the printed circuit board (PCB assembly) and major subassembly level. Once you isolate a suspected PCB assembly, follow the procedures in **Disassembly Guide** section, to replace the PCB assembly with a known good PCB assembly. Check to see if the trouble symptom disappears and that the monitor passes all performance tests.

If the trouble symptom persists, swap back the replacement PCB assembly with the suspected malfunctioning PCB assembly (the original PCB assembly that was installed when you started troubleshooting) and continue troubleshooting as directed in this section.

Obtaining Replacement Parts

Mediana Technical Service Department provides technical assistance information and replacement parts. To obtain replacement parts, contact Mediana. Refer to the part names and part numbers listed in **Spare Parts** section.

Troubleshooting Guide

Problems with the monitor are separated into the categories for further troubleshooting instructions.

Note: Taking the recommended actions discussed in this section will correct the majority of problems you will encounter. However, problems not covered here can be resolved by calling Mediana Technical Service Department.

Power

Power problems are related to AC and/or Battery as follows. If the action requires replacement of the components, refer to **Disassembly Guide** section.



CAUTION: Electrical shock hazard. Disconnect a power cord from the monitor before attempting to open or disassemble the monitor.

1. **When AC power cord is connected to the monitor, AC indicator on the front panel is not lit.**
 - 1-1 Ensure power cord is plugged into the appropriate AC outlet of voltage and frequency.
 - 1-2 Replace the power supply assembly.
 - 1-3 Ensure the key wire is connected to the key board assembly and the main board. If the connection is good, replace the key board assembly.

2. **The monitor fails to power-up when the power button is pressed.**
 - 2-1 Ensure power cord is plugged into operational AC outlet of the appropriate voltage and frequency. Ensure AC power indicator is lit. If indicator is not lit, replace the power supply assembly.
 - 2-2 Check fuses located on the power supply assembly above AC inlet receptacle. Replace fuses if necessary.
 - 2-3 Inside monitor, check the main cable and ensure that it is connected to the main board and the power supply assembly.
 - 2-4 Ensure the key board assembly is connected to the main board. If connection is good, replace the key board assembly.
 - 2-5 If the problem persists, replace the main board.

3. **The monitor turns on, then sounds an alarm and shuts off and no error code is displayed.**
 - 3-1 Replace the main board.

4. The monitor does not operate when disconnected from AC power.

4-1 The battery may be discharged. To recharge the battery, connect a power cord to AC mains over 12 hours (refer to Battery charge in **Performance Verification** section).

Note: DC power should not charge the battery. The monitor may be used with a less than fully charged battery but with a corresponding decrease in operating time from that charge. The battery may be defective.

4-2 If problem persists, replace battery fuse on the power supply assembly.

4-3 Inside monitor, check the main cable and ensure that it is connected to the main board and the power supply assembly

4-4 If the problem persists, replace the main board.

5. Battery does not charge.

5-1 If the battery fails to hold a charge, replace the battery as indicated in **Battery Replacement**.

5-2 Open the monitor as described in **Disassembly Guide**. Verify the power supply's LED flashes while on AC. While charging battery, the red LED is flashed until battery charging is completed, then the red LED is lit continuously. Replace the power supply assembly if the LEDs are not lit.

Display

The followings are symptoms of problems relating to non-functioning displays, and recommended actions. If the action requires replacement of a PCB assembly or module, refer to **Disassembly Guide**.

1. LCD screen is totally black after system powers-up.

1-1 Ensure the inverter wire is connected to the LCD assembly.

1-2 Ensure the LCD cable is connected to the LCD assembly and the main board.

1-3 If the problem persists, replace LCD assembly.

1-4 If the problem persists, replace the main board.

2. LCD screen is illuminated, but no data is visible after system powers-up.

2-1 Ensure LCD cable is connected to the main board and LCD assembly.

2-2 If the problem persists, replace the LCD cable.

2-3 If the problem persists, replace the main board.

2-4 If the problem persists, replace the LCD assembly.

3. Display values are missing or erratic.

3-1 If the measurement cables are connected, ensure that the cables are properly connected.

3-2 If the problem persists, replace the measurement cables.

3-3 If the problem persists, replace LCD cable.

3-4 If the problem persists, replace the main board.

4. Display pixels do not light.

- 4-1 Check the connection of the main board.
- 4-2 If the problem persists, replace the LCD cable.
- 4-3 If the problem still persists, replace the main board.
- 4-4 If the problem still persists, replace the LCD assembly.

Sound

The followings are symptoms of problems and recommended actions relating to sound. If the action requires replacement of a PCB assembly or module, refer to **Disassembly Guide**.

1. Alarm sounds for no apparent reason.

- 1-1 Moisture or spilt liquids can cause an alarm not to sound. Allow the monitor to dry thoroughly before using.
- 1-2 If the problem persists, replace the main board.

2. Audible alarm does not sound.

- 2-1 Check alarm silence status.
- 2-2 Check alarm volume setting in Setup menu.
- 2-3 Ensure the speaker is connected to the power supply assembly.
- 2-4 If problem persists, replace speaker.
- 2-5 Check the main cable between main PCB and the power supply assembly.
- 2-6 If problem persists, replace the power supply assembly.
- 2-7 If problem persists, replace the main board.

3. The monitor responds to button press, but key press tone fails to sound.

- 3-1 Check the key beep volume setting in Setup menu.
- 3-2 Ensure speaker is connected to main the power supply assembly.
- 3-3 If problem persists, replace speaker.
- 3-4 Check the main cable between the main board and the power supply assembly.
- 3-5 If problem persists, replace the power supply assembly.
- 3-6 If problem persists, replace the main board.

Trim knob and Buttons

The following are symptoms of problems and recommended actions relating to buttons, If the action requires replacement of a PCB assembly, refer to **Disassembly Guide**.

1. **The monitor fails to power-up when Power button is pressed.**
 - 1-1 Take steps as noted in paragraph Power in **Performance Verification** section.
2. **The monitor powers-up, but some or one of the other buttons respond.**
 - 2-1 Ensure the key wire is connected to the key board assembly and the main board. If connection is good, change the key board assembly.
 - 2-2 If problem persists, change the main board.
3. **When the trim knob is rotated, no highlight appears on display screen, and/or the monitor does not respond to the trim knob presses.**
 - 3-1 Ensure the encoder wire is plugged into the main board. If connection is good, change encoder assembly.
 - 3-2 If problem persists, replace the main board.

NIBP

The following are symptoms of problems relating to NIBP functions and recommended actions. If the action requires replacement of a PCB or module, refer to **Disassembly Guide**.

1. **The cuff does not inflate.**
 - 1-1 Ensure the cuff hose is not folded and NIBP tube inside of the monitor is not blocked or kinked.
 - 1-2 If problem persists, replace the NIBP module.
 - 1-3 If problem persists, replace the main board.
2. **Alarm Message C11 or E03**
 - 2-1 Ensure the cuff hose is not folded and NIBP tube inside of the monitor is not blocked or kinked.
 - 2-2 If problem persists, replace the NIBP module.
 - 2-3 If problem persists, replace the main board.
3. **The NIBP value is not reliable.**
 - 3-1 Perform the pressure sensor accuracy test and air leakage test.
 - 3-2 If problem persists, replace the NIBP module.

SpO₂

The following are symptoms of problems relating to SpO₂ functions and recommended actions. If the action requires replacement of a PCB or module, refer to **Disassembly Guide**.

- 1. The SpO₂ sensor does not light.**
 - 1-1 Ensure the connection between the sensor and extension cable is not loose. If the connection is good, replace the SpO₂ module.
 - 1-2 If problem persists, replace the main board.
- 2. The SpO₂ sensor is lighting but no display on SpO₂ screen.**
 - 2-1 Replace the SpO₂ module.
 - 2-2 If problem persists, replace the main board.

Temperature

The following are symptoms of problems relating to Temperature functions and recommended actions. If the action requires replacement of a PCB or module, refer to **Disassembly Guide**.

- 1. The measurement values are too low.**
 - 1-1 Check the temperature probe is properly located at patient's body.
- 2. TEMP 1 or 2: Internal error is displayed.**
 - 1-1 Replace the main board.

Respiration

The following are symptoms of problems relating to Respiration functions and recommended actions. If the action requires replacement of a PCB or module, refer to **Disassembly Guide**.

- 1. Respiration signal (imRR) is noisy or the value is not displayed.**
 - 1-1 Ensure ECG accessories are connected properly or the electrode is not contaminated.
 - 1-2 If problem persists, replace the ECG board.
 - 1-3 If problem persists, replace the main board.
- 2. The monitor does not measure the respiration (awRR).**
 - 2-1 Ensure the hose inside the monitor
 - 2-2 If problem persists, replace the EtCO₂ module
 - 2-3 If problem persists, replace the main board.

IBP

The following are symptoms of problems and recommended actions relating to IBP functions. If the action requires replacement of a PCB assembly, refer to **Disassembly Guide**.

1. The ABP and CVP window does not exist on the LCD.

1-2 Please go to the Main screen of setup icon and set the display window to 6-ch Wave.

2. The IBP values are erroneous.

2-3 Check the IBP accessories connection between patient and each end.

2-4 Ensure the "Pressure Zero setting" is done or perform the "Pressure Zero setting" at IBP menu.

2-5 If the problem persists, replace the IBP accessories.

2-6 If the problem persists, replace the IBP board.

EtCO₂

The following are symptoms of problems and recommended actions relating to EtCO₂ functions. If the action requires replacement of a PCB assembly, refer to **Disassembly Guide**.

1. The EtCO₂ is not displayed at LCD

1-1 Please remind that it takes 3 minutes to warm up the EtCO₂ module after turning on the monitor.

1-2 Please go to the Main screen of setup icon and set the display window to 6-ch Wave.

1-3 Check the accessory is properly connected to the monitor and measurement circuit.

1-4 If the problem persists, replace the EtCO₂ accessories.

1-5 If the problem persists, replace the EtCO₂ module.

Firmware Download

If the error code appears during the firmware downloading, take the following action.

Table 16. Firmware Downloading Error Codes

Code	Description	Action
I. 1. 3 / E. 1. 3	When an error is occurred during transferred data.	Check the LAN cable and settings then Retry boot download
I. 2. 3 / E. 2. 3	When an error is occurred during transferred of data.	Check the LAN cable and settings then Retry boot download
E. 1. 5	When an error is occurred during recorded data.	Retry boot download
E. 2. 5	When an error is occurred during recorded data.	Retry main download
E. 1. 7	When the data is broke down during boot download.	Retry boot download
E. 2. 7	When the data is broke down during main download.	Retry main download

Note: If the Error Code still appears, take monitor out of service and contact Mediana Technical Service Department for advice on remedial action

Technical Alarm Conditions

When the monitor detects the technical alarm condition, the monitor will display the alarm message on the screen.

If the alarm message occurs, follow the check items in the below table to remove the alarm condition after checking the alarm message in the alarm message area or informative message area.

Table 17. Alarm Messages and Check Items

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 (C12)	Blood pressure could not be measured even after cuff pressure 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 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 artifact (C13)	Measurement failed because of patient movement during 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	Measurement failed because of insufficient pressurizing. There is

Alarm Messages	Check Items
pressure (C14)	<p>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.</p> <p>When it occurs in the initial measurement in continuous mode, the second measurement will start unless Stop button is pressed.</p>
NIBP: Cuff irregular pulses (C15)	<p>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.</p> <p>When it occurs in the initial measurement in continuous mode, the second measurement is continued unless Stop button is pressed.</p>
NIBP: Cuff motion artifact (C16)	<p>Blood pressure could not be measured because noises interrupted pulse waveform signal. Check for motion artifacts, or external vibration and then, measure again.</p> <p>When it occurs in the initial measurement in continuous mode, the second measurement will start unless Stop button is pressed.</p>
NIBP: Cuff time-out (C17)	<p>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.</p>
NIBP: Cuff time-out, over 160 pulses (C18)	<p>Pulse waveform signal more than 160 beats are detected during measurement. There is possibility that noises might interrupt signal.</p> <p>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.</p>
NIBP: Cuff pressure failure (C19)	<p>Cuff pressure exceeded more than 300 mmHg (adult) during measurement.</p> <p>There is a possibility that the patient moved during measurement or strong pressure from outside might be added to cuffs. Considering above, measure again.</p>
NIBP: Cuff weak pulse (C20)	<p>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.</p> <p>Wrap cuffs around properly, then, measure again.</p>
NIBP: Check cuff, hose and mode (C21)	<p>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.</p> <p>Check the measurement mode setting and application of the cuff, and measure again.</p>
NIBP: Internal error (E03)	<p>NIBP module error BPM pressure sensor fault.</p> <p>Pump operated for ten seconds, however pressure does not change. Check the connection of the cuff hose.</p>
NIBP: Internal error (E07)	<p>Reboot the monitor. If the problem persists, cease use immediately and contact qualified service personnel or your local supplier.</p>

Alarm Messages	Check Items
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 supplier.
NIBP: Internal error (RAM)	Reboot the monitor. If the problem persists, cease use immediately and contact qualified service personnel or your local supplier.
NIBP: Internal error (COM)	Reboot the monitor. If the problem persists, cease use immediately and contact qualified service personnel or your local supplier.
ECG: Internal error.	Reboot the monitor. If the problem persists, cease use immediately and contact qualified service personnel or your local supplier.
RESP: Internal error.	Reboot the monitor. If the problem persists, cease use immediately and contact qualified service personnel or your local supplier.
[label]: 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. There may be a problem with fitting of the SpO ₂ sensor, or blood flow at the sensor site may be unsatisfactory. Check the condition of the patient and fitting of the sensor, and measure again.
SpO ₂ : Internal error.	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. Stop using the monitor immediately and contact qualified service personnel or your local supplier.
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-battery condition.	Connect the AC power cord of the monitor to the AC main to recharge the battery.
SYSTEM: Real time clock error.	Reboot the monitor. If the problem persists, cease use immediately 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

Alarm Messages	Check Items
	local supplier.
SYSTEM: RAM error.	Reboot the monitor. If the problem persists, cease use immediately and contact qualified service personnel or your local supplier.
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 & electrodes.	ECG error is detected, Electrodes or lead wires may not be 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 disconnected.	Cable not connected to the main unit. If connected, the cable 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.
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 probe disconnected.	Sensor not connected to the main unit. If connected, the cable 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.
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.	Electrodes or lead wires may not be 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}: Unable to zero calibration.	Could not zero calibrate pressure. Check that the transducer is open to the atmosphere, and check the three-way tap. As it is also possible that the measured pressure incorporates noise, check the measurement circuit.
{label}: Out of range.	A value outside the measurement range was obtained. As the transducer has been subject to abnormal pressure, check the measurement circuit.
SpO ₂ : 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.
SpO ₂ : 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

Alarm Messages	Check Items
	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 45.0°C). Adjust the ambient temperature and measure again.
CAPNO: Zero calibration range error	Try again the gas calibration.
CAPNO: Zero calibration signal unstable error	Try again the gas calibration.
CAPNO: High calibration range error	Try again the gas calibration.
CAPNO: High calibration signal unstable error	Try again the gas calibration.
SYSTEM: Low battery.	Plug the AC power cord to the AC main to recharge the battery.
NIBP: Retry, Check cuff/Patient (C12)	Blood pressure could not be measured even after cuff pressure 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 excessive artifact (C13)	Measurement failed because of patient movement during 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: Retry, Cuff insufficient pressure (C14)	Measurement failed because of insufficient pressurizing. There is a 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 pulses (C15)	Blood pressure could not be measured because oscillation graph was not normal. There is a possibility that patient movement or external vibration interrupted the measurement. Check whether a 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 artifact (C16)	Blood pressure could not be measured because noises interrupted 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 Stop button is pressed.
NIBP: Retry, Cuff time-out, over 160 pulses (C18)	Pulse waveform signal more than 160 beats are detected during 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 failure (C19)	Cuff pressure exceeded more than 300mmHg during measurement. There is a possibility that the patient moved during measurement or strong pressure from outside might be

Alarm Messages	Check Items
NIBP: Retry, Check cuff, hose and mode (C21)	added to cuffs. Considering above, measure again. 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 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.
SpO ₂ : Motion artifact.	SpO ₂ 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 paper.	In case the recorder door is open, close the door. In case the recorder paper is empty, insert new paper and close the door.
SYSTEM: Abnormally shut down last time.	The monitor has been abnormally shut down last time. When power is lost for less than 30 seconds, the monitor will preserve the current settings and trend data restored automatically before the power loss. However, if the power loss is over 30 seconds, the monitor will be back to the previous user settings (or the factory default settings) as per the 'save settings on power off' in the service menu. Contact qualified personnel in your facility or your local supplier.
SYSTEM: No recorder installed.	The recorder is not installed in your monitor. If required, contact your local supplier.

Note: If the alarm message still appears, contact your local supplier for advice on remedial action.

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Disassembly Guide



WARNING: Performance Verification. Do not place the monitor into operation after repair or maintenance has been performed, until all Performance Tests and Safety Tests listed in Performance Verification section of this service manual have been performed. Failure to perform all tests could result in erroneous monitor readings.



WARNING: Before attempting to open or disassemble the monitor, disconnect the power cord from the monitor.



CAUTION: Observe ESD (electrostatic discharge) precautions when working within the unit.

General

This section describes disassembly procedures with detailed disassembly instructions and illustrations. Disassembly Sequence Flow Chart that is used to access replaceable parts of the monitor is illustrated in Figure 15. The boxes on the flow chart represent the various components or sub-assemblies. A complete listing of the available spare parts and part numbers is in **Spare Parts** section. Follow the reverse sequence of the disassembly procedures for reassembly.

The monitor can be disassembled down to all major component parts, including:

- PCB assemblies
- acquisition modules, the power supply assembly & the LCD assembly
- battery
- cables & wires
- cases
- printer

The following tools are required:

- small, Phillips-head (+) screwdriver
- medium, Phillips-head (+) screwdriver
- needle-nose pliers

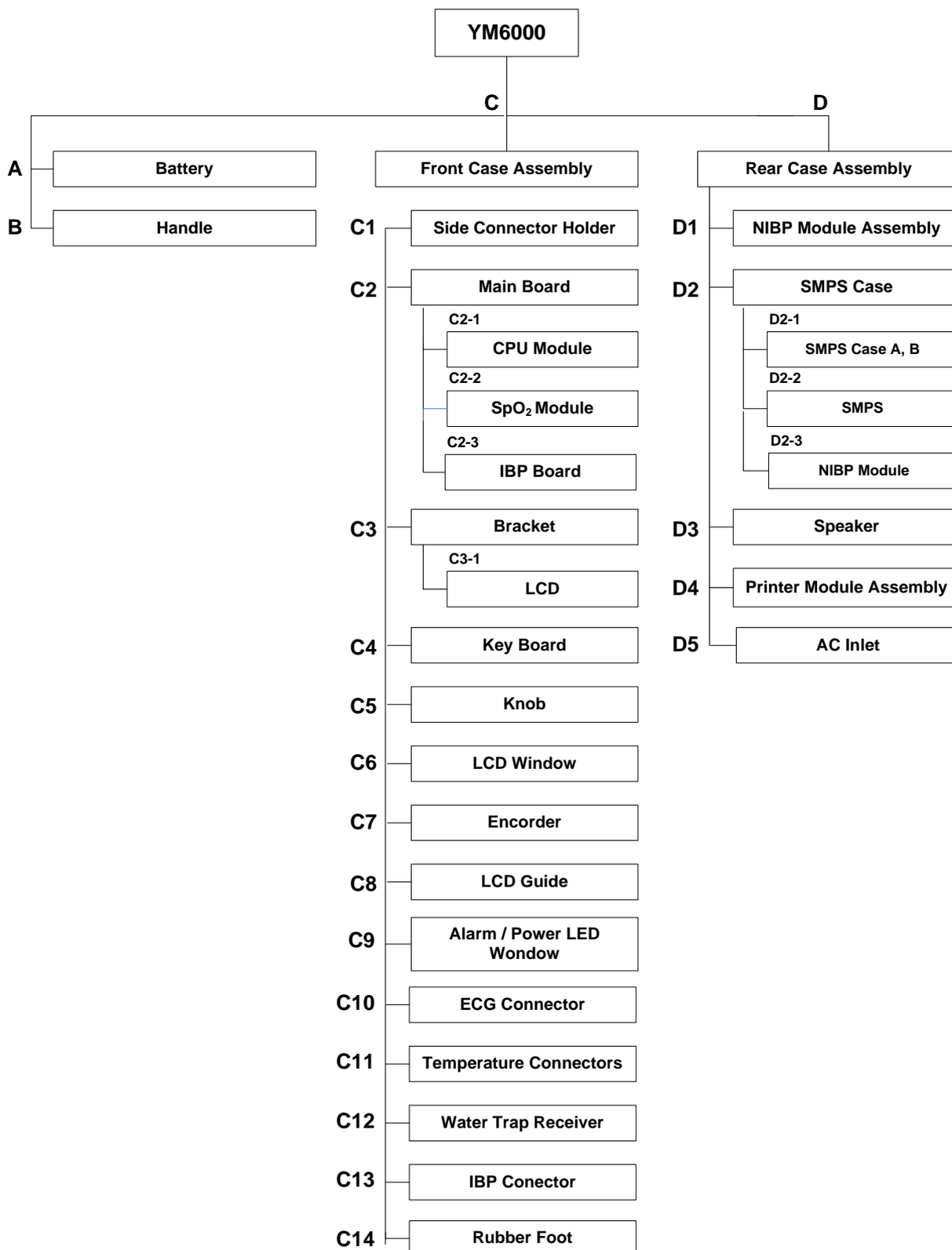


Figure 15. Disassembly Sequence Flow Chart

Replacement Level Supported

The replacement level supported for this product is to the printed circuit board (PCB) and major subassembly level. Once you isolate a suspected PCB, follow the procedures in Disassembly Guide, to replace the PCB with a known good PCB. Check to see if the trouble symptom persists, swap back the replacement PCB with the suspected malfunctioning PCB (the original PCB that was installed when you started trouble shooting) and continue troubleshooting as detected in this section.

Prior to Disassembly

1. Turn off the monitor by pressing *Power button*.
2. Disconnect the monitor from the AC power source.

Fuse Replacement

1. After step D2, remove 2 AC main fuses (F1, F2: 250V/6.3A) out of the socket if required.
2. Remove 2 battery fuses (F3, F4: 250V/6.3A) out of the socket if required.
3. Replace (a) new fuse(s).
4. Reassembly the monitor.

Battery Replacement

This section describes the steps to remove the battery from the monitor for replacement without disassembling the main case of the monitor

1. After step A, replace new batteries.
2. Reassemble the monitor.

Monitor Disassembly

This section describes the steps to remove battery and the handle and to separate the front and rear case assemblies.

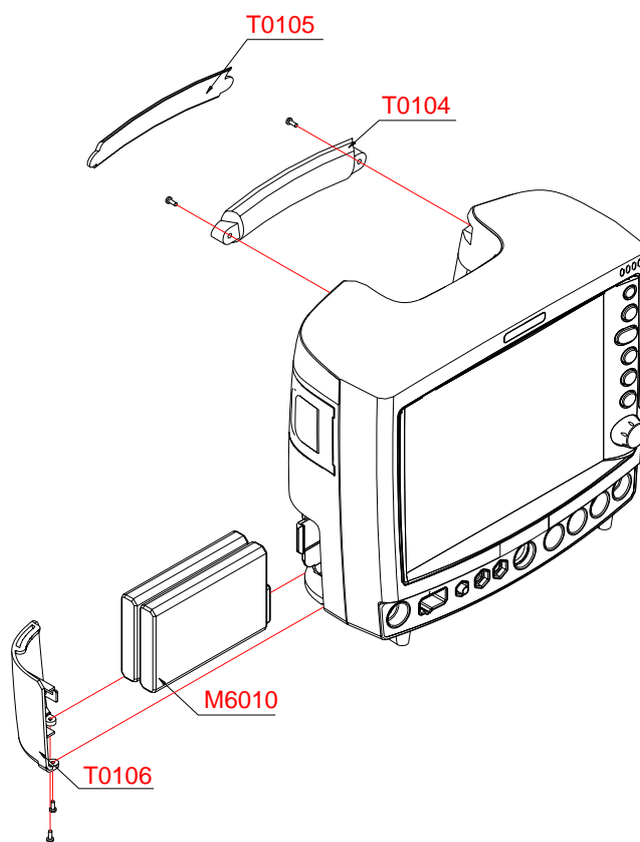


Figure 16. Battery and Handle Disassembly

Table 18. Part Descriptions – Battery and Handle

Part Codes	Descriptions
T0106	Battery Cover
M6010	Battery
T0104	Handle A
T0105	Handle B

A. Battery disassembly

1. Remove 2 screws (3 × 6) fastening the battery cover on the bottom of the monitor.
2. Remove the battery cover.
3. Remove a battery carefully from the monitor.

B. Handle disassembly

1. Pull the handle B out from the handle A.
2. Remove 2 screws (4 × 10) on the handle A.
3. Remove the handle A from the rear case.

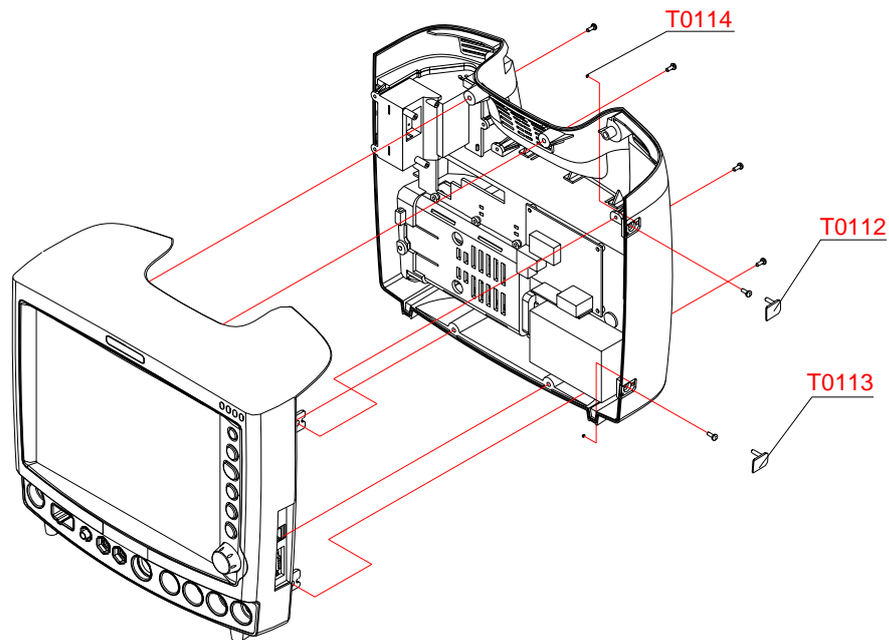


Figure 17. Monitor Disassembly

Table 19. Part Descriptions – Monitor

Part Codes	Descriptions
T0112	Side cap top
T0113	Side cap bottom
T0114	Side cap stopper

Before the steps C and D

1. Remove side caps top, bottom and stopper on the monitor.
2. Remove 6 screws (4 × 10) on the monitor.
3. Separate the front case assembly from the rear case assembly.
4. Disconnect the SMPS wire connected to the main board
5. Disconnect the main cable connected to the SMPS.
6. Disconnect the NIBP hose connected to the NIBP connector.

Front Case Disassembly (C)

This section describes the items that may be removed on the front case assembly.

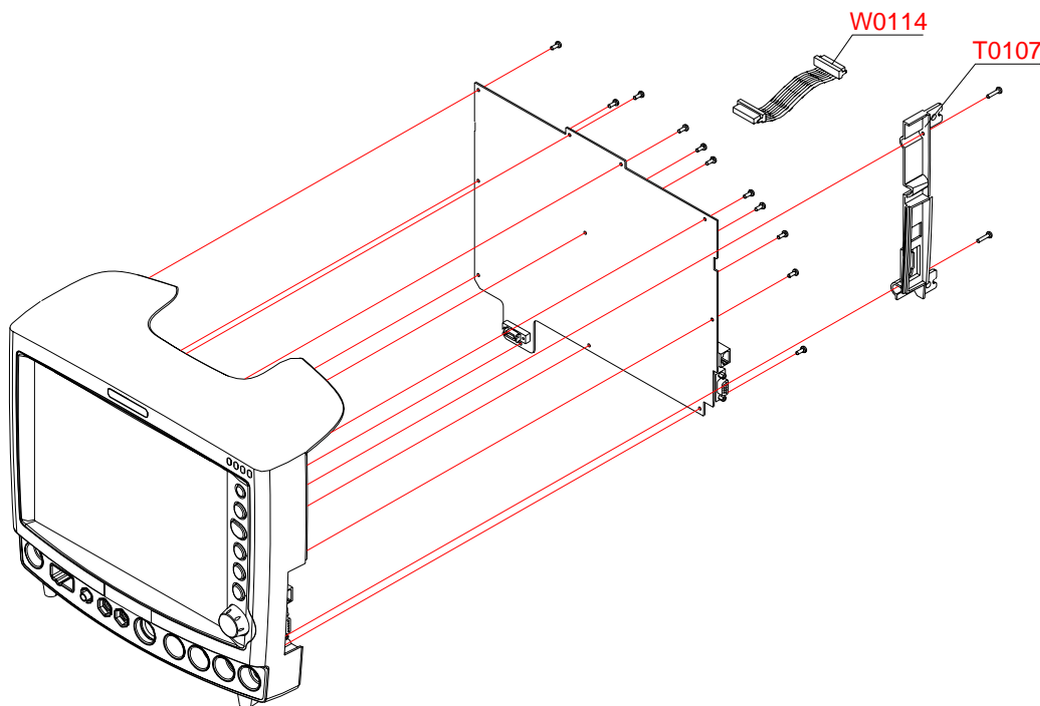


Figure 18. Front Case Disassembly – Side connector holder and Main board

Table 20. Part Descriptions – Side connector holder and Main board

Part Codes	Descriptions
T0107	Side connector holder
W0114	Main cable / 24 pin

C1. Side connector holder disassembly

1. Remove 2 screws (3 × 6) on the side connector holder.
2. Separate the side connector holder from the front case assembly.

C2. Main board disassembly

1. Disconnect the LCD cable, the inverter wire, the key wire, the encoder wire, ECG connector wire and two temperature wires connected to the main board.
2. Remove the main cable from the main board.
3. Remove 11 screws (3 × 6) on the main board.
4. Separate the main board from the front case assembly.

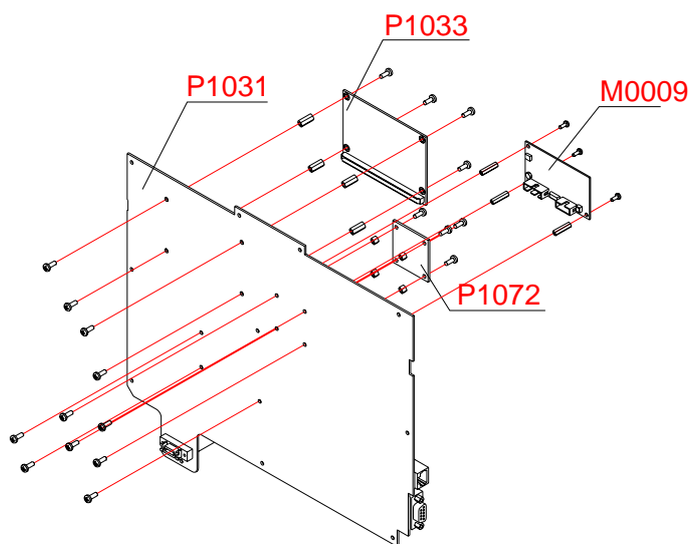


Figure 19. Front case disassembly – CPU module, SpO₂ module and IBP board

Table 21. Part Descriptions – CPU module, SpO₂ module and IBP board

Part Codes	Descriptions
P1031	Main board
P1033	CPU module
M0009	SpO ₂ module
P1072	IBP board

C2-1. CPU module disassembly

1. .After step C2, remove 4 screws (3 × 4) on the CPU module.
2. Remove the CPU module from the main board.
3. Remove 8 screws (3 × 4) to separate 4 supporters (5mm) from the main board.

C2-2. SpO₂ module disassembly

1. .After step C2, remove 3 screws (3 × 6) on the SpO₂ module.
2. Remove the SpO₂ module from the main board.
3. Remove 6 screws (3 × 4) to separate 3 supporters (12mm) from the main board.

C2-3. IBP board disassembly

4. .After step C2, remove 2 screws (3 × 6) on the IBP board..
1. Remove the IBP board from the main board.
2. Remove 2 screws (3 × 4) to separate 2 supporters (6mm) from the main board.

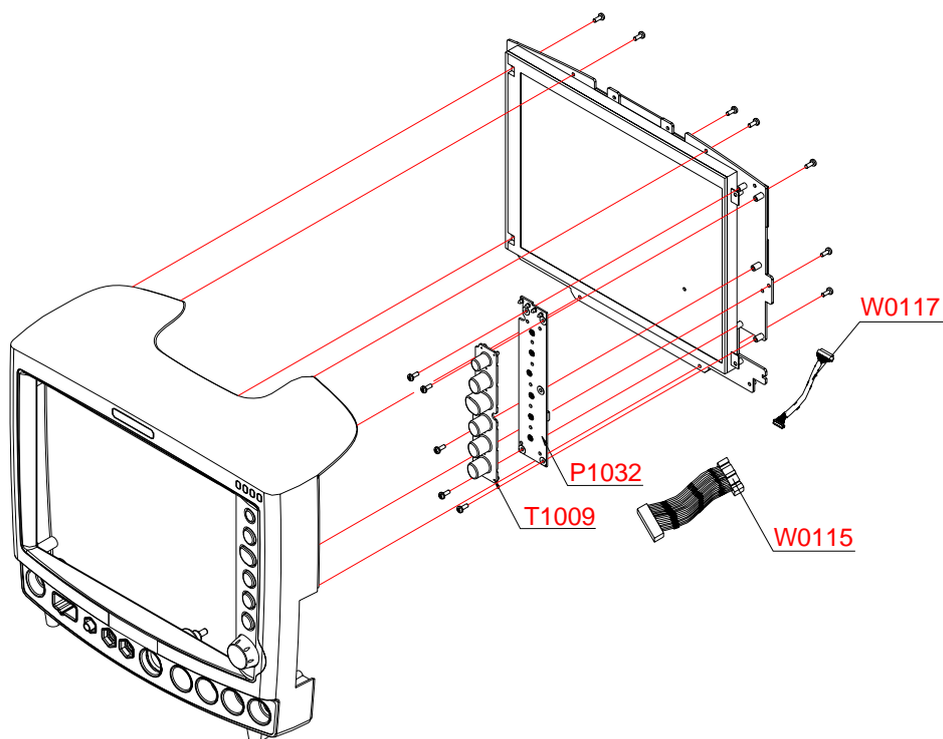


Figure 20. Front Case Disassembly –Bracket and Key board

Table 22. Part Descriptions –Bracket and Key board

Part Codes	Descriptions
T1009	Rubber Key
W0115	Key wire
P1032	Key board
W0117	LCD cable

C3. Bracket disassembly

1. Remove 7 screws (3 × 6) on the bracket.
2. Separate the bracket from the front case assembly.

C4. Key board disassembly

1. Remove the rubber key from the key board.
2. Remove 5 screws (3 × 6) on the key board.
3. Remove the key board from the bracket.
4. Remove the key wire form the key board.

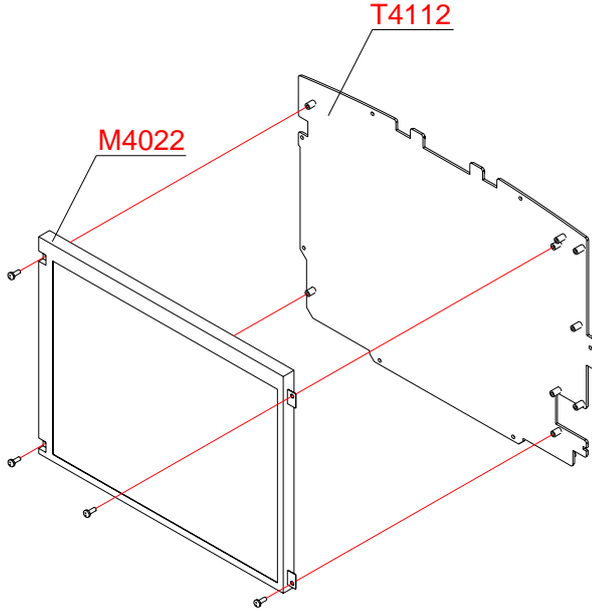


Figure 21. Front Case Disassembly – sLCD and Bracket

Table 23. Part Descriptions – LCD and Bracket

Part Codes	Descriptions
M4022	LCD
T4112	Bracket

C3-1. LCD disassembly

- 1. After step C3, remove 4 screws (3 × 6) on the LCD.
- 2. Remove the LCD from the bracket.

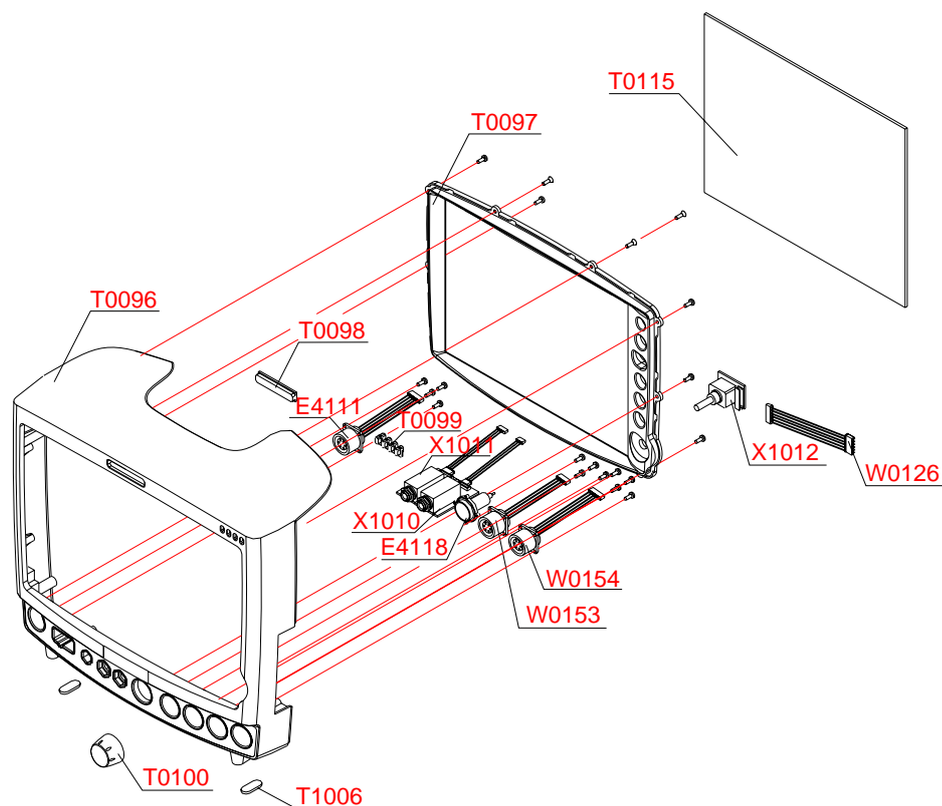


Figure 22. Front Case Disassembly – Knob, LCD window, Encoder assembly and Connectors

Table 24. Part Descriptions – Knob, LCD window, Encoder assembly and Connectors

Part Codes	Descriptions
T0096	Front case
T0100	Knob
T0115	LCD window
X1012	Encoder board assembly
W0126	Encoder wire
T0097	LCD guide
T0098	Alarm LED window
T0099	Power LED window
E4111	ECG connector / 6 pin
X1010	Temperature assembly / 3 pin
X1011	Temperature assembly / 4 pin
E4118	Water trap receiver assembly
W0153	IBP wire / 4 pin
W0154	IBP wire / 5 pin
T1006	Rubber foot

C5. Knob disassembly

1. Pull the trim knob straight out to remove from the front case.

C6. LCD window disassembly

1. Remove the LCD window from the front case.

C7. Encoder board disassembly

1. Turn the encoder board assembly to the left to remove from the front case.
2. After separate the encoder board, remove the encoder wire.

C8. LCD guide disassembly

1. Remove 8 screws (3 × 8) on the LCD guide.
2. Turn the encoder board assembly to the left to remove from the front case.

C9. Alarm LED window and Power LED window disassembly

1. Remove the alarm LED window form the front case.
2. Remove the power LED window from the front case.

C10. ECG connector disassembly

1. Remove 4 tapping screws (3 × 8) on the ECG connector.
2. Remove the ECG connector from the front case.

C11. Temperature connector disassembly

1. Remove two nuts (14mm) fastening the temperature connector from the front case.
2. Remove two temperature connectors from the front case.

C12. Water trap receiver disassembly

1. Pull the water trap receiver straight out to remove from the front case assembly.

C13. IBP connector disassembly

2. Remove 2 screws (3 × 8) on the IBP connector (4 pin).
1. Remove the IBP connector form the front case.
2. Remove 2 screws (3 × 8) on the IBP connector (5 pin)
3. Remove the IBP connector from the front case.

C14. Rubber foot disassembly

1. Remove two rubber foots from the front case.

Rear Case Disassembly (D)

This section describes the items that may be removed on the rear case assembly.

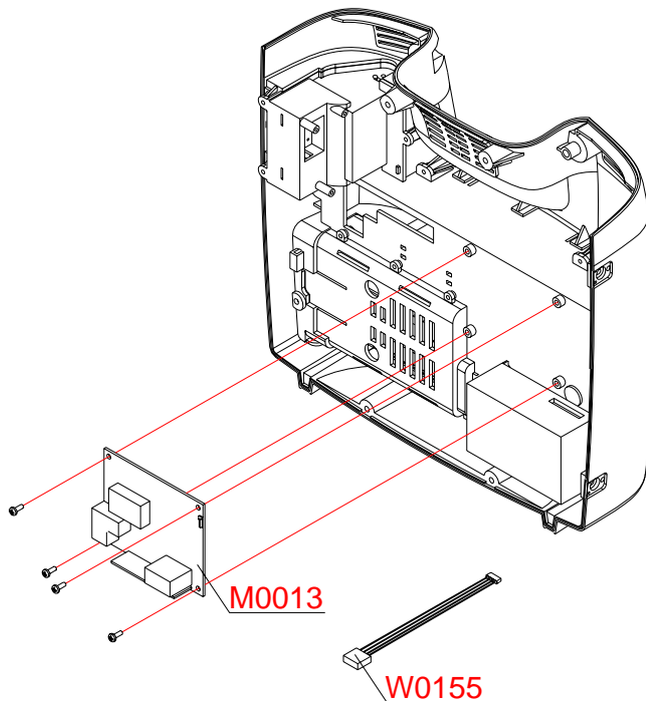


Figure 23. Rear case disassembly – EtCO₂ module

Table 25. Part Descriptions – EtCO₂ module

Part Codes	Descriptions
M0013	EtCO ₂ module
W0155	EtCO ₂ wire / 4 pin

D1. EtCO₂ module disassembly

1. Remove the EtCO₂ wire from the EtCO₂ module and the SMPS.
2. Remove 4 screws (3 × 6) on the EtCO₂ module.
3. Remove the EtCO₂ module from the SMPS.

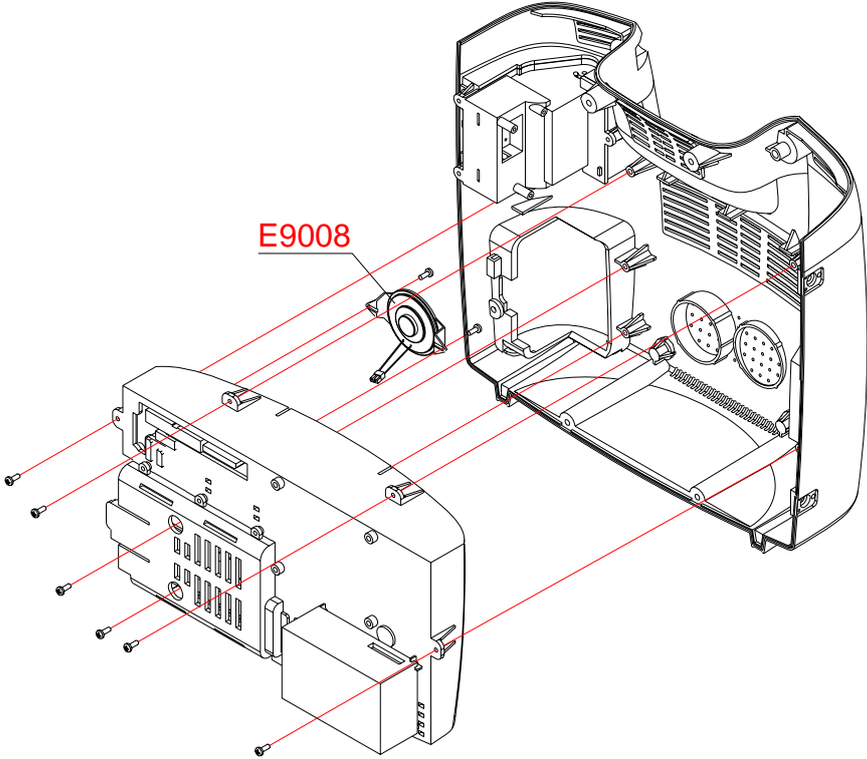


Figure 24. Rear case disassembly – Speaker

Table 26. Part Descriptions – Speaker

Part Codes	Descriptions
E9008	Speaker

D2. SMPS case disassembly

- 1. Remove 4 screws (3 × 6) on the SMPS case A.
- 2. Disconnect the AC inlet wire and the printer wire connected to the SMPS.
- 3. Separate the SMPS case from the rear case assembly.

D3. Speaker disassembly

- 1. Disconnect the speaker wire connected to the SMPS.
- 2. Remove 2 tapping screws (3 × 8) on the speaker.
- 3. Remove the speaker from the SMPS case A.

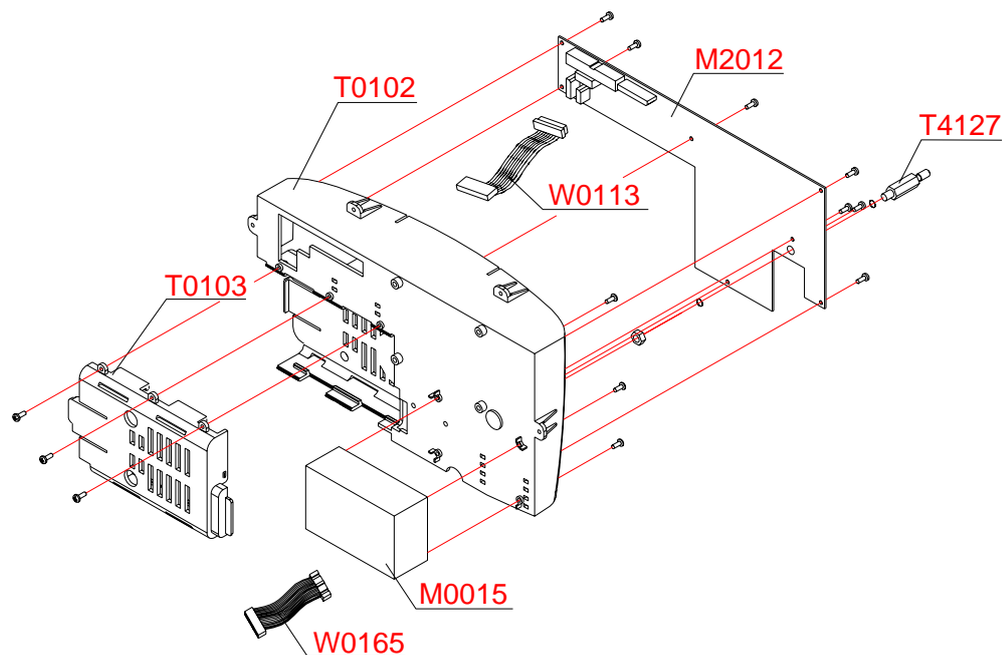


Figure 25. Rear case disassembly –SMPS case, SMPS and NIBP module

Table 27. Part Descriptions –SMPS case, SMPS and NIBP module

Part Codes	Descriptions	Qty
T0102	SMPS case A	1
T0103	SMPS case B	1
M2012	SMPS	1
W0113	SMPS wire	1
T4127	Ground terminal	1
M0015	NIBP module	1
W0165	NIBP cable	1

D2-1. SMPS case A, B disassembly

1. After step D2, remove 3 screws (3 × 8) on the SMPS case B.
2. Remove the SMPS case B from the SMPS case A.

D2-2. SMPS disassembly

1. After step D2, remove 7 screws (3 × 6) on the SMPS.
2. Remove the SMPS from the SMPS case A.
3. Remove the SMPS wire from the SMPS.
4. Remove the ground terminal from the SMPS.

D2-3. NIBP module disassembly

4. After D2, remove the remove 4 screws (3 × 6) on the NIBP module.
5. Remove the NIBP module from the SMPS case A

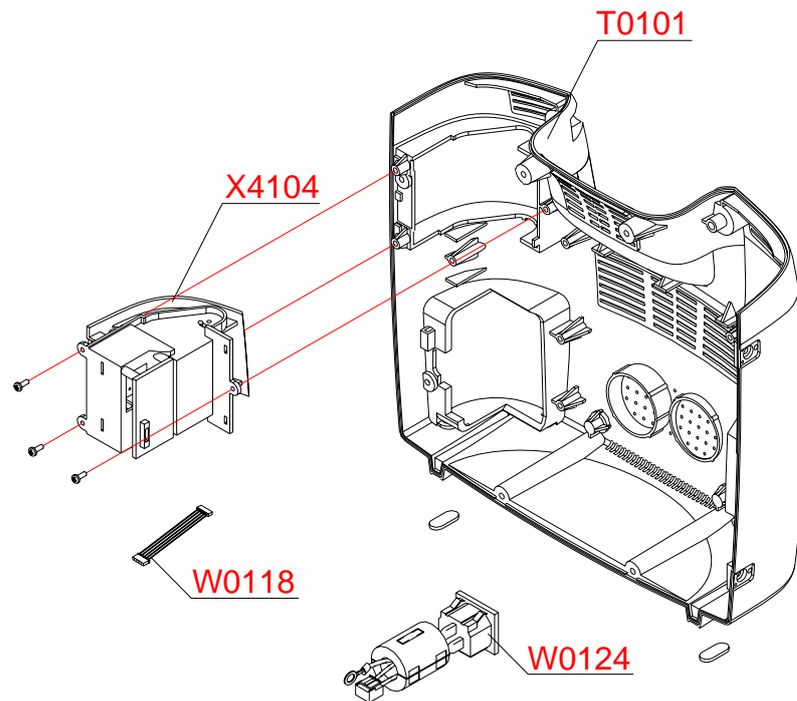


Figure 26. Rear case disassembly – Printer module and AC inlet

Table 28. Part Descriptions – Printer module and AC inlet

Part Codes	Descriptions	Qty
T0101	Rear case	1
X4104	Printer module	1
W0118	Printer wire	1
W0124	AC inlet	1

D4. Printer module disassembly

1. Remove the printer wire from the printer module.
2. Open the printer door to separate the printer module from the rear case.
3. Remove 3 screws (3 × 6) on the printer module.
4. Separate the printer module from the rear case.

D5. AC inlet disassembly

1. Remove the wire connected to the AC inlet.
2. Pull out the AC inlet to separate from the rear case.

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Spare Parts



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

General

Spare parts, along with part numbers, are shown in Table 29. “Item No.” corresponds to the circled callout numbers in Figure 27.

Obtaining Replacement Parts

Mediana Technical Service Department provides technical assistance information and replacement parts. To obtain replacement parts, contact Mediana. Refer to parts by the part names and part numbers.

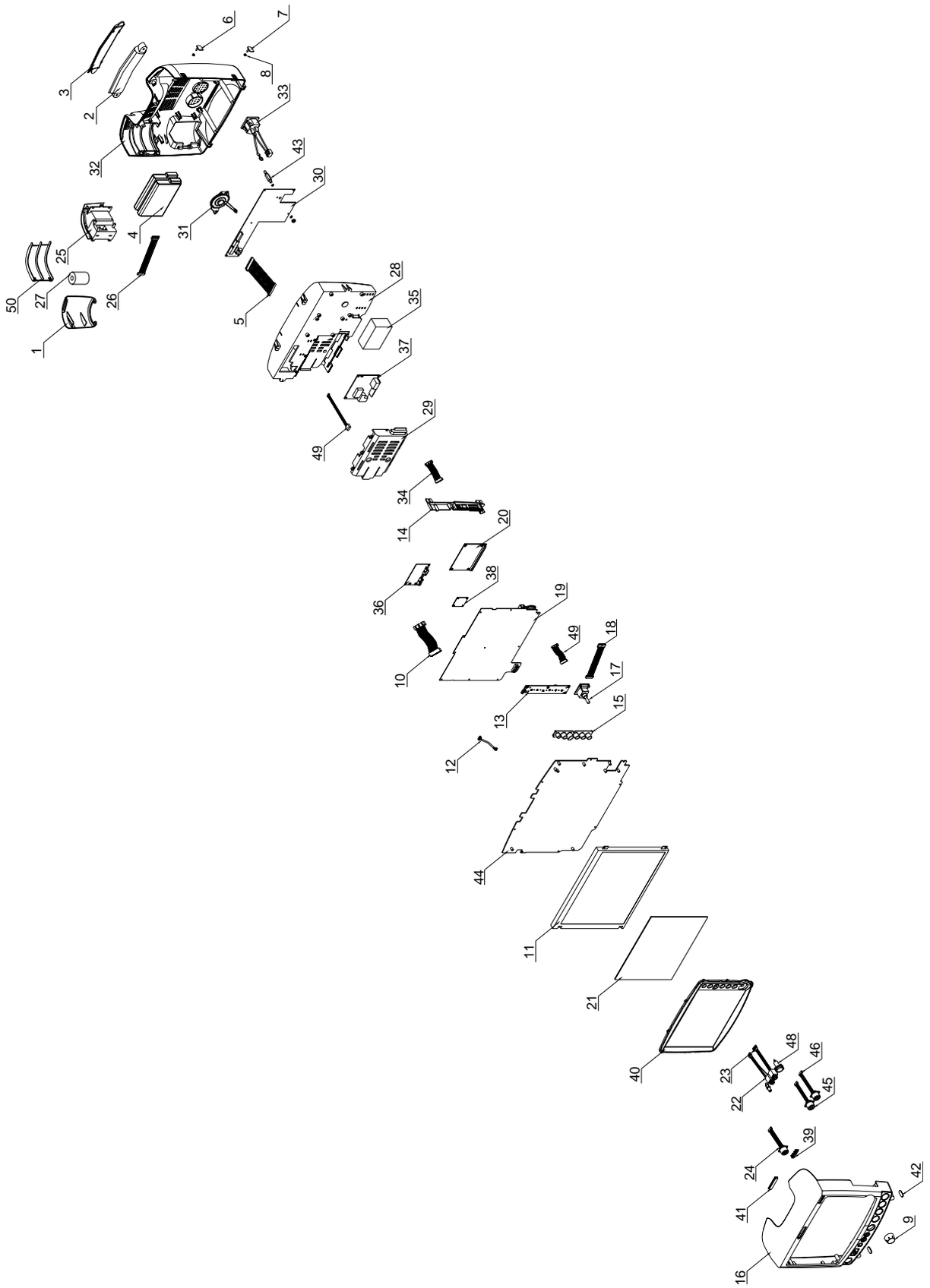


Figure 27. YM6000 Exploded View – Spare Parts

Table 29. Parts List

Item	Part Code	Description
1	T0106	Battery cover
2	T0104	Handle A
3	T0105	Handle B
4	M6010	Battery
5	W0113	SMPS wire / 12 pin
6	T0112	Side cap top
7	T0113	Side cap bottom
8	T0114	Side cap stopper
9	T0100	Knob
10	W0114	Main cable / 24 pin
11	M4022	LCD
12	W0117	LCD cable
13	P1032	Key board
14	T0107	Side connector holder
15	T1009	Rubber key
16	T0096	Front case
17	X1012	Encoder board assembly
18	W0126	Encoder wire
19	P1031	Main board
20	P1033	CPU board
21	T0115	LCD window
22	X1010	Temperature board assembly / 3 pin
23	X1011	Temperature board assembly / 4 pin
24	E4111	ECG connector / 6 pin
25	X4104	Printer module assembly
26	W0118	Printer wire / 6 pin
27	A0062	Printer paper
28	T0102	SMPS case A
29	T0103	SMPS case B
30	M2012	SMPS
31	E9008	Speaker
32	T0101	Rear case
33	W0124	AC inlet wire
34	W0165	NIBP cable / 16 pin
35	M0015	NIBP module
36	M0009	SpO ₂ Module
37	M0013	EtCO ₂ module
38	P1072	IBP board
39	T0098	Alarm LED window
40	T0097	LCD guide
41	T0099	Power LED window

Item	Part Code	Description
42	T1006	Rubber foot
43	T4127	Ground terminal
44	T4112	Bracket
45	W0153	IBP wire / 4 pin
46	W0154	IBP wire / 5 pin
47	W0155	EtCO ₂ wire / 4 pin
48	E4118	Water trap receiver assembly
49	W0115	Key wire / 12 pin
50	T0111	Printer cover

Note: You can use the assembly kit (SS8000-0) supplied by the manufacturer for screws, washers and supporters.

Packing for Shipment

General Instructions

To ship the monitor for any reason, follow the instructions in this section.

Pack the monitor carefully. Failure to follow the instructions in this section may result in loss or damage not covered by the Mediana warranty. If the original shipping carton is not available, use another suitable carton; North American customers may call Mediana Technical Service Department to obtain a shipping carton.

Prior to shipping the monitor, contact your supplier or the Mediana office (Technical Service Department) for a returned goods authorization (RGA) number. Mark the shipping carton and any shipping documents with the returned goods authorization number.

Pack to shipping the monitor, contact your supplier or the Mediana office (Technical Service Department) for a returned goods authorization number. Mark the shipping carton and any shipping documents with the returned goods authorization (RGA) number. Return the monitor by any method that provides proof of delivery.

Returning the YM6000 monitor

Contact Mediana Technical Service Department for shipping instructions, including a Returned Goods Authorization (RGA) number. Unless otherwise instructed by Mediana Technical Service Department, it is not necessary to return the sensor or other accessory items with the monitor. Pack the monitor in its original shipping carton. If the original carton is not available, use a suitable carton with appropriate packing material to protect it during shipping.

Repacking In Original Carton

If available, use the original carton and packing materials. Pack the monitor as follows:

1. Place the monitor and, if necessary, accessory items in original packaging.
2. Place in shipping carton and seal carton with packaging tape.
3. Label carton with shipping address, return address and RGA number, if applicable.

Repacking In a Different Carton

If the original carton is not available, use the following procedure to pack the monitor:

1. Place the monitor in a plastic bag.
2. Locate a corrugated cardboard shipping carton with at least 200 pounds per square inch (psi) bursting strength.
3. Fill the bottom of the carton with at least 2 inches of packing material.
4. Place the bagged unit on the layer of packing material and fill the box completely with packing material.
5. Seal the carton with packing tape.
6. Label the carton with the shipping address, return address, and RGA number, if applicable.

System Processing Description

System Overview

The YM6000 monitor is a multi-function monitor for use on adult, pediatric and neonatal patients; ECG, heart rate, non invasive blood pressures, arterial oxygen saturation, pulse rate, respiration rate and temperature.

In addition to monitoring and displaying the status of these physiological parameters, the instrument performs various microprocessor-programmed analytical functions;

- Creating both visual and audible alarm signals when settable limits are violated;
- Creating and displaying warning/error messages when conditions are detected that would degrade or prevent valid measurements;
- Creating and displaying graphical or tabular trend data;
- Providing input to an optional recorder for printout of current or tabular trend data.

The monitor is essentially a battery-powered instrument. An internal charging unit is designed to accept only an AC line voltage.

System Block Diagram

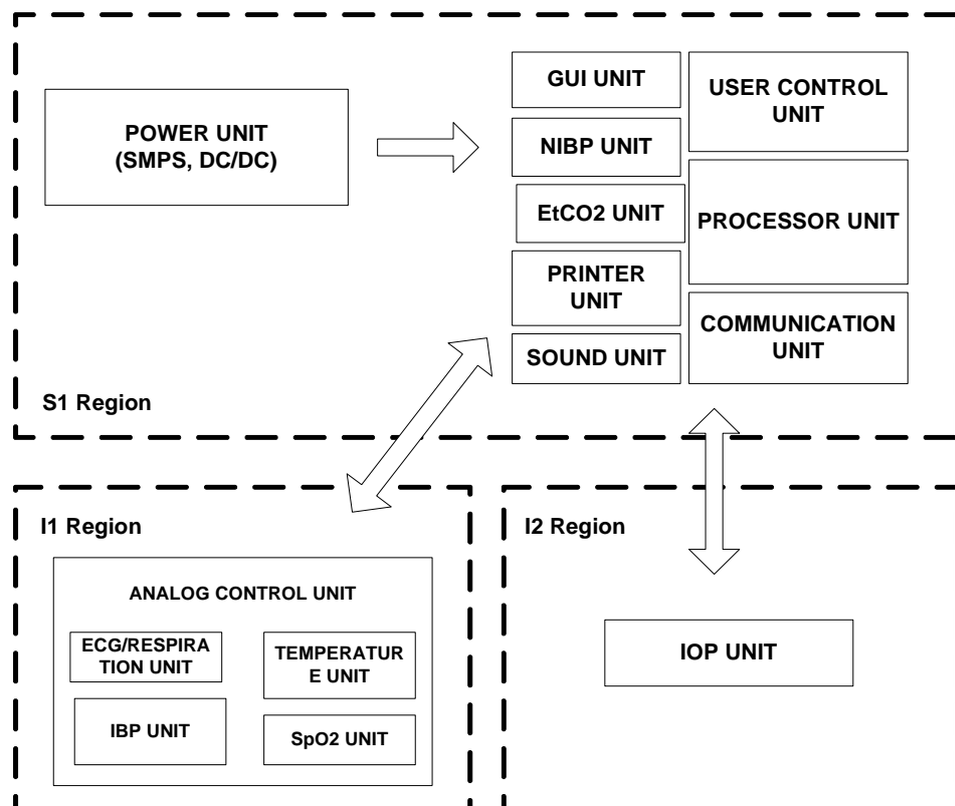


Figure 28. System Block Diagram

Unit Description

- **Power unit:** consists of power entry module, power supply, battery charger, battery, external DC input and DC/DC unit.

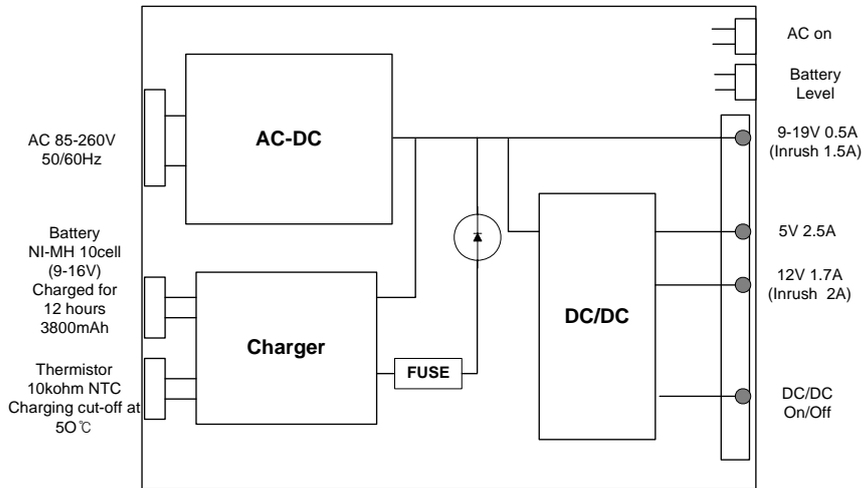


Figure 29. Power Unit Block Diagram

- **Processor unit:** consists of Samsung S3C2440 CPU, SDRSM, Boot ROM and Flash.

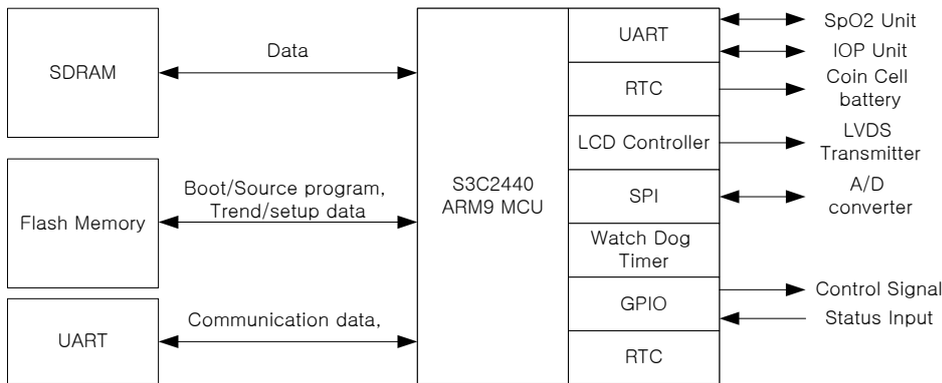


Figure 30. Process Unit Block Diagram

- **User-control unit:** consists of trim knob, 6 functional button, optical encoder, power on indicator LED, AC indicator LED and battery charging status indicator LED.

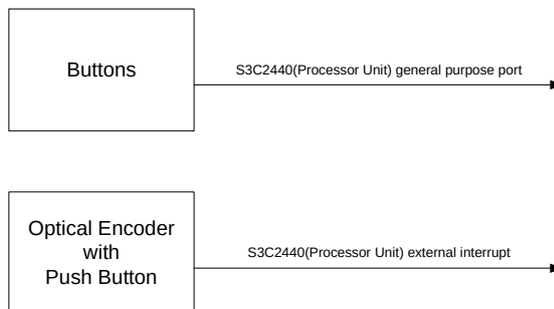


Figure 31. User-Control Unit Block Diagram

- **Sound unit:** consists of 8bit PIC micom, 2-channel amplifiers and speaker.

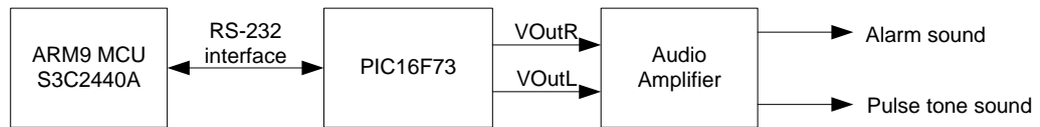


Figure 32. Sound Unit Block Diagram

- **Communication unit :** 4-channel UART

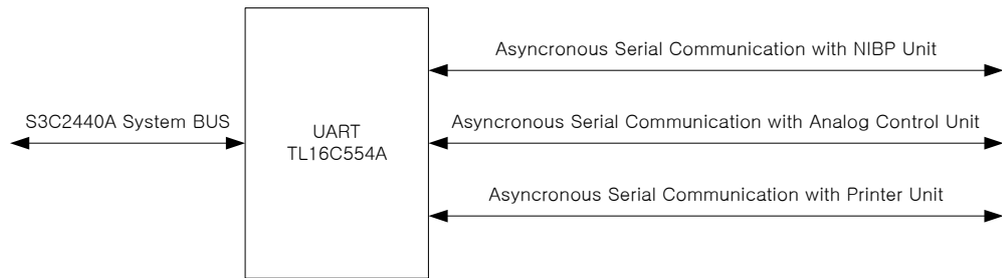


Figure 33. Communication Unit Block Diagram

- **GUI (graphic user interface) unit:** consists of TFT LCD, inverter for backlit and internal video controller.

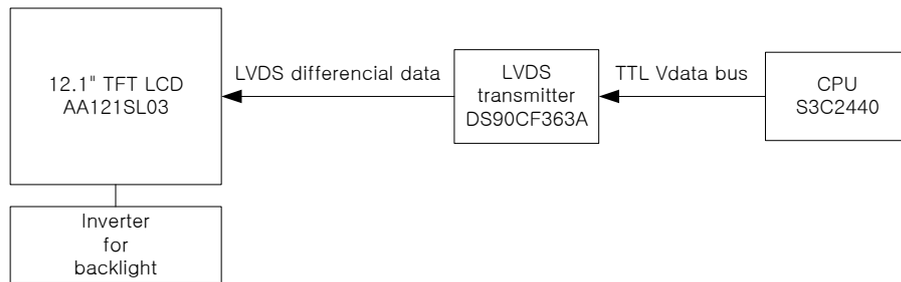


Figure 34. GUI Unit Block Diagram

- **Thermal Printer unit:** prints data records.

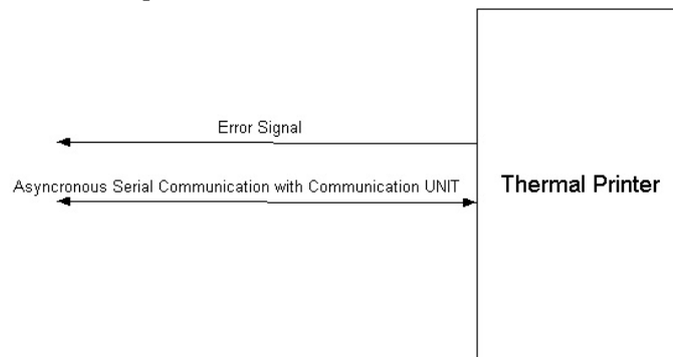


Figure 35. Thermal Printer Unit Block Diagram

- **NIBP unit:** measures non-invasive blood pressure data.

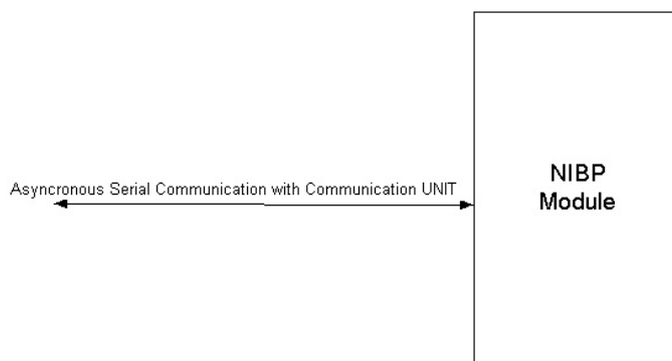


Figure 36. NIBP Unit Block Diagram

- **EtCO₂ unit:** measures EtCO₂ data.

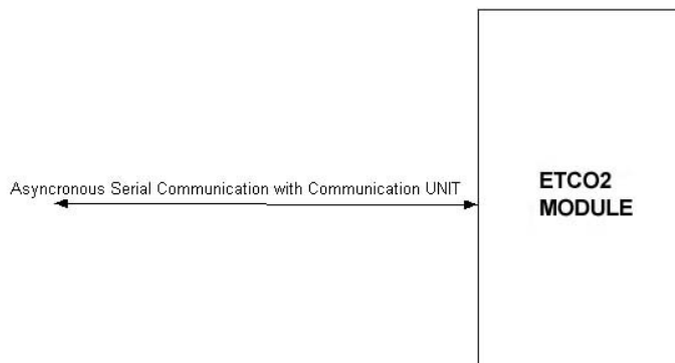


Figure 37. EtCO₂ Unit Block Diagram

- **ECG unit:** measures electrocardiographic waveform data.

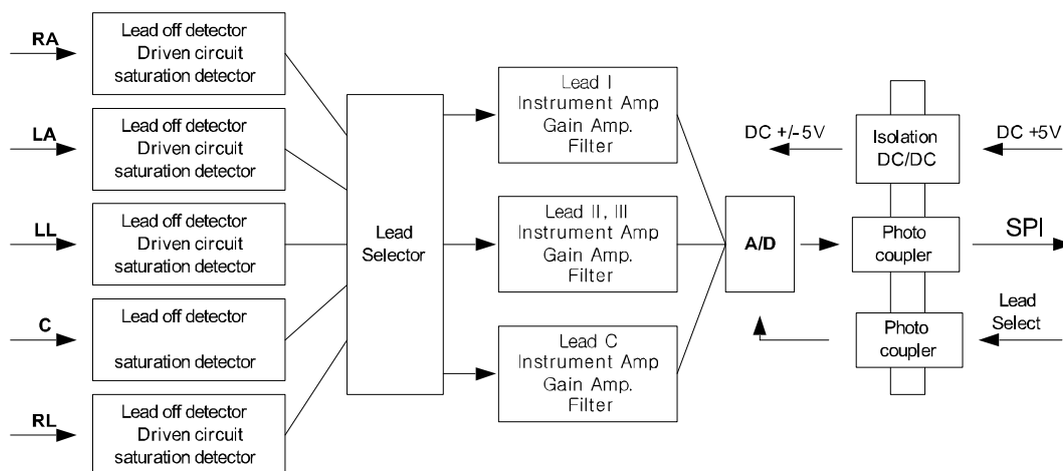


Figure 38. ECG Unit Block Diagram

- **Respiration unit:** measures respiration rate data.

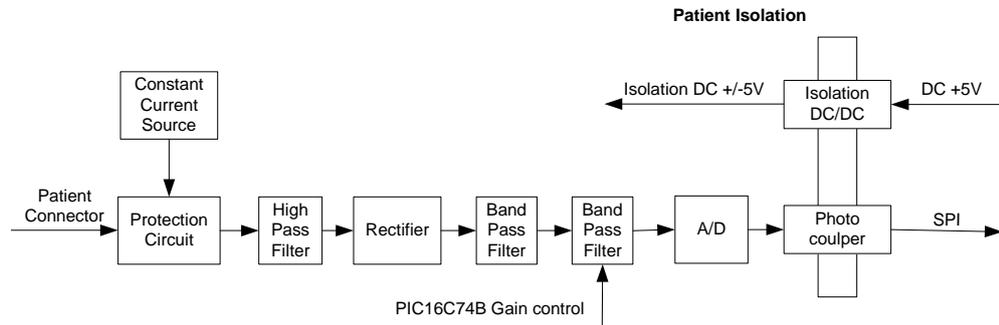


Figure 39. Respiration Unit Block Diagram

- **SpO₂ unit:** measures oxygen saturation data

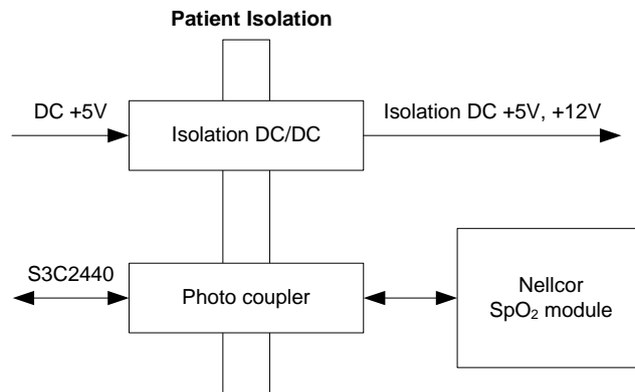


Figure 40. SpO₂ Unit Block Diagram

- **Temperature unit:** measures temperature data.

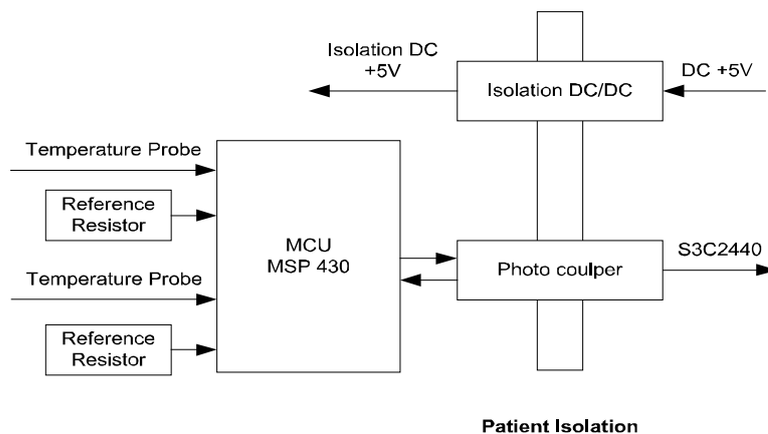


Figure 41. Temperature Unit Block Diagram

- **IBP unit:** measures Invasive blood pressure waveform data.

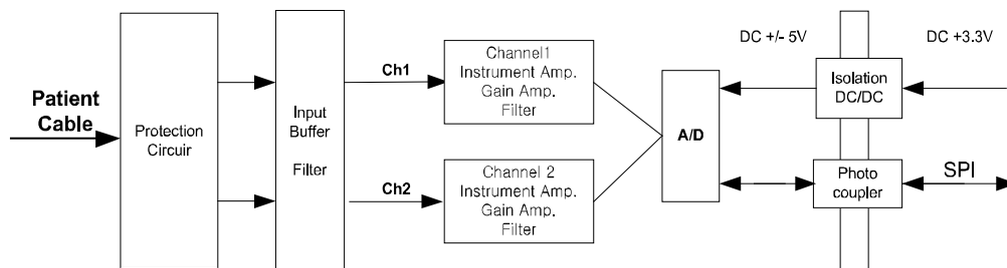


Figure 42. IBP Unit Block Diagram

- **Analog control unit:** analog circuit control, PIC16C74B – selection of ECG channels and filter, size adjustment, verification of lead off, QRS, heart rate, etc.

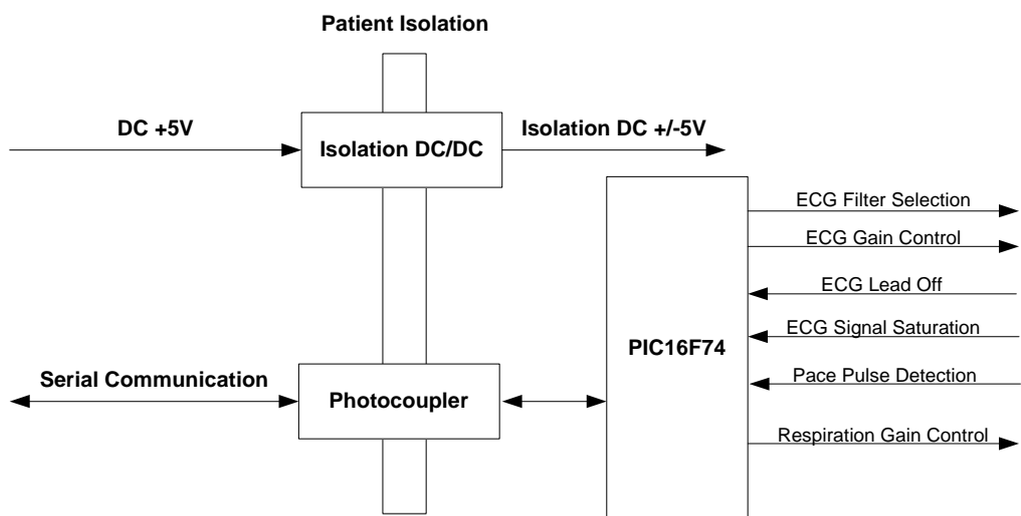


Figure 43. Analog Control Unit Block Diagram

External Interface

- **TCP/IP :** Network connector
- **Serial I/O :** a female 9 pin sub-miniature D shell connector

ECG Processing

The measurement of the skin surfaces electrocardiogram is based on the electrical signals on the skin surface, produced as the heart muscle contracts and relaxes. The signals are detected by electrodes placed on the patient body. The information on heart activity carried by these signals varies with the placing of the electrodes.

The technique used in ECG senses the varying potential difference between two points at the skin surface which respond to the chemical actions of the muscular activity of the heart.

Three electrodes are attached to the patient's right arm (RA), left arm (LA) and left leg (LL). The varying potentials at these locations are cable-connected to the ECG circuit inputs where they are conditioned, and the difference of potential between two selected leads is digitized before transmitting through opto-isolators to the processor. The processor-installed algorithms operate on the signals to develop drivers for the graphic display and to compute the heart rate in beats per minute (bpm).

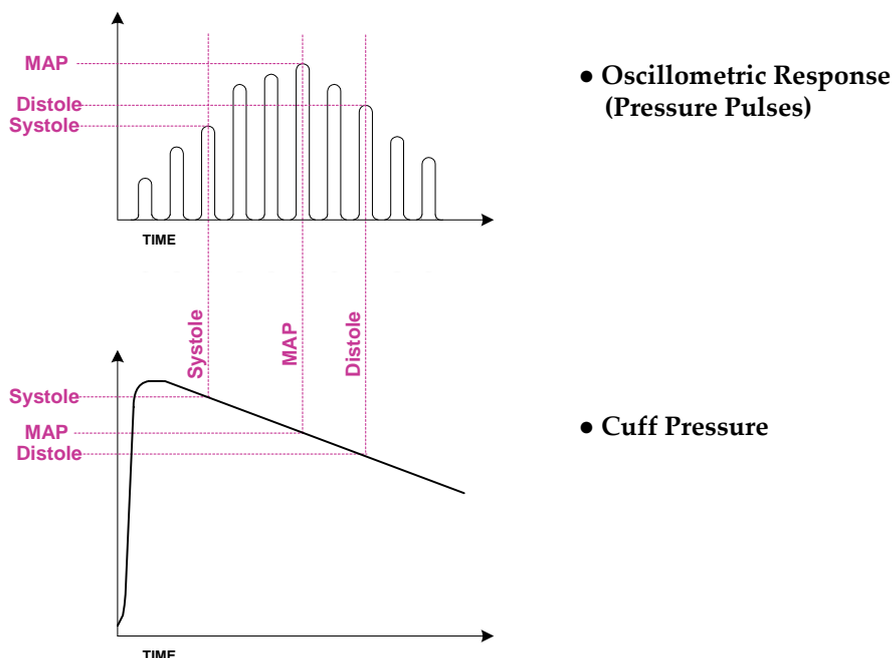
In addition to the acquisition of the QRS waveform complex, the ECG input and subsequent signal processing computing circuitry perform a number of other functions:

- They detect a lead-off condition if one of the electrode connections is disrupted.
- They detect the presence of pacemaker signals within the QRS waveform complex of the ECG.

NIBP Processing

Overview

The oscillometric technique does not use Korotkoff sounds to determine blood pressure. The oscillometric technique monitors the changes in cuff pressure caused by the flow of blood through the artery. The monitor inflates the cuff to a pressure that occludes the artery. Even when the artery is occluded, the pumping of the heart against the artery can cause small pressure pulses in the cuff baseline pressure. The monitor lowers cuff pressure at a controlled rate. As the cuff pressure goes down, blood starts to flow through the artery. The increasing blood flow causes the amplitude of the pressure pulses in the cuff to increase. These pressure pulses continue to increase in amplitude with decreasing cuff pressure until they reach a maximum amplitude at which point they begin to decrease with decreasing cuff pressure. The cuff pressure at which the pulse amplitude is the greatest is known as Mean Arterial Pressure (MAP). The manner in which the pulse amplitudes vary is often referred to as the pulse envelope. The envelope is an imaginary line that connects the peak of each pressure pulse and forms an outline. The shape of the envelope is observed by the monitor using a variety of techniques to determine the diastolic and systolic blood pressure.



Overall Accuracy Discussion

Overall system accuracy shall be determined by considering various influences of the pressure sensor accuracy, motion artifacts, other artifact created by pressure valve, technical errors of electrical components, and the origin error of oscillometric method. The origin error of oscillometric comes from the basic theory of that the MAP is determined by the pulse. Therefore, there might be an error of the time between two pulses. In another words, the greatest amplitude point of pulses could not represent the MAP point exactly.

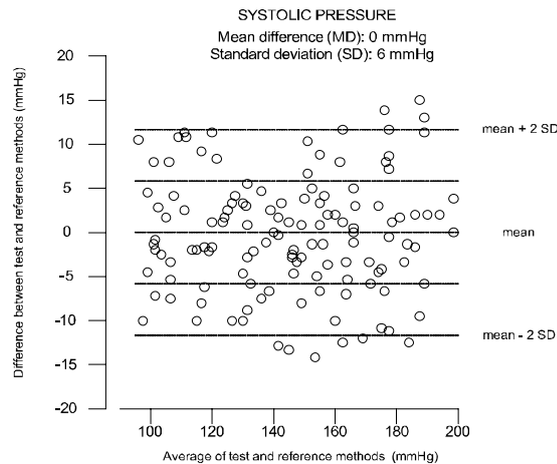
On clinical trial perspective, overall system accuracy is not easy to be determined. The clinical trial test protocols have been tried and have been described in many treatises, and international standards. So, there are many methods to determine the overall system accuracy of Automated Sphygmomanometer using the oscillometric method. But, there are no absolute test protocols to determine the overall system accuracy of the Automated Sphygmomanometer using oscillometric method. Normally, the Gold standards of Blood pressure for the reference are the intra-arterial pressure and the auscultatory method.

The popular standard for the overall system accuracy is AAMI, SP-10 2002+A1;2003 (Electronic or automated sphygmomanometers).

The main test conditions are as follow:

- A. Data comparing the Intra-arterial or the auscultatory by the clinical experts with the automated sphygmomanometer.
- B. For data collection and the data analysis, Bland-Altman Plot is used.
- C. On the systolic, diastolic, and MAP, the Deltas of all measurements shall be met under ± 5 mmHg of mean difference (MD), and ± 8 mmHg of standard deviation (SD).

(Delta = Intra-arterial or Auscultatory – Automated sphygmomanometer)



(EXAMPLE) Agreement between test and reference methods for systolic pressure. Hypothetical data

SpO₂ Processing

Pulse oximetry works by applying a sensor to a pulsating arteriolar vascular bed. The sensor contains a dual light source and photodetector. Bone, tissue, pigmentation, and venous vessels normally absorb a constant amount of light over time. The arteriolar bed normally pulsates and absorbs variable amounts of light during systole and diastole, as blood volume increases and decreases. The ratio of light absorbed at systole and diastole is translated into an oxygen saturation measurement (SpO₂). Because a measurement of SpO₂ is dependent upon light from the sensor, excessive ambient light can interfere with this measurement.

Pulse oximetry is based on two principles: that oxyhemoglobin and deoxyhemoglobin differ in their absorption of red and infrared light (spectrophotometry), and that the volume of arterial blood in tissue (and hence, light absorption by that blood) changes during the pulse (plethysmography). The monitor determines SpO₂ by passing red and infrared light into an arteriolar bed and measuring changes in light absorption during the pulsatile cycle. Red and infrared low-voltage light-emitting diodes (LED) in the oximetry SpO₂ sensor serve as light sources; a photo diode serves as the photo detector. Because oxyhemoglobin and deoxyhemoglobin differ in light absorption, the amount of red and infrared light absorbed by blood is related to hemoglobin oxygen saturation. To identify the oxygen saturation of arterial hemoglobin, the monitor uses the pulsatile nature of arterial flow. During systole, a new pulse of arterial blood enters the vascular bed, and blood volume and light absorption increase. During diastole, blood volume and light absorption reach their lowest point. The monitor bases its SpO₂ measurements on the difference between maximum and minimum absorption (measurements at systole and diastole). By doing so, it focuses on light absorption by pulsatile arterial blood, eliminating the effects of nonpulsatile absorbers such as tissue, bone, and venous blood.

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 SpO₂ sensor's red LED to accurately measure SpO₂. 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.

Measured versus Calculated Saturation

The measured SpO₂ value from an oximeter may differ from the saturation value that is calculated from a blood gas partial pressure of oxygen (PO₂). This usually occurs because the calculated saturation was not appropriately corrected for the effects of variables that shift the relationship between PO₂ and saturation: pH, temperature, partial pressure of carbon dioxide (PCO₂), 2, 3-DPG, and fetal hemoglobin.

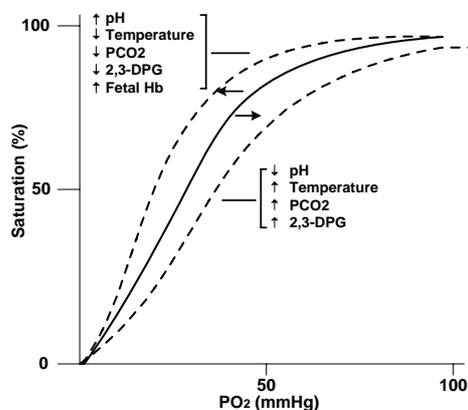


Figure 44. Oxyhemoglobin Dissociation Curve

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:

$$\text{functional saturation} = \frac{\text{fractional saturation}}{100 - (\% \text{carboxyhemoglobin} + \% \text{methemoglobin})} \times 100$$

SpO₂ Accuracy

The saturation (SpO₂) accuracy specification and/or pulse rate (PR) performance were analyzed by comparative oximetry performance (COPS) tests between YM6000 SpO₂ module and MP506 module with the same version of the oximetry algorithm from Nellcor. This was to demonstrate that the performance of YM6000 SpO₂ module was equivalent to that of MP506, which had been validated during both standard motion, combined motion and cold-induced peripheral vasoconstriction (low perfusion) conditions by direct comparison to measurements of arterial oxygen saturation (SpO₂) obtained from arterial blood samples analyzed with Instrumentation Laboratory (IL) CO-Oximeters under Nellcor's clinical protocol, Invasive Controlled Hypoxia Studies.

Respiration Processing

The respiration monitoring is designed to use the variation of this thoracic impedance. The chest contains various materials, ranging from bone to air. Each of these materials has different electrical properties and is located in a different portion of the chest. The materials of the chest vary in electrical resistivity (the amount of electrical resistance between opposite faces of a cube of that material), which is an important determinant of electrical impedance in the body.

Two of the major components of the chest, blood and air, are at opposite ends of the scale. Furthermore, the volume of each of these materials varies with time over the cardiac and breathing cycles. The variation of the thoracic impedance is caused by the difference between air and blood in the thoracic impedance. Blood has relatively low resistivity, which varies over the cardiac cycle owing to changing blood volumes in the heart and in the vascular compartment. Air, on the other hand, has high electrical resistivity and hence impedance, and it undergoes wide volume changes in the lungs during normal breathing. i.e. the impedance of blood is 150 ohm/cm and the one of air is 5000 ohm/cm.

The patient's respiration is detected by using two of the three leads of the ECG electrodes (RA and LA, or RA and LL) and cable. The electrical impedance between a pair of electrodes is determined by dividing the voltage difference between the two electrodes by the current that passes between them. When the electrodes are placed on the actual structure, respective structures change.

A low-level excitation signal is applied to these leads, and the variation of the thoracic impedance caused by the breathing is sensed and processed for display and measurement. This variation is processed to the voltage value for the measurement. In order to transfer the thoracic impedance by a transformer, it is used a minimum constant current of the sine wave carrier signal. The transferred thoracic impedance is changed to the voltage signal by using bridge circuit and differential amplifier. Then, ECG signal is removed by filter, and carrier frequency is removed by full wave rectifier and filter in order to extract only thoracic impedance in amplifying at the definite level of signal. This extracted thoracic impedance signal is used to measure the respiration by digital signal processing.

Temperature Processing

Measurement of patient temperature is accomplished by processing the signal from a probe containing a resistor whose resistance is temperature dependent. The class of such components is called thermistor.

Temperature measurement used by the YM6000 monitor based on a thermistor whose resistance is inversely proportional to its temperature. By measuring the thermistor's resistance, its temperature can be calculated. The resistance of the thermistor is measured by passing a current through it and measuring the voltage developed across it.

The YM6000 monitor is designed to accept the signals from electrically isolated a range of temperature probes from YSI-400 series. The probes may be used for skin or rectal temperature measurement. Probes are furnished with a standard 10-foot lead; extension leads are available. The signal from the probe is conditioned by the monitor input circuitry, processed, and used to drive the numeric display.

Invasive Blood Pressure Processing

The pressure transducer is connected to a pressure line which, by means of a catheter is invasively connected to the patient blood stream. The force of movement of the blood in the patient vessels is transported by the fluid column in the pressure line to the transducer. This movement causes an electrical signal to be generated which is then amplified to display the pressure wave and the numeric for the systolic, diastolic and mean pressure values.

The blood pressure is influenced by the respiratory system. This occurs in spontaneous breathing patients, but is more apparent in positive pressure ventilated patients. To reduce this respiration artifact the module uses a variable weight filter technique in the processing of the pressure values.

Capnography Processing

The capnography parameter provides the continuous, non-invasive monitoring of sidestream end-tidal carbon dioxide (EtCO₂), inspired carbon dioxide (InCO₂), and respiration rate (RR). The measured values for capnography (EtCO₂, InCO₂ and RR) are displayed in the CO₂ parameter box and a CO₂ waveform can be continuously displayed. Nitrous oxide compensation is selectable.