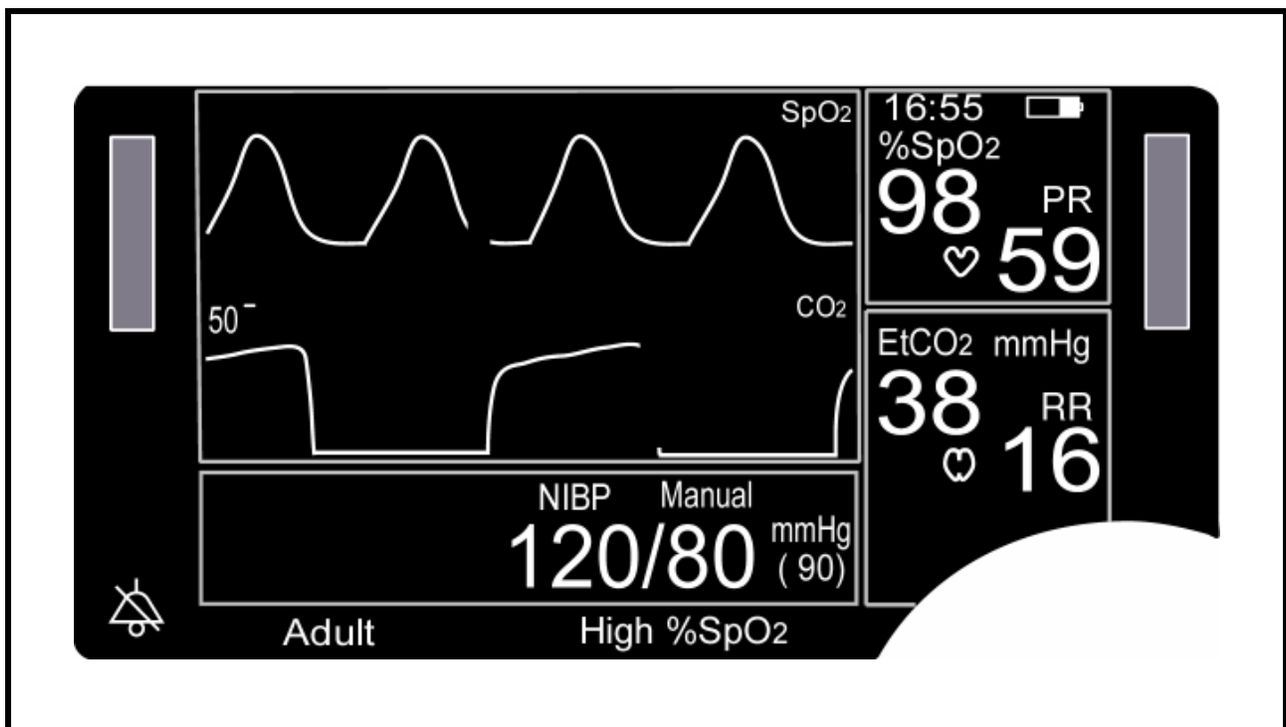

CAS 750C

Multi-Parameter Monitor



User's Manual

CE 0086

 **CAS** MEDICAL SYSTEMS, INC.
TECHNOLOGY APPLIED TO MEDICINE

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Please contact the distributor in the country of purchase if product information or service should be required.

General Information



This manual is an integral part of the product and describes its intended use. Observance of the manual is a prerequisite for proper product performance and correct operation and ensures patient and operator safety.

The symbol means ATTENTION: Consult accompanying documents.

The warranty does not cover damages resulting from the use of accessories and consumables from other manufacturers.

CAS Medical Systems, Inc. is responsible for the effects on safety, reliability, and performance of the product, only if:

- Assembly, operations, extensions, readjustments, modifications, or repairs are carried out by persons authorized by CAS Medical Systems, Inc.
- The electrical installation of the relevant room complies with the requirements of the appropriate regulations.
- The device is used in accordance with the instructions for use.
- All publications conform to the product specifications and applicable IEC publications on safety and essential performance of electro medical equipment as well as with applicable UL requirements and AHA recommendations valid at the time of printing.

For complete Warranty information, refer to the Warranty Policy located on page 110.

The CAS Medical System, Inc. quality management system complies with the international standards ISO 13485 and the Council Directive on Medical Devices 93/42/EEC.

Note: The information in this manual applies only to the CAS 750C software version 2.2. It does not apply to earlier software versions.

Note: Due to continuing product innovation, specifications in this manual are subject to change without notice.

In the U.S. the following caution applies:



Caution: Federal law restricts this device to sale by or on the order of a physician or properly licensed practitioner.



Warning: Before using the monitor for the first time, please read the information given in section "Safety" starting on page 13.

About this Manual

Note: This manual addresses all parameters a CAS 750C Monitor can have installed. It remains suitable for use if the monitor has a sub-set of parameters only. Please refer to those sections that are applicable for the model in use.

Manual Purpose

This manual contains the instructions necessary to operate the CAS 750C monitor safely and in accordance with its functions and intended use.

Intended Audience

This manual is written for clinical professionals. Clinical professionals are expected to have working knowledge of medical procedures, practices, and terminology as required for monitoring of critically ill patients.



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

Conventions



Warning: Directions that warn of conditions that put the patient or the caregiver at risk.



Caution: Directions that help to avoid damaging the monitor or losing data.

Note: Directions that make it easier to use the monitor, something not readily apparent.

Figures: All illustrations in this manual are provided as examples only. They may not necessarily reflect your monitoring setup or data displayed on your monitor.

Revision History

This manual has a revision number located at the bottom of each page. It changes whenever the manual is updated.
First Printing: 08/2005.

Read this manual carefully before patient use of the monitor.

CAS Medical Systems, Inc. reserves the right to make changes to this manual and improvements to the product it describes at any time without notice or obligation.

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1 Safety

The following Warnings, Cautions and Notes have to be obeyed to guaranty a safe operation of the monitor. Additional Warnings, Cautions and Notes, which apply to specific parameters, are listed in the related sections for each parameter.

Indications for Use

The 750C monitor is intended to continuously calculate and display the following physiological vital signs: end tidal carbon dioxide, respiration rate, capnograph waveform, functional arterial oxygen saturation, pulse rate, plethysmograph waveform and an optional non-invasive blood pressure measurement of systolic, diastolic, mean arterial pressures along with pulse rate derived from a NIBP pressure waveform.

The 750C is intended for monitoring of adult, pediatric and neonatal patients in the care of health care professionals.

Contraindications

- Reusable SpO₂ sensors are contraindicated for use for prolonged periods of use. It is not intended for long term monitoring. It must be removed and repositioned every four (4) hours and if indicated by circulatory condition or skin integrity, reapplied to a different monitoring site.
- Disposable SpO₂ sensors are contraindicated for patients that exhibit allergic reactions to adhesive tape. The sensors must be removed and repositioned every eight (8) hours and if indicated by circulatory condition or skin integrity, reapplied to a different monitoring site.
- No other contraindications are known at this time.

Installation and Setup

Follow the instructions given in paragraph “Site Requirements” on page 25 and “Power Requirements” on page 25.



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



Warning: The CAS 750C Monitor is intended only as an adjunct in patient assessment. It must be used in conjunction with clinical signs and symptoms.



Warning: Do not rely exclusively on the audible alarm system for patient monitoring. Adjustment of alarm volume to a low level during patient monitoring may result in a hazard to the patient. Remember that the most reliable method of patient monitoring combines close personal surveillance with correct operation of monitoring equipment.



Warning: Do not use the CAS 750C Monitor for any purpose other than specified in this manual. Doing so will invalidate the monitor’s warranty.



Warning: Do not connect the CAS 750C Monitor to more than one patient.



Warning: The CAS 750C Monitor is not intended to be used in Oxygen Enriched Atmospheres.



Warning: The CAS 750C Monitor is defibrillator proof. It may remain attached to the patient during defib., but the readings may be inaccurate during use and less than ten (10) seconds thereafter.



Warning: Do not use the monitor in the presence of Magnetic Resonance Imaging (MRI) equipment.



Warning: The CAS 750C Monitor is not “Category AP or APG Equipment”.



Warning: *Explosion Hazard* - Do not use the monitor in the presence of a flammable Anesthetic Mixture with Air or with Oxygen or Nitrous Oxide.



Warning: *Explosion Hazard* - Do not use this equipment in the presence of flammable anesthetics, vapors or liquids.



Warning: *Electromagnetic Compatibility (EMC)* - The equipment needs special precautions if it is placed close to a strong transmitter such as X-ray equipment, MRI devices, TV, AM/FM radios, police/ fire stations, a HAM radio operator, an airport, or cellular phone. Their signals could interfere with the monitor, which may result in disruption of performance of this device or prevents the clear reception of signals by the monitor.



Caution: Qualified biomedical engineering personnel only must interface monitoring equipment with other types of medical equipment. Be certain to consult manufacturers' specifications to maintain safe operation.



Caution: *Leakage Current Test* - The interconnection of auxiliary equipment with this device may increase the total leakage current. When interfacing with other equipment, qualified biomedical engineering personnel must perform a test for leakage current before using it with patients. Serious injury or death could result if the leakage current exceeds applicable standards.



Caution: Measurements may be affected in the presence of strong electromagnetic sources such as electro surgery equipment.

Note: The CAS 750C Monitor is designed for continuous operation.

Note: The CAS 750C Monitor is suitable for use in the presence of electro surgery.

Note: The CAS 750C Monitor can remain connected to the patient during Cardio Defibrillation. All applied parts are "Type BF Defibrillation Proof".

The monitor has been designed to promote patient safety. All equipment parts are protected against the effects of the discharge of a defibrillator. No separate actions are required when using this equipment with a defibrillator.

Device Handling



Warning: To ensure patient safety, do not place the monitor in any position that might cause it to fall on the patient.



Warning: Do not lift the monitor by any sensor cable or line as they could disconnect from the monitor, causing the monitor to fall on the patient.



Warning: To avoid electric shock or device malfunction, liquids must not be allowed to enter the device. If liquids have entered a device, take it out of service and have it checked by a service technician before it is used again.



Warning: The CAS 750C Monitor provides “DRIP-PROOF” level of protection from ingress to moisture.



Warning: Do not place liquids on top of the monitor. Do not immerse the monitor or power cord in water or any liquid.



Warning: Do not gas sterilize or autoclave the monitor.



Warning: Do not touch part of non-medical electrical equipment in the patient environment after removal of covers, connectors etc... without the use of a tool which operate at voltages not exceeding 25 VAC or 60 VDC and the patient at the same time.



Warning: Where the integrity of the external protective conductor in the installation or its arrangement is in doubt, EQUIPMENT shall be operated from its INTERNAL ELECTRICAL POWER SOURCE.



Warning: Isolation of product from mains can only be achieved by removal of external power cord.



Warning: Route all cables away from patient’s throat to avoid possible strangulation.



Caution: Pressing the front panel keys with a sharp or pointed instrument may permanently damage the switch membrane. Press the keys using only your finger.



Caution: If the monitor is accidentally wetted, take it out of operation. It should be thoroughly dried. To verify the absence of water, a qualified service technician can remove the rear cover.



Caution: To avoid the risk of electrical shock, do not remove the back cover. Refer all servicing to qualified personnel.

Note: There are no known risks with common disposal of equipment or accessories; however, the disposing of accessories should follow in accordance with local hospital policies. The user should ensure these policies do not conflict with any local, state or federal guidelines.

Safety Checks



Warning: Do not, under any circumstances, perform any testing or maintenance on the monitor or power cord while the unit is being used to monitor a patient. Unplug the power cord before cleaning or servicing the monitor. The operator should not perform any servicing except as specifically stated in this manual.



Warning: The functions of the alarm system for monitoring of the patient must be verified at regular intervals.



Warning: Periodically, and whenever the integrity of the product is in doubt, test all functions.



Warning: Do not use a frayed or damaged power supply cord or any accessory if you notice any sign of damage. Contact CAS Medical Systems for assistance.



Caution: Inspect the monitor, patient cables, sensors and air hose for damage prior to operation. If any damage is noted, the monitor should not be used until it has been serviced. Only personnel authorized to do so by CAS Medical Systems, Inc. should repair the monitor.



Caution: If the monitor fails to respond, do not use it until the situation has been corrected by qualified personnel.

Monitoring



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



Warning: If the accuracy of any value displayed on the monitor or printed on a graph strip is questionable, determine the patient's vital signs by alternative means. Verify that all equipment is working correctly.



Caution: Use only CAS approved accessories and sensors to ensure patient safety and to preserve the integrity, accuracy and the electromagnetic compatibility of the monitor.

Initial Inspection

Before unpacking the monitor, inspect the packaging for damage. If there are any signs of damage to the package, a claim should be filed immediately with the shipping agent. It is the receiver's responsibility to notify the carrier's local office to arrange for the pickup of the damaged items. Save the damaged shipping carton as evidence.

Contact your distributor, CAS sales representative, or call CAS Medical Systems, Inc. to report external damage and to arrange for repair or replacement of damaged equipment.

The shipping carton should contain the items listed below. Unpack the monitor and account for each item. Inspect each item for signs of external damage, dents, cracks, scratches, etc. If an item is missing or damaged, contact your distributor, CAS sales representative, or CAS Medical Systems, Inc.

Record the monitor model, serial number and date of purchase at the back of this manual.

Monitor Checklist

- (1) CAS 750C Monitor
- (1) Hospital Grade AC Power Cord or DC Power Cord
Depending on model ordered
- (1) SpO₂ Interconnect Cable
- (1) SpO₂ Finger Sensor
- (2) FilterLine Set, Adult/Pediatric
- (2) Smart CapnoLine, Adult
- (1) Ten (10) Foot Coiled Inflation Hose
For models with NIBP installed
- (1) Tuff-Cuff® Blood Pressure Cuffs, Adult
For models with NIBP installed
- (1) Tuff-Cuff® Blood Pressure Cuffs, Child
For models with NIBP installed
- (1) P9 Calibration Kit
For models with NIBP installed
- (1) CAS 750C Monitor User's Manual

Note: The monitor is shipped with the appropriate line cord for the country and or voltage being used.

Patient Environment

The CAS 750C Monitor has been tested with specific parts of the “system” used within the Patient Environment. These parts are:

- The CAS 750C Monitor
- Appropriate Accessories as listed in section “Accessories” at the back of this manual.
- Line Cord
- Citizen CMP-10 Mobile Printer
- RS232 Interconnect Cable (supplied with printer)
- AC Adapter / Charger, Model TRC-09-1100-M from Group West or equivalent (supplied with printer)

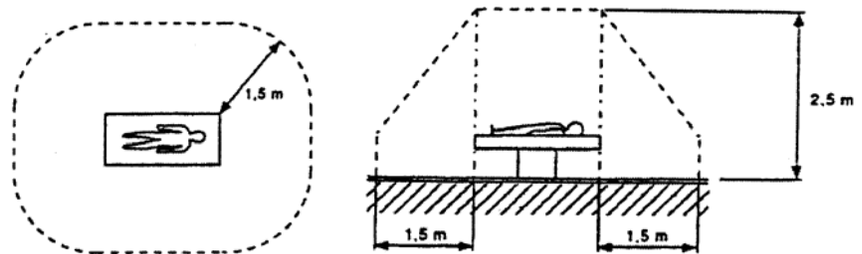


Figure 1: Patient Environment

Monitor Electronics Classifications

The **CAS 750C** Monitor (monitor version with integrated AC Power Supply) is a Class I device.

The **CAS 750CM** Monitor (monitor version with external DC Power Supply) is a Class II device.

2 Basic Operations

Introduction

The CAS 750C Monitor is a pre-configured monitor that can include the following measurement functions:

- Capnography (CO₂)
- Pulse Oximetry (SpO₂)
- Non Invasive Blood Pressure (NIBP)

The MicroStream Capnography provides continuous monitoring of the EtCO₂ value and the Respiration Rate (RR) of intubated and non-intubated patients. The CO₂ signal can be displayed as a waveform.

The Pulse Oximeter function continuously monitors and displays values for functional arterial hemoglobin saturation (%SpO₂) and the pulse rate (PR). The dynamic pulse signal can be displayed as a waveform (Plethysmogram) or as a bar graph indicator.

The non-invasive blood pressure measurement uses the oscillometric method and applies for neonatal, pediatric and adult patients. The user can select between manual, cycle or continuous (STAT) mode.

The monitor is equipped with a rechargeable battery pack and can be used independently from an external power source.

Front View

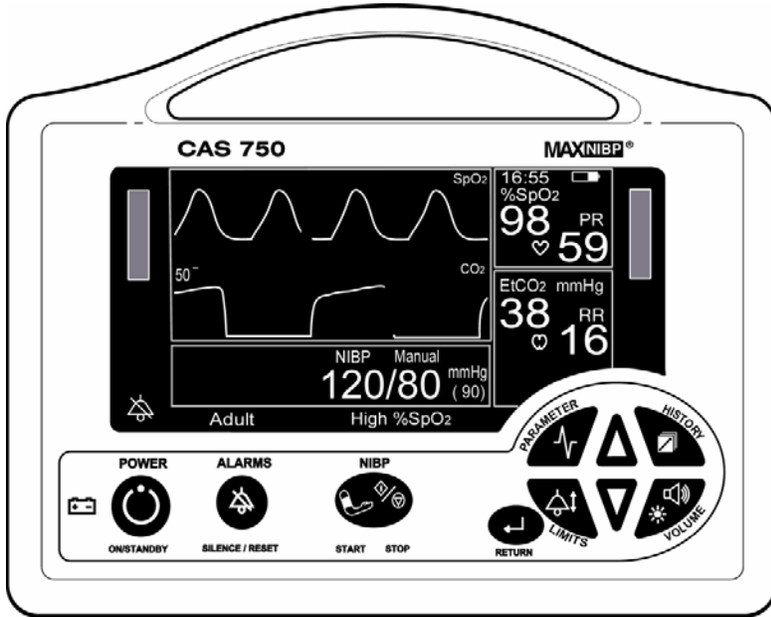


Figure 2: Front View

Rear View

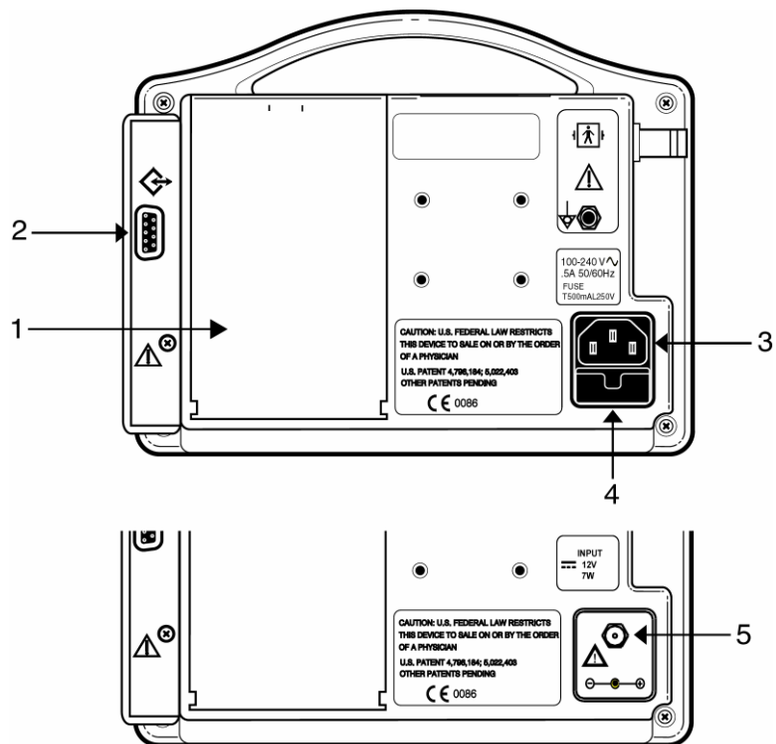


Figure 3: Rear View of Monitors

(M-Version shown below)

1. Battery Compartment.
2. External Device Interface (RS232)
3. Receptacle for the AC power cord
4. Fuse Compartment
5. Receptacle for the 12 VDC cable

Bottom View

Infrared Communication Port

An Infrared (Ir) output port, located on the bottom panel of the monitor's front cover, is available to print Waveforms, History and Alarm data to the optional external printer or other data collection device(s). Maximum distance is approximately 3 feet, direct line of sight operation.

Serial Number Label

The serial number label is located on the bottom of the monitor.

Getting Started

Site Requirements

Note: Put the monitor in a location where you can *easily* see the screen and access the operating controls.



Caution: Stacking - Where monitor is used adjacent to or stacked with other equipment, the monitor should be observed to verify normal operation in the configuration in which it will be used.



Caution: Negligence - CAS Medical Systems Inc. does not assume responsibility for damage to the equipment caused by improperly vented cabinets, improper or faulty power, or insufficient wall strength to support equipment mounted on such walls.

Power Requirements

The external power for the CAS 750C Monitor can be either AC mains power or DC power. The two different models are:

- Model **CAS 750CM** for +12V VDC
- Model **CAS 750C** for 110 to 240 VAC

The following applies for the CAS 750C model with AC power supply:



Warning: Do not plug the monitor into an outlet controlled by a wall switch or dimmer.



Warning: Where the integrity of the external protective conductor in the installation or its arrangement is in doubt, EQUIPMENT shall be operated from its INTERNAL ELECTRICAL POWER SOURCE.



Warning: Isolation of product from mains can only be achieved by removal of external power cord.



Caution: Do not defeat the three-wire grounding feature of the power cord by means of adaptors, plug modifications, or other methods. Do not use extension cords of any type.

Note: The monitor is suitable to be connected to public AC mains power.

Turning the Monitor On

Press the POWER key on the front panel to turn the monitor on.

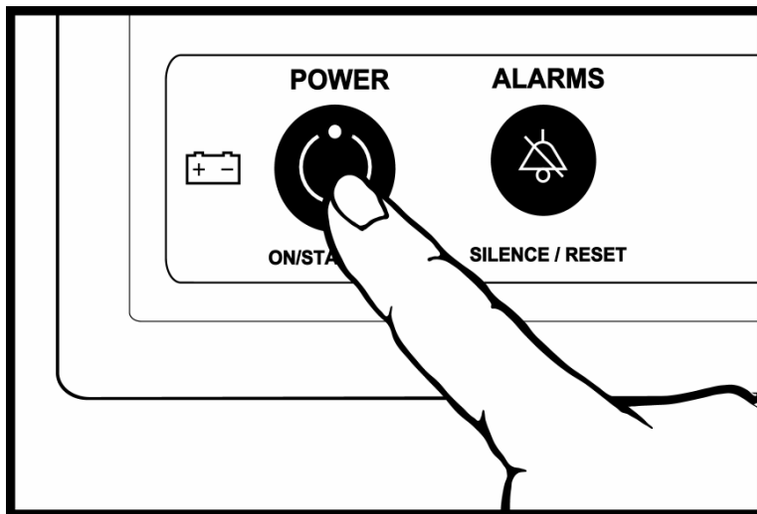


Figure 4: Turning the Monitor On

Each time the monitor is turned on an internal self-test is conducted to ensure that all circuits are functioning properly. Verify the following:

- The monitor produces 3 one-second audio beeps.
- Both, the Equipment and the Alarm Indicators are lit for one-half second and cycled one after the other.
- The monitor displays temporarily its current configuration data:
 - Model number (750C)
 - Power source (C=External AC, CM=External DC)
 - Number of installed parameters
 - Brand of installed parameters (MS = Masimo, NL = Nellcor, C = Oridion)



Warning: The monitor must be briefly turned off and back on once every 24 hours of continuous operation to exercise the start-up self-test function of the optional NIBP module.

Note: If after the Power-On-Self-Test is completed the monitor displays “NIBP Cal” in the Equipment Message area, the monitor should be returned to CAS Medical Systems for service.

Once the Power-On-Self-Test is completed, the monitor’s Main screen is activated and the monitor is ready for use.

Turning the Monitor Off

When the monitor is not being used, press the POWER key can turn it off. The display will turn blank and the unit is no longer monitoring the patient.

Note: The internal power supply is not switched off! It remains connected to the external power source (AC or DC) to enable battery charging.

Power Fail Message

During the internal self-test the monitor checks whether it was previously turned off correctly or if it was disconnected from power in any other way. The user will be alerted about such a power loss after the monitor is turned on again.

The message "Power Failure" is displayed in the Equipment Message Window, the Equipment Alarm LEDs are activated and 3 beeps followed by 2 beeps are heard every 10 seconds.

- Press the SILENCE/RESET key to clear the Power Fail condition.

Note: Only the SILENCE/RESET and the ON/STANDBY keys are operable, all other keys are disabled while the message is displayed.

Battery Operation

The monitor is equipped with a rechargeable battery. The battery is charging whenever the monitor is connected to an external power source (AC Line Power or +12 VDC).

Batteries will self-discharge when they are not used. It is recommended leaving the monitor connected to an external power source whenever possible.

The monitor will operate on a completely charged battery for 3 to 5 hours depending of its configuration and the use of the NIBP function.

Battery Status

There are several means to get an indication about the battery charge status:

- When the monitor is connected to external power and turned off, the Main screen displays the Battery Indicator icon with a moving bar from left to right within the indicator signifying the battery is being charged. Once charged, the moving bar will stop and the battery icon will be completely filled in.
- When the monitor is being powered from the battery, the moving bar within the Battery Indicator icon will be moving from right to left signifying the battery is being discharged.



Caution: Under various state and local laws, it may be illegal to dispose of the battery into the municipal waste stream. Check with your local authorities for instructions on recycling options in your area.

Battery Conditions

The user will be alerted in case the battery charge level gets low:

- The "Low Battery" or "Dead Battery" message is displayed.
- The Power Indicator changes from orange to red.
- The Equipment Alarm LEDs are activated.
- An audible tone is generated



Warning: After approx. 3 minutes in "Dead Battery" condition, the monitor turns off and the patient is no longer being monitored.



Warning: If the battery is not charged, the monitor may no longer function as intended.

Note: When the "Low Battery" or "Dead Battery" message appears, the monitor should be connected to an external power source. A depleted battery may be fully recharged in 5 hours.

Note: The monitor can remain in normal operation while the battery is charging. During charging, the case may feel warm to the touch.

Connecting the Accessories

Connect the various accessories to the appropriate input connector at the left side of the monitor.

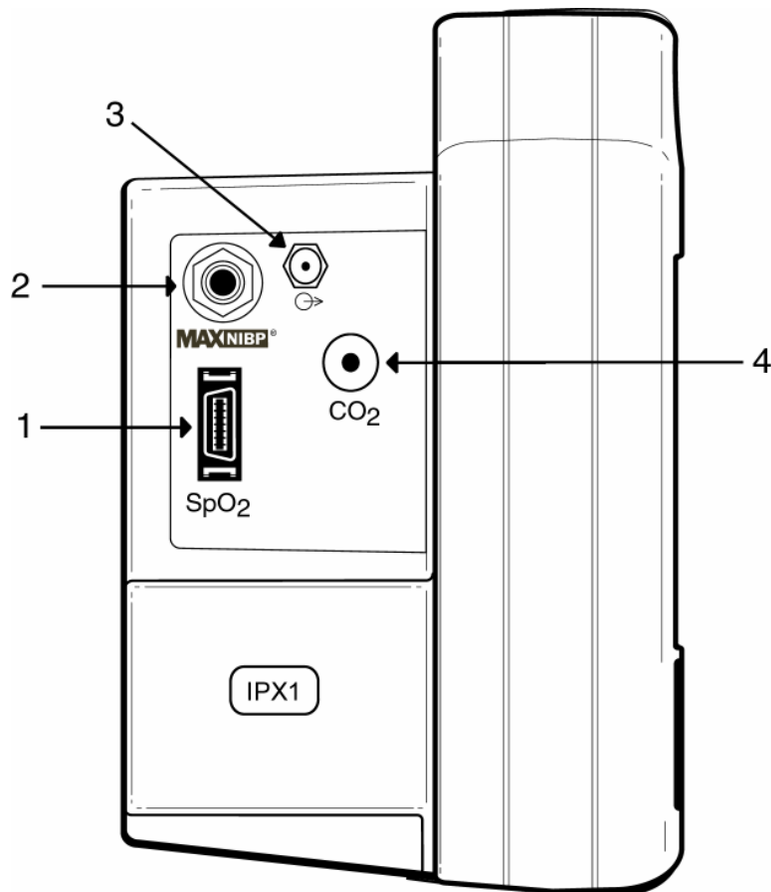


Figure 5: Left Side View

1. SpO₂ Probe Connector
2. NIBP Hose Connection
3. CO₂ Scavenger Exhaust
4. MicroStream™ CO₂ Input Connector

Front Panel Controls

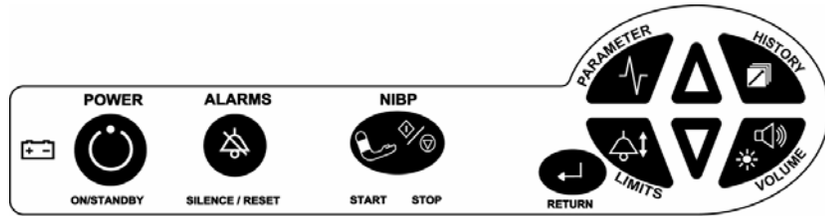


Figure 6: Front Panel Controls



Power Indicator.

- Green:* Monitor operates on external power.
- Yellow:* Monitor operates on battery power.
- Red:* Low or dead battery.



POWER

- On:* Turns on the monitor's display.
- Standby:* Switches monitor to standby mode when pressed again.



ALARM

- Reset:* When depressed once during an active patient alarm, silences the audio portion of that alarm for fifteen (15) seconds.

When depressed during an active equipment alarm, the alarm condition shall be acknowledged along with the audio and visual shall be removed.

- Silence:* Starts a period of silence when pressed while no alarms are active. The duration of the silence period depends on its previously made configuration.

Note: Press the key twice to enter a silence period while an alarm is active.

The LED lighting scheme indicates the selected functionality:

- Continuous:* 2-Minute audio silence.
- Flashing, 1s:* Permanent audio silence.



NIBP

- Start:* Initiates a NIBP measurement.
- Stop:* Terminates any active NIBP measurement and immediately deflates the cuff.
- NIBP Menu:* Activates the NIBP menu when pressed and held for 2 seconds.



RETURN

- Main Screen:* Returns to the Main screen when pressed while any other screen or menu is being displayed.
- Freeze:* Freezes all traces when pressed while the Main screen is active. Press again to un-freeze. Traces will un-freeze automatically after 60 seconds.
- Print:* Activates a print of the actual screen content (Traces, Trend or Alarm History) when key is pressed for 2 seconds.

Refer to paragraph “Main Screen” on page 33 for more information.



PARAMETERS

Activates the Parameters menu, which provides an overview of parameters and screen layout settings and gives access to change these settings. Refer to paragraph “Parameters Menu” on page 38 for more information.



LIMITS

Activates the Alarm Limits menu, which provides an overview of all actual limit settings and gives access to change, save and restore alarm limits. Refer to section “Alarms” starting on page 41 for more information.



HISTORY

Activates the Trend History and the Alarm History screen. Gives access to erase History data. Refer to paragraph “History Screens” on page 34 for more information.



INDICATORS

Activates the Audio/Visual menu that allows configure the audio and visual signals the monitor can generate. Refer to paragraph “Audio/Visual Menu” on page 39 for more information.

UP



Moves cursor upward or scrolls through menu options, press and hold for quicker advance.

Sets patient mode to Adult when pressed and hold while the monitor is being turned on.

DOWN



Moves cursor downward or scrolls through menu options, press and hold for quicker advance.

Sets patient mode to Neonate when pressed and hold while the monitor is being turned on.

NEXT Function

In the menus: The HISTORY and INDICATORS keys are programmed to advance horizontally to the **next** parameter selection.

PREVIOUS Function

In the menus: The PARAMETERS and LIMITS keys are programmed to move backwards horizontally to the **previous** parameter selection.

Main Screen

Note: When switching from the Main screen to any other screen or menu, the monitor will continue to update and display the numeric values of the parameters being monitored.

Note: The actual displayed information depends on the parameter configuration of the monitor and the user defined screen layout.

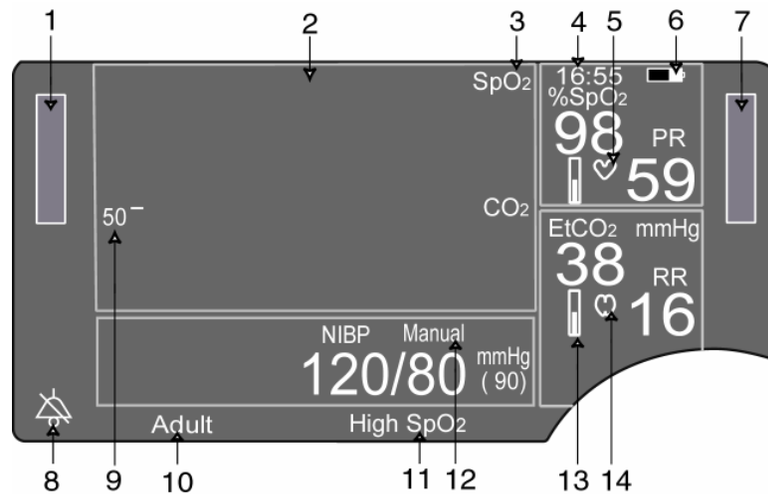


Figure 7: Main Screen

1. Equipment Alarm Indicator
2. Main Display Screen
3. Waveform Selections
4. Time
5. Heart Beat Icon, flashes for every detected pulse beat.
6. Battery Indicator
7. Patient Alarm Indicator
8. Silence Indicator
9. CO₂ size indicator
10. Equipment Message Window
11. Patient Message Window
12. NIBP Mode
13. Bar Graph Indicator for signal strength when traces not displayed.
14. Respiration Breath Icon, flashes for every detected breath.

Freeze Traces

While the Main screen is being display the user can freeze the traces.

- Press the RETURN key.

The message “Traces Frozen” appears at the top of the Main screen. While the traces are frozen, the numerics continue to update. Traces will automatically un-freeze if no key is pressed for 60 seconds or in case any other screen or menu is entered.

- Press the RETURN key again to manually un-freeze traces.

History Screens

The monitor collects History data over a 24-hour period. Continuously measured parameters as %SpO₂, PR, CO₂ and RR are stored as one-minute averages. Sporadically measured NIBP readings and alarm events are stored as they occur.

Note: Turning the power off does not clear the stored data. The stored data will remain in memory for 24 hours. Older data is deleted automatically. It is suggested to manually clear History Data between patients. Refer to paragraph “Erase Trend History Data” on page 37.

Note: The monitor uses an internal Real Time Clock to time stamp all entries. Changes made to either the time or date settings, should be performed in-between patients being monitored.

Note: The information being displayed depends on the monitor configuration and patient specific data.

History data is presented in two screens, the Trend History and Alarm History screen.

- Once the History screen has been displayed, press and hold the HISTORY key for 2 seconds to toggle between the Trend History and the Alarm History screens.
- Press the RETURN key to exit from the History screen.

Note: If no key is pressed for 30 seconds, the monitor will automatically exit to the Main screen.

Trend History

The Trend History screen shows patient data in a tabular form. All continuously monitored parameters are listed as 1-minute average values. Each line represents one minute. Sporadically measured NIBP parameters are displayed in an additional row positioned at the time of occurrence.

- Press the HISTORY key to enter this screen.

Note: If there is no data available, the message “No Trend History” is displayed at the top of the screen.

<u>History</u>				
HR:MN	%O ₂	PR	CO ₂	RR
13:40	98	59	38	16
13:39	NIBP=120/ 80(90)PR= 59			
13:39*	98	59	38	16
13:38	98	59	38	16
13:37	NIBP=112/ 78(89)PR= 61			
13:37	98	59	38	16
13:36	98	59	38	16
13:35	NIBP=120/ 80(90)PR= 58			
13:35	98	59	38	16
Erase No				

Figure 8: Trend History Screen

The cursor position is initially set to the first line of data, which is the most recent data.

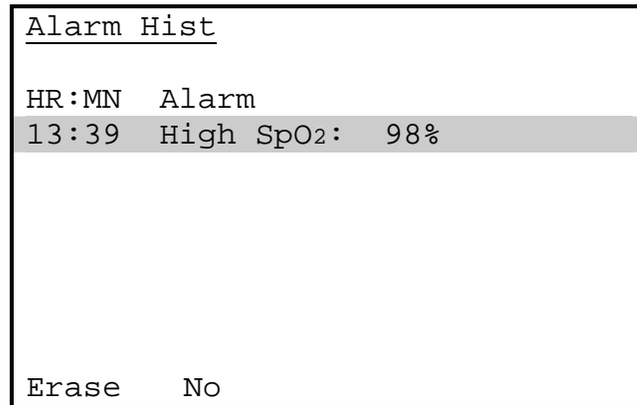
- Press the DOWN key to scroll down (back in time)
- Press the UP key to scroll up (forward time).

Alarm History

The Alarm History screen shows patient alarms in a tabular form as they occurred. Each line represents one alarm event.

- Press the HISTORY key to access the Trend History screen first.
- Then press and hold the HISTORY key for 2 seconds to enter the Alarm History screen.

Note: If there is no data available, the message “No Alarm History” is displayed at the top of the screen.



<u>Alarm Hist</u>	
HR:MN	Alarm
13:39	High SpO ₂ : 98%
Erase No	

Figure 9: Alarm History Screen

The cursor position is automatically set to the first alarm, which represents the most recent event.

- Press the DOWN key to scroll down (back in time)
- Press the UP key to scroll up (forward in time).

Printing Trend or Alarm History

To print the Trend or Alarm History data, proceed as follows:

- Enter the Trend or Alarm History screen by pressing the HISTORY key.
- Press and hold the RETURN key for 2 seconds. The message “Printing” appears in the Equipment Message Window.

Note: Refer to section “Printer” for more information about the printer and a sample printout of the History screen.

Erase Trend History Data

Note: In-between patients, the previously stored data should be erased to avoid any potential misinterpretation of data.

Note: Trend and Alarm History data are not erased at the same time.

Follow these steps to erase Trend History data:

- Press the HISTORY key to access the Trend History screen.
- Press the NEXT key to move the cursor to the line labeled Erase No.
- Use the UP and DOWN key to scroll through the available options. Select “Erase Yes” and press the HISTORY key.

The monitor erases all data in Trend History and returns to the Main screen.

Note: If the monitor contained Alarm History data, the monitor displays the Alarm History screen.

Note: Turning the monitor off or disconnecting it from all power sources does not erase the History data.

Erase Alarm History Data

Note: In-between patients, the previously stored data should be erased to avoid any potential misinterpretation of data.

Note: Trend and Alarm History data are not erased at the same time.

Follow these steps to erase Alarm History data:

- Press the HISTORY key to access the Trend History screen.
- Then press and hold the HISTORY key for 2 seconds to enter the Alarm History screen.
- Press the NEXT key to move the cursor to the line labeled Erase No.
- Use the UP and DOWN key to scroll through the available options. Select “Erase Yes” and press the HISTORY key.

The monitor erases all data in Alarm History and returns to the Trend History screen.

Note: Turning the monitor off or disconnecting it from all power sources does not erase the History data.

Parameters Menu

The Parameters menu shows the settings of all parameters and allows changing them.

- Press the PARAMETERS key to enter this menu.

Note: The information being displayed depends on the monitor configuration and patient specific settings.

Note: The monitor will automatically return to the Main screen if no key is pressed for 30 seconds.

<u>Parameters</u>		
		mm/Sec
Trace 1	SpO ₂	25.0
Trace 2	EtCO ₂	12.5
Trace 3	OFF	
Print Speed		25.0
Print On Alarm	OFF	
EtCO ₂ Scale	0-50 mmHg	
EtCO ₂ Print	OFF	
EtCO ₂ Trace	Line	

Figure 10: Parameters Menu

The cursor position is initially at the first item in the left column.

- Use the UP and DOWN keys to select the item that needs to be changed.
- Use the NEXT key to move the cursor to the next column to the right.
- Use the UP and DOWN key to scroll through the available options.

Note: Several changes may be done in one session. Use the PREVIOUS key to return the cursor back to the left column and repeat the steps as before.

- Press the RETURN key when finished. All changes will be saved and become immediately effective.

Note: A continuously updating Trend History or a menu of NIBP event readings can be chosen, in the Parameters Setup menu as Trace 3.

CO₂ Library

If CO₂ is selected as Trace 2 or 3, a library of 10 CO₂ traces can be selected for educational viewing. Use the UP key to enter the library, and then select other traces by using the UP or DOWN keys. Traces are displayed at 3 mm/Sec. The CO₂ Library will be displayed for 30 seconds and then automatically return to the Main screen.

Audio/Visual Menu

The Audio/Visual menu shows the actual settings and allows changing them.

- Press the VOLUME key to enter this menu.

Note: The monitor will automatically return to the Main screen if no key is pressed for 30 seconds.

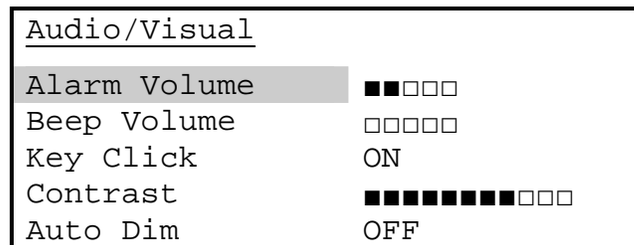


Figure 11: Audio/Visual Menu

The cursor position is initially at the first item in the left column.

- Use the UP and DOWN keys to select the item that needs to be changed.
- Use the NEXT key to move the cursor to the next column to the right.
- Use the UP and DOWN key to scroll through the available options.

Note: Several changes may be done in one session. Use the PREVIOUS key to return the cursor back to the left column and repeat the steps as before.

- Press the RETURN key when finished. All changes will be saved and become immediately effective.

Auto Dim

The CAS 750C Monitor incorporates an Auto Dim feature that, when enable, turns the display's backlight automatically off after 30 seconds. This feature can be helpful when the monitor is being used at night in a patient's room. The Auto Dim feature is enabled when the user selects on in the Auto Dim parameter selection of the Audio/Visual menu.

3 Alarms

The monitor distinguishes between patient and equipment alarms.

Patient Alarms



Warning: Before each use, verify that the alarm limits are appropriate for the patient being monitored.

Note: All patient alarms, based on **continuously** monitored parameters (e.g. CO₂ and SpO₂, etc.) will clear automatically when the alarm cause is no longer persistent.

Note: All patient alarms, based on **discontinuously** monitored parameters (e.g. NIBP readings) will clear only after acknowledged by the user.

Manifestation of Patient Alarms

- Numeric display of the alarming parameter flashes.
- Patient Alarm LED's are activated.
- Audible tone is generated (3 beeps followed by 2 beeps every 10 seconds).
- Flashing text in Patient Message Window explains the cause.

Note: When an ongoing patient alarm is acknowledged by pressing the ALARMS key, only the audible signal will be silenced for 15 seconds. All other alarm indicators will remain active until the alarm condition no longer exists.

Equipment Alarms

Note: All equipment alarms will clear automatically when the alarm cause is no longer persistent.

Manifestation of Equipment Alarms

- Alarming parameter display flashes.
- Equipment Alarm LED's are activated.
- Audible tone is generated (3 beeps every 25 seconds)
- Flashing text in Equipment Message Window explains the cause.

Note: When an ongoing equipment alarm is acknowledged by pressing the ALARMS key, only the audible signal will be disabled. All other alarm indicators will remain active until the alarm condition no longer exists.

Note: Low Battery and Dead Battery alarms cannot be silenced.

Silencing Alarms

In case the monitor should temporarily not generate any alarm tone, the user can activate a Silence Period.

Note: When a Silence Period is being entered, the monitor will not generate an audible signal for patient or equipment alarms with the exception of “Low Battery” and “Dead Battery” conditions.

No alarm active

Pressing the ALARMS key will immediately activate a Silence Period.

Alarm is active

Pressing the ALARMS key will only acknowledge the active alarm and disable the audio signal for 15 seconds. A second keystroke is required to activate a Silence Period.

Silence Indication

The duration of the Silence Period can be selected in the monitors Configuration menu. Refer to paragraph “Configure Audio Silencing” on page 77 for more information. Depending on the selection made, the indication for a Silence Period will be as following:

Silence Period is set to “2-Minute”

- The Silence Indicator will illuminate continuously.
- The message “2 Minute” will be displayed in the Equipment Message Window.
- The Silence Period will be terminated automatically after 2 minutes.

Silence Period is set to “Permanent”

- The Silence Indicator will flash (1 second on/ 1 second off).
- The message “Permanent” will be displayed in the Equipment Message Window.
- To re-activate the audio alarm, the ALARMS key has to be pressed manually.

Alarm Limits

The monitor can store two independent sets of alarm limit values for both Adult and Neonate modes. These sets and the factory defaults can be restored when needed.

Note: Switching between Adult and Neonate mode automatically activates the most recently stored set for patient alarm limits of the appropriate type.

Alarm Limits Menu

The Alarm Limits menu shows the actual settings of all alarm limits and allows changing them.

- Press the LIMITS key to enter this screen.

Note: The information being displayed depends on the monitor configuration and patient specific data.

Note: The monitor will automatically return to the Main screen if no key is pressed for 30 seconds.

Adult 1	Limits		
	Low	High	
%SpO ₂	88	OFF	%
PR	OFF	220	BPM
SatSeconds ⁽¹⁾	OFF		
EtCO ₂	OFF	OFF	mmHg
RR	OFF	OFF	BrPM
No RESP	30		Sec
SYS	OFF	240	mmHg
DIA	OFF	130	mmHg
Save	No		
Restore	No		

Figure 12: Alarm Limits Menu

(1) SatSeconds Limit selection only available with Nellcor SpO₂ oximetry option.

Changing Alarm Limits

The cursor is automatically positioned at the first item in the left column.

- Use the UP and DOWN keys to select the alarm limit that needs to be changed.
- Use the NEXT key to scroll to the next “Low” column or advance to the “High” column by pressing the key again.
- Use the UP or DOWN keys to increase or decrease the alarm limit value.

Note: Several changes may be done in one session. Use the PREVIOUS key to move the cursor back to the left column and repeat the steps as before.

- Press the RETURN key when finished.

The monitor returns to the Main screen, all changes will become effective immediately and remain valid until the monitor is turned off.

Saving Alarm Limits

To save the alarm limit changes, proceed as follows:

- Use the DOWN key and move the cursor to the “Save” position.
- Use the NEXT key and move the cursor to the “No” position.
- Use the UP or DOWN keys and select the appropriate alarm limit set “Adult 1” or “Adult 2” for saving.

Note: Saving options will be “Neo 1” and “Neo 2” when the monitor is in Neonate mode.

- Press the LIMITS key to start the saving process.

The message “Saving” appears and the alarm limit values will be stored in memory.

If a second set of alarm limits is required, repeat these steps and save the changes under a different name (e.g. “Adult 2” or “Neo 2”).

- Press the RETURN key when finished.

The monitor returns to the Main screen, all changes will become effective immediately and remain valid until alarm limits changed manually or a different set of alarm limits is selected.

Restore Alarm Limits

To restore a set of alarm limits, proceed as follows:

- Use the DOWN key and move the cursor to the “Restore” position.
- Use the NEXT key and move the cursor to the “No” position.
- Use the UP or DOWN keys and select the appropriate alarm limit set “Factory”, “Adult 1” or “Adult 2” to restore.

Note: Restore options will be “Factory”, “Neo 1” and “Neo 2” when the monitor is in Neonate mode.

- Press the LIMITS key to start the restore process.

The message “Restoring” appears and the alarm limit values are overwritten with the restored values.

- Press the RETURN key when finished.

The monitor returns to the Main screen and the new limit set will become effective immediately.

Factory Default Alarm Limits

Factory Default Alarm Limits are different for Adult and Neonatal patients.

Parameter	Adult		Neonate	
	Low	High	Low	High
%SpO ₂	88	OFF	88	96
Pulse Rate	OFF	220	OFF	220
SatSeconds	OFF	N/A	OFF	N/A
EtCO ₂	OFF	OFF	OFF	OFF
FiCO ₂ ⁽¹⁾	N/A	3	N/A	3
Resp Rate	OFF	OFF	OFF	OFF
No Resp	30	N/A	20	N/A
Systolic	OFF	240	OFF	120
Diastolic	OFF	130	OFF	80

Table 1: Factory Default Alarm Limits

(1) Factory Set. Not user adjustable

Alarm Volume

The alarm tone volume can be changed but not completely turned off.

- To change the alarm tone volume follow the steps described in paragraph “Audio/Visual Menu” on page 39.

Alarm Delays

To reduce the number of false-positive alarms, some patient alarms will be activated after a short delay. These are:

- EtCO₂ alarms: 10 seconds
- FiCO₂ alarm: 10 seconds
- RR alarms 10 seconds

When the monitor incorporates *Nellcor* SpO₂:

- %SpO₂ alarms: 0 or SatSeconds (configurable)

To change the alarm delay, follow the steps described in paragraph “Changing Alarm Limits” on page 44.

- SpO₂ Pulse Rate alarms: 0 or 10 seconds (configurable)

To change the alarm delay, follow the steps described in paragraph “Selecting Alarm Delay” on page 79.

When the monitor incorporates *Masimo* SpO₂:

- %SpO₂ and Pulse Rate alarms: 0 or 10 seconds (configurable)

To change the alarm delay, follow the steps described in paragraph “Selecting Alarm Delay” on page 79.

Note: All other alarms are generated without any delay.

Stand By mode

This mode can be used just prior to connecting the monitor to a patient, but all the monitor's cables are connected.

- Depress and holding for two (2) seconds the SILENCE/RESET pushbutton.

When enabled, all Equipment alarms are silenced and the message "Stand By" is displayed on the Main display.

For the Stand by feature to work properly, the following conditions must apply;

- The monitor cannot be actively monitoring any patient parameter.

Note: Temperature monitoring is not effected by the Stand By mode.

To return the monitor to normal operation, connect any of the patient leads to the patient.

4 SpO₂

Pulse Oximetry

Introduction

The Pulse Oximeter parameter (SpO₂) determines arterial oxyhemoglobin saturation by measuring the absorption of red and infrared light passing through the tissue. Changes in absorption caused by pulsations of blood are used to determine Arterial Saturation (%SpO₂) and Pulse Rate (PR).

If SpO₂ was not selected as a trace to be displayed, a SpO₂ signal bar graph will be displayed indicating the relative signal strength and signal quality at the sensor site.

Note: The bar graph indicator is not proportional to the pulse volume.

An audio “beep” can be enabled that is generated each time the SpO₂ module detects a pulsation.

Safety



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



Warning: Accurate oxygen saturation measurements cannot be obtained when the oximeter is not measuring the pulse properly. If the SpO₂ waveform, perfusion bar graph or the Pulse Rate be erratic or inaccurate, first examine the patient for any sign of distress and only then re-examine sensor placement.



Warning: A pulse oximeter should be considered an early warning device. As a trend toward patient deoxygenation is indicated, blood samples should be analyzed by a laboratory co-oximeter to completely understand the patient's condition.



Warning: Various patient or mechanical conditions may cause inaccurate measurements. These conditions include but are not limited to: venous pulsations, hypotension, severe vasoconstriction, severe anemia, hypothermia, shock, cardiac arrest, sensor tension, sensor placement on the same extremity as a blood pressure cuff, arterial catheter or intravascular line, or arterial occlusion proximal to the sensor.



Warning: *MRI Scanning* - Do not use oximetry sensors during magnetic resonance imaging (MRI) scanning. Conducted current could cause burns.



Warning: Do not expose probe detector to strong ambient light while monitoring a patient. A poor signal may result.



Warning: If a probe is damaged in anyway, discontinue use immediately.



Caution: Cardiogreen and other intravascular dyes, depending on the concentration, may affect the accuracy of the oximeter measurement.



Caution: Significant levels of dysfunctional hemoglobins such as carboxyhemoglobin or methemoglobin may affect the accuracy of the measurement.



Caution: Certain antibiotics, such as the Sulfas, can create high levels of methemoglobin. Methemoglobin is unable to bind O₂ and will absorb light similarly to reduced hemoglobin, thus giving an artificially low SpO₂.



Caution: *Reusable* SpO₂ sensors - Prolonged monitoring may require changing the probe site periodically. The sensor must be removed and repositioned every 4 hours and if indicated by circulatory condition or skin integrity, reapplied to a different monitoring site.



Caution: *Disposable* SpO₂ sensors - Prolonged monitoring may require changing the probe site periodically. The sensors must be removed and repositioned every 8 hours and if indicated by circulatory condition or skin integrity, reapplied to a different monitoring site. This sensor type should not be used with patients that exhibit allergic reactions to adhesive tape.



Caution: *Masimo LNOP probe* - Tissue damage can be caused by incorrect application or use of an LNOP probe, for example by wrapping the probe too tightly. Inspect the probe site as directed in the probe's directions for use to ensure skin integrity and correct positioning and adhesion of the probe.

SpO₂ Sensors

The monitor can be equipped to use SpO₂ sensors manufactured by Masimo or Nellcor. The appropriate manufacturer's logo is shown next to the SpO₂ input connector.

Note: No other manufacturer's sensors should be used.

Select a sensor based on the patient size and monitoring conditions and properly attach the sensor to the patient.

Note: Consult instructions enclosed with each sensor for proper application.

There are a variety of SpO₂ sensors designs for a specific clinical application. Please refer to the Instructions of Use, which are supplied with each type of sensor.

Preparations

Follow the instructions provided by the sensor manufacturer (Masimo or Nellcor) and prepare for SpO₂ monitoring as follows:

- Select a sensor that has the appropriate size for the patient's digit or extremity.
- Clean the surface of the probe before and after each patient use.
- Correctly position and attach the probe to the patient.
- Connect the sensor cable to the SpO₂ patient cable.
- Verify a secure connection and gently tug on the patient cable connector.
- Plug the SpO₂ patient cable into the SpO₂ connector on the left side panel of the monitor.

Note: The SpO₂ probe must be kept as motionless as possible to make a proper determination. To minimize motion artifacts, secure the sensor cable independently from the sensor. Tape may be used to secure the cable to the patient, e.g. around the base of the finger. Make sure that the tape being used does not restrict the blood flow.

SpO₂ Monitoring

Once the sensor has been attached to the patient and the cable to the monitor, proceed as follows:

- Press the POWER key to turn the monitor on.
- If required, configure the appropriate waveform to be displayed on the Main Screen.
- Verify the signal quality and strength with the help of the SpO₂ waveform or the bar graph indicator.
- Wait for the monitor to determine the initial %SpO₂ and Pulse Rate values.



Caution: Some sensors may not be appropriate for a particular patient. If at least ten seconds of one bar pulses cannot be observed for a given sensor, change sensor location or sensor type until this condition is achieved.

- Check the alarm limits and configure them appropriately for the patient.

Note: Inspect the SpO₂ sensor site every 2 to 4 hours or per hospital protocol. If there is any skin irritation caused by the sensor, remove the sensor and apply it to a different location.

SatSeconds™ Alarm Management

Note: The SatSeconds™ alarm management feature is only available in monitors with Nellcor's Oximax technology.

False or nuisance alarms are a common concern with pulse oximetry monitoring. Nuisance alarms are often triggered by minor and brief desaturation events that are clinically insignificant. Clinicians tend to manage these alarms by ignoring them, turning off the alarm or monitor, or widening the alarm limits. The CAS 750C Monitor incorporates Nellcor's SatSeconds Alarm Management feature that offers a better way to manage nuisance alarms without sacrificing patient safety.

The SatSeconds function can be activated from the Alarm Limits menu by selecting a SatSeconds limit, or "clock" of 10, 25, 50, 100 or OFF (0) SatSeconds. Clinicians who choose to employ the SatSeconds function should select a limit suited to their clinical environment and patient conditions. Think of SatSeconds as the product of magnitude and time a patient exceeds SpO₂ alarm limits. For example, 3 points below the alarm limit for 10 seconds equals 30 SatSeconds. An alarm is only triggered if a desaturation event occurs that reaches the SatSeconds limit you selected. As a safety net, when three or more SpO₂ alarm violations occur within 60 seconds, an alarm will sound even if the SatSeconds limit has not been reached.

To activate the SatSeconds function, proceed as follows:

- Press the LIMITS key to enter the Alarm Limits menu.
- Use the DOWN key to select "SatSeconds".
- Use the NEXT key to scroll to the next column to the right.
- Use the UP or DOWN keys to increase or decrease the SatSeconds value.
- Press the RETURN key when finished. All changes will be saved and become immediately effective.

Note: When SatSeconds is set to OFF, the alarm delay for %SpO₂ is based on the current configuration. Refer to paragraph "Selecting Alarm Delay" on page 79.

Disconnecting SpO₂ Accessories

When SpO₂ monitoring is not required, disconnect the patient cable from the monitor by squeezing the tabs with your thumb and index finger while pulling the connector away from the monitor.

Note: To avoid damage to the Interface Cable, always hold it by the connector rather than the cable when connecting or disconnecting either end.

When the probe is disconnected from the monitor, an Equipment Alarm is activated and the message “No SpO₂ Probe” appears. Press the ALARMS key to acknowledge the alarm.

SpO₂ Display Window



Oxygen Saturation (%SpO₂) and Pulse Rate (PR) are displayed in the green window at the upper right area of the Main screen.

Note: The bar graph indicator is only presented if the SpO₂ waveform is not displayed in one of the trace windows.

SpO₂ related Settings

Access the Parameters menu as described on page 38 to review or change the following settings:

- SpO₂Trace Speed selection

Access the Alarm Limits menu as described on page 43 to review or change the following settings:

- SpO₂ Alarm Limits

Access the Audio/Visual menu as described on page 39 to review or change the following settings:

- Beep Volume

SpO₂ Troubleshooting

Note: If the accuracy of any measurement does not seem reasonable, first check the patient's vital signs by an alternate method.

Note: The %SpO₂ and related Pulse Rate numeric displays will show dashes "--" and the signal strength window will be blank when no SpO₂ probe is attached to the monitor.

Note: Inaccurate SpO₂ measurements may be caused by:

- Anemia or low hemoglobin concentrations.
- Electro surgical interference.
- Excessive ambient light.
- Excessive patient movement.
- Incorrect sensor application or use.
- Intravascular dyes such as indocyanine green or methylene blue.
- Moisture in the sensor.
- Placement of a sensor on an extremity with a blood pressure cuff, arterial catheter, or intravascular line.
- Venous pulsations.

Note: The loss of a pulse signal can occur in any of the following situations:

- A blood pressure cuff is inflated on the same extremity as the one with the SpO₂ sensor attached.
- Excessive ambient light such as from a surgical lamp, a bilirubin lamp, or sunlight is present.
- The patient has hypotension, severe vasoconstriction, severe anemia, or hypothermia.
- The patient is in cardiac arrest or is in shock.
- The sensor is too tight.
- There is arterial occlusion proximal to the sensor.

SpO₂ related Messages

The following SpO₂ related messages may occur:

Chk SpO₂ Probe

Note: This message relates to Masimo technology only.

The monitor is questioning the quality of the signal being received by the SpO₂ probe or the probe is receiving too much ambient light.

- Press the ALARMS key. The monitor silences the audio alarm tone, but the visual messages remain.
- Verify that the probe being used is the correct one for the monitor's SpO₂ configuration.
- Replace a wrong or defective probe.
- Verify that the probe is being used according to the manufacturer's recommendations.
- Verify that the probe emitter and detector are parallel to and directly opposing each other.

Low Perfusion

Note: This message relates to Masimo technology only.

The perfusion level being received by the SpO₂ probe is low.

- Determine if an extreme change in the patient's physiology and blood flow at the monitoring site occurred (e.g. an inflated blood pressure cuff, a squeezing motion).
- Try to warm the patient or the probe site.
- Move probe to a site with better perfusion.

Low Signal IQ

Note: This message relates to Masimo technology only.

The quality of the signal level being received by the SpO₂ probe is in question.

- Ensure proper probe type and application.
- Verify that the probe emitter and detector are parallel to and directly opposing each other.
- Clean or replace the probe.

Note: When the message "Low Perfusion" or "Low Signal IQ" appear and the monitor is displaying valid %SpO₂ numerics, no audible or visual alarms will be generated. In case the monitor is not displaying valid %SpO₂ numerics, an audible and visual alarm will be generated.

No SpO₂ Probe

The monitor is not detecting the SpO₂ probe. The probe was disconnected from either the Interface Cable or from the monitor.

- Press the ALARMS key. The monitor silences the audio alarm tone, but the visual messages remain.
- Reconnect probe to monitor.

Probe Error

The SpO₂ probe being used is not the correct one for the monitor's configuration.

- Press the ALARMS key. The monitor silences the audio alarm tone, but the visual messages remain.
- Verify the probe being used is the correct probe.

Pulse Search

The monitor is searching for a Pulse signal. This is normal at power-up as the monitor searches for a pulse or the probe position may have changed.

- Check the probe site.

SpO₂ Error

A failure occurred on the SpO₂ board. The monitor needs to be serviced.

- Press the ALARMS key. The monitor silences the audio alarm tone, but the visual messages remain.

SpO₂ Module

A failure occurred on the SpO₂ board. The monitor needs to be serviced.

- Press the ALARMS key. The monitor silences the audio alarm tone, but the visual messages remain.

SpO₂ Probe OFF

The monitor is no longer receiving a patient signal from the SpO₂ probe. The probe is no longer in contact with the patient.

- Press the ALARMS key. The monitor silences the audio alarm tone, but the visual messages remain.
- Check if probe is properly applied to the patient.
- Check if probe cable is connected to extension cable.
- Check if cable is attached to monitor.
- Replace probe and/or cable.

5 CO₂ Capnography

Introduction

The monitor uses Microstream non-dispersive infrared (NDIR) spectroscopy to continuously measure the amount of CO₂ during every breath. The amount of CO₂ at the end of exhalation (EtCO₂) and the Respiratory Rate (RR) are measured.

The CO₂ measurement is intended for use in any environment where continuous, noninvasive monitoring is desired, including hospital and mobile use (when protected from excessive moisture such as direct rainfall). It is intended for use on adult, pediatric, and infant/neonatal patients. It is intended for use on intubated and non-intubated patients.

A CO₂ library can be selected for educational viewing. Refer to paragraph "CO₂ Library" on page 39.

Safety



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



Warning: When monitoring CO₂ in the presence of flammable gases or anesthetics, such as high concentrations of oxygen or nitrous oxide, connect the gas outlet to a scavenger system.



Warning: CO₂ readings and respiratory rate can be affected by certain ambient environmental conditions, sensor application errors, and certain patient conditions.



Caution: Microstream consumables are latex free, non-sterile and non-pyrogenic.



Caution: During MRI scanning, the monitor must be placed outside the MRI suite. When the monitor is used outside the MRI suite, CO₂ monitoring can be implemented using the FilterLine XL.

Preparations



Caution: Prior to patient monitoring, ensure the monitor is configured to the appropriate patient mode – Neonate or Adult. Refer to paragraph “Selecting Patient Mode” on page 76.

Microstream CO₂ Consumables

Note: The Microstream CO₂ technology requires the use of Microstream consumables. No other manufacturer’s products should be used.

Refer to paragraph “Capnography Accessories” on page 126 for CO₂ consumable types and order information.

When choosing CO₂ consumables for a particular patient, the following should be considered:

- Intubated or non-intubated.
- Whether the patient is on mechanical ventilation.
- Duration of use.
- Patient’s size and weight



Caution: Before use, carefully read the Directions for Use that is enclosed with the Microstream consumable.



Caution: Microstream consumables are designed for single patient use, and are not to be reprocessed.



Warning: Do not attempt to disinfect or flush the FilterLine as the monitor can be damaged.

Non-Intubated Application

For non-intubated patients the continuous gas sampling is taken from the patient’s nose and/or mouth. The following consumables are available:

- CapnoLine: Cannula with nasal patient adapter.
- Smart CapnoLine Oral Nasal Cannula – for use in procedural sedation. Also available with O₂ delivery.
- CapnoLine H – for use in ICU for patients receiving hi-flow oxygen by mask or on long term CPAP or Bi-PAP. Also available with O₂ delivery.
- NIV-Line – for use under oxygen CPAP, Bi-PAP or NPPV mask and to avoid intubation during EMS transport.

Select the appropriate CapnoLine and connect it to the monitor before attaching it to the patient’s airway. Be sure to follow Microstream CO₂ Consumables’ *Directions for Use* for proper connection.



Caution: Loose or damaged connections may compromise ventilation or cause an inaccurate measurement of respiratory gases. Securely connect all components and check connections for leaks.



Warning: Carefully route the CapnoLine to reduce the possibility of patient entanglement or strangulation.

Note: Ensure the CapnoLine is not twisted or crimped and the cannula tips are in the nostrils.

Note: If the CapnoLine includes an additional oxygen connection, connect the oxygen connector to the standard outlet.

Intubated Applications

For intubated patients the continuous gas sampling is taken from the Airway Adapter directly at the intubation tube. The following consumables are available:

- FilterLine set for non-humid environments.
- FilterLine H set for humid environments.

Select the appropriate FilterLine and connect it to the monitor before attaching it to the patient's airway. Be sure to follow Micro-stream CO₂ Consumables' *Directions for Use* for proper connection.



Warning: When used with closed suction system, do not place the airway adapter between the suction catheter and endotracheal tube.



Caution: Loose or damaged connections may compromise ventilation or cause an inaccurate measurement of respiratory gases. Securely connect all components and check connections for leaks.

Note: Ensure the consumable is not twisted or crimped.

Note: During nebulization or suction (when not using closed suction system), in order to avoid moisture buildup and FilterLine occlusion, disconnect the Airway Adapter from the patient's endotracheal tube.

CO₂ Monitoring

Once the appropriate consumable has been applied to the patient, proceed as follows:

- Press the POWER key to turn the monitor on.

Note: While the CO₂ module is initializing (typically 30 to 40 seconds) the monitor shall display the message “CO₂ Warm-Up” and dashes “- -” will remain in the numeric section.

- Check that the monitor is accurately detecting the CO₂ signal and wait for the monitor to determine the initial EtCO₂ and Respiration Rate values.
- If the lung visual indicator is not corresponding to the patient’s respiration, reposition the sensor until the indicators flash in synch with the patient’s breathing. This will help to minimize false alarms.

Check the alarm limits and configure them appropriately for the patient. Refer to paragraph “Alarm Limits Menu” on page 43.

Removing the CO₂ Consumables

When CO₂ monitoring is no longer required, disconnect the consumable by carefully removing the connector from the CO₂ input receptacle.

When the consumable is disconnected from the monitor, the message “EtCO₂ OFF” is displayed, the Equipment Alarm LEDs are flashing and an audible alarm sounds, indicating a connection has been broken.

Note: If CO₂ is selected as a waveform, messages appear in the CO₂ waveform window - otherwise messages appear in the Equipment Message Window of the Main screen.

- To acknowledge the alarm, press the ALARMS key.



Caution: Dispose of Microstream consumables according to standard operating procedures or local regulations for the disposal of contaminated medical waste.

CO₂ Display Window



End Tidal CO₂ (EtCO₂) and Respiration Rate (RR) are displayed in the yellow window at the lower right area of the Main screen.

Note: The bar graph indicator is only presented if the CO₂ waveform is not displayed in one of the trace windows.

CO₂ related Settings

Access the Parameters menu as described on page 38 to review or change the following settings:

- EtCO₂ Trace speed selection
- EtCO₂ Scale selection
- EtCO₂ Print selection
- EtCO₂ Trace Type selection

Access the Alarm Limits menu as described on page 43 to review or change the following settings:

- EtCO₂ Alarm Limits

Access the Monitor Setup menu as described on page 79 to change the following settings:

- EtCO₂ Unit selection (mmHg, kPa or %).

CO₂ Troubleshooting

Note: If the accuracy of any measurement does not seem reasonable, first check the patient's vital signs by an alternate method.

CO₂ related messages

- Pressing the ALARMS key will silence the audio alarm tone for the following CO₂ related messages.

Blocked Line

The CO₂ module has attempted to purge the line with no success.

- Disconnect the FilterLine.
- Check the FilterLine for kinks and/or blockages.

Cal EtCO₂

The CO₂ module's calibration timer has reached zero hours.

- Perform a CO₂ calibration check.

Clearing Line

The CO₂ module has detected an occlusion and is attempting to purge the line.

- Check the FilterLine tube for kinks and/or blockages.

Chk EtCO₂ Flow

The CO₂ module has detected a blockage in the exhaust port.

- Check the exhaust port and associated tubing for kinks and blockages.

EtCO₂ Error

A failure occurred on the CO₂ Board.

- To reset the CO₂ board turn the monitor off and disconnect it from external power (AC Line or 12VDC). Remove the battery, and then reconnect the battery and external power. Turn the monitor on.
- If the message remains, the monitor needs to be serviced.

EtCO₂ Module

A failure occurred on the CO₂ Board. The monitor needs to be serviced.

EtCO₂ OFF

The CO₂ module is no longer receiving a signal from the patient or the FilterLine is no longer connected to the monitor. The consumable is no longer connected to the monitor.

- Press the ALARMS key. The monitor silences the audio alarm tone, but the visual messages remain.
- Check the external tubing for disconnections.

Service EtCO₂

The CO₂ module's service timer has reached zero hours. The monitor will continue to function.

- Arrange for service as soon as possible.

6 NIBP

Non-Invasive Blood Pressure

Introduction

The Non-Invasive Blood Pressure (NIBP) parameter is using the oscillometric measurement technique. It automatically inflates an occluding cuff and then releases the cuff pressure step by step. It determines systolic, diastolic and mean arterial pressure. The NIBP function is suitable for neonatal, pediatric and adult patients.

Safety



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



Warning: Do not place the cuff on a limb being used for A-V fistulas, intravenous infusion or on any area where circulation is compromised or has the potential to be compromised.



Warning: When monitoring over an extended period of time, or at frequent intervals, periodically observe the patient's limb to make sure that the circulation is not impaired for a prolonged period of time.



Warning: The position of subject, physiological condition and other factors affect the NIBP readings. Blood pressure and pulse rate can fluctuate greatly between measurements; the monitor cannot alert the user to changes in vital signs occurring between measurement cycles.



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



Warning: Do not apply external pressure against the cuff while monitoring. Doing so may cause inaccurate blood pressure values.



Caution: Always be sure to check that there are no signs of prolonged impairment of circulation regardless of automatic safety features that restrict the over all measurement time.



Caution: In shock conditions, the low amplitude of the blood pressure waveform may make it difficult for the monitor to accurately determine the systolic and diastolic pressures.



Caution: The monitor does not operate effectively if a patient is having seizure activity, tremors or is connected to a heart/lung machine.



Caution: Consult a physician for interpretation of blood pressure measurements.



Caution: As with any non-invasive oscillometric blood pressure monitor, the accuracy of the measurements obtained may be adversely affected by the presence of agents, which alter the patient's cardiovascular system.



Caution: Do not operate the monitor's NIBP function unless it has been properly calibrated. Inaccurate blood pressure readings may result.



Caution: When a patient is experiencing arrhythmias during a measurement, the accuracy of the pulse determination may be affected or the time needed to complete a measurement may be extended.



Caution: Do not alter the monitor's air hose. CAS Medical Systems, Inc. cannot ensure proper monitor performance if the tubing is altered. Modification of the air hose will void the warranty. Avoid compression or restriction of pressure tubes.

Note: A patient's vital signs may vary dramatically during the use of cardiovascular agents such as those that raise or lower blood pressure or those that increase or decrease heart rate.

Note: The maximum amount of time allowed completing a NIBP measurement is 120 seconds in Adult mode and 90 seconds in Neonate mode. If the measurement has not been completed within that time, the cuff is deflated automatically and a message is displayed indicating the problem.

Note: To prevent exposure of the extremity to an inordinately high pressure, the cuff is deflated automatically when the pressure in the system is greater than 290 mmHg in the Adult mode or 145 mmHg in the Neonate mode.

Note: In the event of a microprocessor failure, the cuff will be deflated automatically within 10 seconds.

Preparations

Patient Mode Selection



Caution: Prior to patient monitoring, ensure the monitor is configured to the appropriate patient mode – Neonate or Adult. Refer to paragraph “Selecting Patient Mode” on page 76.

Select the Neonate mode when measuring on newborn patients and for all babies where the systolic pressure is expected to be below 120 mmHg. Use the Adult mode for infant and adult patients.

Switch from Neonate to Adult mode if one or more of the following conditions occur:

- Systolic pressure is expected to be above 120 mmHg.
- A cuff width of 6 cm or larger is required.

NIBP Hose Selection

Two different inflation hoses are available.

- The coiled ten-foot NIBP hose is recommended for adult and pediatric patients.
- The straight six-foot NIBP hose is recommended for neonatal patients and small infants.

Cuff Selection

The use of the correct size of cuff is essential for the accurate measurement of the blood pressure.

Note: CAS recommends the use of its Tuff-Cuff reusable cuffs, Safe-Cuff disposable cuffs and Pedisphyg neonatal cuffs.

Width: The widest cuff that can be placed around the upper arm or thigh should be used. A cuff that is too small for the arm will not supply sufficient pressure to the artery. This can cause an erroneously high blood pressure reading.

Length: The edge of the cuff is marked with a white arrow. It should fall within the white range marking when the cuff is applied to the patient. Otherwise the cuff is too long or too short.

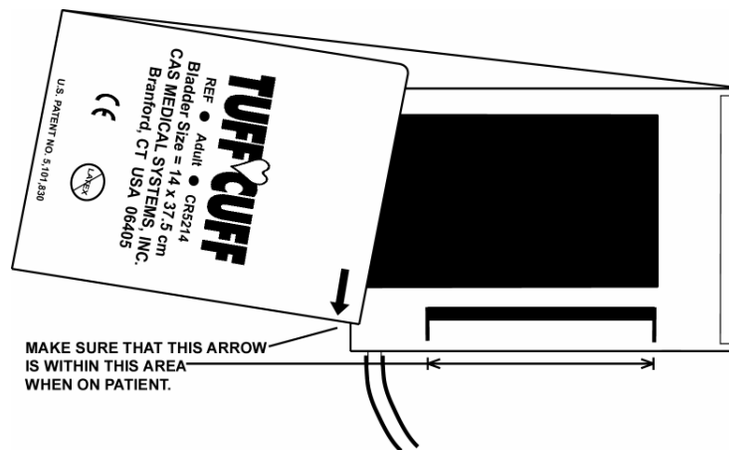


Figure 13: Cuff Application Range Marker

Note: Overlapping the cuff will not affect the measurement results.

Cuff Application

Measurements made above the level of the heart will give reduced blood pressure readings while measurements made below the heart level will give increased readings. These errors are mainly due to the weight of the blood.



Figure 14: Cuff Application

CAS cuffs do not require an exact placement of the hose over the brachial artery. The design of a fully encircling bladder ensures that the artery is properly compressed independent of the hose placement. It can be placed in any position.

- Wrap the cuff for a snug fit to avoid prolonged pumping time. Do not wrap a cuff over the patient's clothing.
- Verify the hook and loop sections of the cuff are fully engaged when it is wrapped around the limb.
- The limb should be positioned to be at heart level.
- Do not compress the cuff or the cuff hose. The hose must not be kinked or pinched.



Warning: The cuff should not be applied on a limb being used for an intravenous infusion. Do not place the cuff on any extremity being used for SpO₂ monitoring.

Note: Remember that there may be a marked difference between readings taken from the left and the right arms. Be consistent with each patient.

NIBP Measurements



Warning: When monitoring over an extended period of time, or at frequent intervals, periodically observe the patient's limb to make sure that the circulation is not impaired for a prolonged period of time.



Caution: Make sure the monitor is set to the correct Patient Mode. Refer to page 30 or to paragraph "Selecting Patient Mode" on page 76 to change the patient mode.

Note: The cuff target pressure must be higher than the patient's systolic pressure to obtain an accurate systolic and diastolic reading.

The initial cuff inflation pressure is 150 mmHg in Adult mode and 85 mmHg in Neonate mode. After the first measurement, the monitor will adapt the cuff inflation pressure automatically based on the previously measured systolic value (approximately 30 mmHg above). This will continue until the monitor is turned off and on again or the inflation pressure is selected manually.



Caution: When measuring blood pressure on a pediatric patient using the Adult mode, it is recommended that the Initial Inflation Pressure be set to a lower value (e.g. 120 mmHg).

Once the cuff is selected and applied to the patient, proceed with the following steps:

- Connect the cuff to the end of the NIBP hose and make sure the hose is connected to the NIBP connector on left side of the monitor.

Starting a Blood Pressure Reading



Caution: Excessive motion can contribute to inaccurate measurements. It is important that the patient be kept still during a measurement. Make every attempt to alleviate patient's fear, anxiety and pain.

Note: The NIBP key will start a NIBP measurement regardless of the type of screen that is currently displayed. The monitor will switch to the Main screen if this is not already on display.

- To select an alternate Initial Inflation Pressure, enter the NIBP menu as described in this section on page 71.
- Press the NIBP key to start a measurement.

The cuff inflation pressure will be displayed until the measurement is completed.

The measurement typically takes less than 30 seconds to complete. In no case will the cuff remain pressurized for more than 120 seconds for Adult/Pediatric patients and no more than 90 seconds for Neonates.

When the measurement is completed, the cuff will automatically deflate, the monitor will provide an audio indication and display the NIBP values: Systolic/Diastolic and (MAP) or Systolic/Diastolic and Pulse Rate.

Note: If any displayed NIBP measurement were to be left on the screen for up to 24 hours, the monitor will automatically blank the NIBP displays to all dashes "- - -".

Stopping a Blood Pressure Reading

A blood Pressure Reading can be interrupted for any reason at any time.

- Pressing the NIBP key during an ongoing NIBP measurement will interrupt the measurement and immediately deflate the cuff.

Entering the Cycle Mode

If a cycle time is selected the monitor will automatically take blood pressure measurements at pre-selected time intervals.

- Enter the NIBP Menu as described in this section on page 71.
- Select the desired cycle time and return to the Main Screen by pressing the RETURN key. The selected cycle time will be displayed in the NIBP window.
- Press the NIBP key to begin the first measurement.

The measurement results are displayed in the NIBP Numeric Area of the Main Screen until the end of the next Automatic measurement or the start of the next Manual measurement. If applicable, the selected cycle time and the remaining time until the next measurement will be displayed as well.

Note: If a measurement is desired between measurement cycles, press the NIBP key. After this measurement, the monitor will continue the Automatic Cycle mode.

Terminating the Cycle Mode

To return to the Manual NIBP mode, proceed as follows:

- Enter the NIBP Menu as described in this section on page 71.
- Change the “NIBP Cycle” entry back to Manual.

STAT Mode

The monitor will take a series of automatically started blood pressure measurements for a period of 5 minutes when STAT mode is activated. Each measurement is followed immediately by the next one after a 10 seconds pause that allows venous blood to return.



Warning: Readings obtained during STAT mode may not meet the stated accuracy of this monitor.



Warning: In some cases, rapid, prolonged cycling of an oscillometric, noninvasive blood pressure monitor cuff has been associated with any or all of the following: ischemia, purpura, or neuropathy. Apply the oscillometric cuff appropriately, according to instructions, and check the cuff site and cuffed extremity regularly when blood pressure is measured at frequent intervals or over extended periods of time.

Entering STAT Mode

To activate the STAT mode, proceed as follows:

- Enter the NIBP Menu as described in this section on page 71.
- Move the cursor to STAT and then select ON.
- Return to the Main Screen by pressing the RETURN key. The message "STAT" appears in the NIBP window.
- Press the NIBP key to begin the first measurement.

Between readings, the NIBP window will display the remaining seconds until the next measurement.

Note: Selecting STAT mode will override any time interval selected for the NIBP measurement cycle.

Exiting STAT Mode

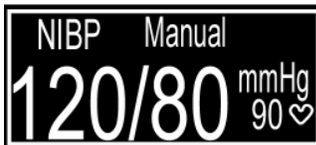
The monitor will stop the STAT mode automatically after 5 minutes and returns to the Manual NIBP mode.

- Press the NIBP key to terminate the current NIBP measurement.
- Enter the NIBP Menu as described in this section on page 71.
- Select STAT mode OFF.

NIBP Display Window



The NIBP numeric values (SYS, DIA and MAP) are displayed in the blue window at the lower center of the Main screen.



Note: The monitor may be configured to display the Pulse Rate (PR) instead of the Mean Arterial Pressure (MAP).

NIBP Menu

To enter the NIBP Menu

Press and hold the NIBP key for 2 seconds to access the NIBP menu. The following screen will be displayed:

<u>NIBP</u>	
NIBP Cycle	Manual
Init Inflate	---
STAT	OFF
(MAP)/PR	(MAP)

Figure 15: NIBP Menu

Note: While in the NIBP menu and no key is pressed for 30 seconds, the monitor will automatically save all changes made and return to the Main Screen.

Menu Options

The NIBP menu allows the user to:

- Select the cycle time. Available selections are: Manual, 1, 2, 3, 4, 5, 10, 15, 30, 60 or 90 Minutes.
- Select the initial Inflation Pressure for the next NIBP measurement. Available selections are:
 - Neonate mode: 60, 80, 85, 100 or 120 mmHg
 - Adult mode: 80, 100, 120, 140, 150, 160, 180 or 200 mmHg
- Select the STAT mode to be OFF or ON
- Select MAP or Pulse Rate to be displayed in the NIBP display window.

Operating the NIBP Menu

To change settings, proceed as follows:

- Use the UP and DOWN keys to select the desired row.
- Move the cursor to the right with the use of the NEXT keys.
- Use the UP and DOWN keys to select the desired function or setting.

NIBP related Settings

Access the Parameters menu as described on page 38 to review or change the following settings:

- NIBP History selection for trace 3.

Access the Alarm Limits menu as described on page 43 to review or change the following settings:

- NIBP Alarm limits

NIBP Troubleshooting

Note: If the accuracy of any measurement does not seem reasonable, first check the patient's vital signs by an alternate method.

NIBP related Messages

- Pressing the ALARMS key will cause the following NIBP related messages to be removed from the display and silence the audio alarm tone.

Air Leak

Air leak in the cuff, the NIBP hose or in the internal pneumatic system of the monitor.

- Check that the cuff/hose/monitor connection is secure.
- Check cuff for leaks. Do not use a known leaky cuff.

Appl Error

Neonate cuff is detected in Adult Mode.

- Check cuff. Replace cuff or change operating mode.

Flow Error

Stable cuff pressure cannot be maintained by the NIBP pneumatic system.

- Check the external tube for kinks.
- Perform a Pneumatic Pressure Check as described on page 90.
- Replace cuff.

Loose Cuff

Cuff applied too loosely.

- Check cuff for proper fit on patient.

Motion

There was too much extremity motion for the monitor to accurately complete the NIBP measurement in 120 seconds. Measurements can be obtained when there is limited extremity movement, but the measurement time may be extended.

- Restrain patient extremity motion.

NIBP Cal

Pressure calibration data corrupted within NIBP module. Pressure module needs recalibration. The monitor needs to be serviced.

NIBP Error

A failure occurred on the NIBP board. The monitor needs to be serviced.

NIBP Module

A failure occurred on the NIBP board. The monitor needs to be serviced.

Over Press

Cuff pressure exceeded 290 mmHg in the Adult mode or 145 mmHg in the Neonate mode. Very rapid squeezing of the cuff can cause this error.

- Repeat the measurement.
- If this message repeatedly occurs during normal use, the monitor must be serviced.

Range Error

The systolic reading exceeds the measurement range of 255 mmHg in the Adult mode or 135 mmHg in the Neonate mode.

- Repeat measurement.
- If the message is displayed again, use another method to measure the patient's blood pressure.

Signal Sat

Signal Saturation or motion pulses too strong.

- Repeat measurement. Limit patient activity; the arm must be still and/or relaxed.

Time Out

The monitor was unable to complete a measurement within 120 seconds in the Adult mode or 90 seconds in the Neonate mode. An extremely long measurement can be due to a loose cuff, high blood pressure, or monitor re-pumps.

- Repeat measurement.
- Try higher initial cuff pressure.
- If message consistently reappears try using another means to obtain patient's blood pressure.

Weak Signal

The monitor did not detect any pulses during a NIBP measurement.

- Check the fit of the cuff.
- Repeat measurement.

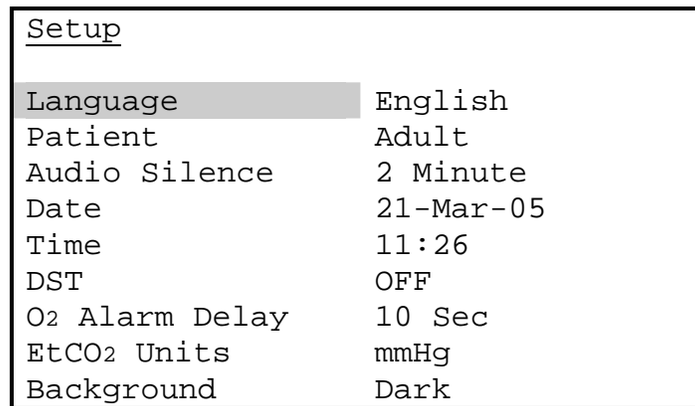
7 Setup

Configuring the Monitor

Entering the Setup Menu

The Setup menu shows all configuration options in a tabular form. To enter the Setup menu, proceed as follows:

- Press and hold the LIMITS and VOLUME keys while the monitor is being turned on.



The screenshot shows a terminal-style window titled "Setup". The window contains a table of configuration options. The first row, "Language", is highlighted with a grey background. The table lists various settings such as Patient type, Audio Silence duration, Date, Time, DST status, O2 Alarm Delay, EtCO2 Units, and Background color.

<u>Setup</u>	
Language	English
Patient	Adult
Audio Silence	2 Minute
Date	21-Mar-05
Time	11:26
DST	OFF
O2 Alarm Delay	10 Sec
EtCO2 Units	mmHg
Background	Dark

Figure 16: Setup Menu

The cursor is initially positioned at the first item in the left column.

- Use the UP and DOWN keys to select the item that needs to be changed.
- Use the NEXT key to move the cursor to the right column.
- Use the UP and DOWN key to scroll through the available options.
- Press the RETURN key to exit the Setup menu.

Note: If no key is pressed for 60 seconds, the monitor automatically exits the Setup menu.

All changes will be saved and the monitor returns to the Main screen.

Selecting the Language

To select the monitor's language, proceed as follows:

- Enter the Setup menu as described on page 75 and follow the general instructions given there.
- Press one of the NEXT keys to move the cursor to the next column to the right.
- Use the UP or DOWN keys to select the desired language.

Note: Once a new language is selected, all text in the Setup menu (and in all other menus and screens) will be displayed in that language. This may cause difficulties if an unknown language is selected.

Selecting Patient Mode

Caution: Prior to patient monitoring, ensure the monitor is configured to the appropriate Patient mode.

To select the Patient mode, proceed as follows:

- Enter the Setup menu as described on page 75 and follow the general instructions given there.
- Press the DOWN key until the parameter "Patient" is selected.
- Press one of the NEXT keys to move the cursor to the next column to the right.
- Use the UP or DOWN keys to select Adult or Neonate mode.

Or alternatively:

- Keep the UP keys pressed when the monitor is turned on to set the monitor to Adult mode.
- Keep the DOWN keys pressed when the monitor is turned on to set the monitor to Neonate mode.

The Patient mode is being displayed in the Equipment Message Window in the lower left area of the Main screen.

Note: CO₂, and NIBP functions are affected by selecting Adult or Neonatal operating modes. The factory default is the Adult mode.

Configure Audio Silencing

The audio signal associated with patient alarms can be silenced by pressing the ALARM key. The functionality of this key depends on the selected configuration:

To configure the monitor's silencing function, proceed as follows:

- Enter the Setup menu as described on page 75 and follow the general instructions given there.
- Press the DOWN key until the parameter "Audio Silence" is selected.
- Press one of the NEXT keys to move the cursor to the next column to the right.
- Use the UP or DOWN keys to select the desired option.

Select "2 Minute" when the silence period should automatically terminate after 2 minutes.

Select "Permanent" when the silence period should continue permanently until it is terminated manually.



Warning: In Permanent Silence mode the user is responsible to reactivating (enabling) the audio alarm signal by pressing the ALARMS key again.

Setting the Date

To set the monitor's Date function, proceed as follows:

- Enter the Setup menu as described on page 75 and follow the general instructions given there.
- Press the DOWN key until the parameter "Date" is selected.
- Press one of the NEXT keys to move the cursor to Day-Month-Year.
- Use the UP or DOWN keys to increase or decrease the values for Day-Month-Year.

Note: Altering the monitors' date will affect the History data.

Setting the Time

To set the monitor's Time function, proceed as follows:

- Enter the Setup menu as described on page 75 and follow the general instructions given there.
- Press the DOWN key until the parameter "Time" is selected.
- Press one of the NEXT keys to move the cursor to Hours : Minutes.
- Use the UP or DOWN keys to increase or decrease the values for hours or minutes.

Note: Altering the monitors' time will affect the History data.

Daylight Saving Time Option

To set the monitor's Daylight Saving Time function, proceed as follows:

- Enter the Setup menu as described on page 75 and follow the general instructions given there.
- Press the DOWN key until the parameter "DST" is selected.
- Press one of the NEXT keys to move the cursor to the next column to the right.
- Use the UP or DOWN keys to select one of the Daylight Saving Time options.

The possible options have the following meaning:

OFF: Daylight Saving Time is not activated. The monitor will not change the time automatically. It is up to the user for changing the time if needed. This is the monitor's default setting.

N AMERICA: Use this setting and the monitor will automatically change time accordingly to the Daylight Saving Time for countries in North America.

EU 01:00: Daylight Saving Time for all European countries using Greenwich Mean Time.

EU 02:00: Daylight Saving Time for all European countries 1 hour ahead of Greenwich Mean Time.

EU 03:00: Daylight Saving Time for all European countries 2 hours ahead of Greenwich Mean Time.

Note: Enabling Daylight Saving Time will affect the History data.

Selecting Alarm Delay

To change the SpO₂ Alarm Delay setting, proceed as follows:

- Enter the Setup menu as described on page 75 and follow the general instructions given there.
- Press the DOWN key until the parameter “O₂ Alarm Delay” is selected.
- Press one of the NEXT keys to move the cursor to the next column to the right.
- Use the UP or DOWN keys to select 10 Sec or 0 Sec for the delay.

Selecting CO₂ Units

To select the display unit for CO₂ waveform and EtCO₂ values, proceed as follows:

- Enter the Setup menu as described on page 75 and follow the general instructions given there.
- Press the DOWN key until the parameter “EtCO₂ Unit” is selected.
- Press one of the NEXT keys to move the cursor to the next column to the right.
- Use the UP or DOWN keys to select one of the options (mmHg, kPa or %).

Selecting Display Background

To select the display background, proceed as follows:

- Enter the Setup menu as described on page 75 and follow the general instructions given there.
- Press the DOWN key until the parameter “Background” is selected.
- Press one of the NEXT keys to move the cursor to the next column to the right.
- Use the UP or DOWN keys to select Dark or Light.

8 Cleaning



Warning: Do not, under any circumstances, perform any cleaning while the monitor is being used to monitor a patient. The monitor must be turned off. Unplug the monitor from the AC or DC power source and disconnect all accessories.



Caution: Do not open the monitor to clean it.



Caution: Do not immerse any part of the electrical connector of the cable or accessories in the cleaning or disinfection solution at any time. This may cause internal damage and reduce the product life.



Caution: Do not use abrasive cleaners, isopropyl alcohol or organic solvent for cleaning. Use of these cleaners can cause damage, stiffness and brittleness to the monitors' surface and to cables and wires.

Cleaning the Monitor



Caution: Disconnect all accessories from the monitor before cleaning.

Examine the monitor's case for damages and check the AC power cord for bent or broken prongs, cracks or fraying. Neither the monitor nor the power cord should be used if damaged. If any damage is noted, contact the appropriate service personnel.



Caution: Do not spray or pour any water or cleaning solution directly onto the monitor.

- **Housing:** As needed, clean the monitor using a soft cloth dampened with a mild dishwashing detergent solution and gently rub the soiled area until clean. Use a clean soft cloth to dry the monitor.
- **Display:** Clean the display window using a soft, lint-free cloth sprayed with an alcohol free glass cleaner. The use of paper towels is not recommended as it may scratch the surface.
- **Disinfections:** The monitor surfaces may be disinfected using a soft cloth saturated with a 10% (1:10) solution of chlorine bleach in tap water. When all of the surfaces have been disinfected, wipe the entire surface of the monitor using a soft cloth dampened with fresh water to remove any trace amounts of residue and/or fumes.

Note: Thoroughly wipe off any excess cleaning solutions. Care should be taken to prevent water or cleaning solution to run into connector openings or crevices.

Cleaning Cuffs and Pneumatic Hoses

Prior to each patient use, inspect the blood pressure cuff and the pneumatic hose for damage.

- **Tuff-Cuff:** Clean the cuff using a soft cloth dampened with a 70% Isopropyl Alcohol solution. The cuff should be allowed to thoroughly dry before use.
- **Safe-Cuff:** Clean the cuff using a soft cloth dampened with a soap, water-based detergent or chlorinated disinfectant solution. Do not use alcohol.
- **Pedisphyg Cuffs:** A water-based detergent is suitable for wiping the cuff. A damp, detergent-free cloth should then be used to rinse the cuff.
- **Pneumatic Hose:** Clean the pneumatic tubing using a soft cloth dampened with a germicidal solution.

Note: It is not recommend submersing the cuff or hose. Liquid should not be permitted to enter the cuff bladder or hose because instrument damage may occur. Cuffs and hoses should be allowed to thoroughly dry before use.

Cleaning SpO₂ Sensors



Caution: Do not soak or immerse the sensor or its cable in any liquid solution. Do not attempt to sterilize it.

Refer to the Directions For Use pamphlet enclosed with each sensor for more information.

- Clean the SpO₂ sensors with a soft cloth dampened with 70% Isopropyl alcohol solution. Allow the sensor to dry prior to placement on a patient.

Cleaning CO₂ Consumables

Microstream CO₂ consumables are designed for single patient use and are not to be reprocessed.



Caution: Do not attempt to disinfect or flush the FilterLine as the monitor can be damaged.

Cleaning the Printer



Refer to the printer User's Manual for more information.

Caution: Before cleaning the printer, disconnect the AC adapter.

- Wipe with a soft dry cloth.

To remove extreme dirt buildup:

- Soak a cloth with mild detergent, wring well and wipe. Dry by wiping with a soft dry cloth.



Caution: Do not use volatile chemicals such as thinner, benzine, etc.



Caution: Never wet the inside of the printer mechanism.

9 Maintenance



Warning: Do not, under any circumstances, perform any testing or maintenance on the monitor while the monitor is being used to monitor a patient. The monitor must be turned off. Unplug the monitor from the AC or DC power source and remove the internal battery.



Caution: Service performed by unauthorized personnel could be damaging to the monitor and may void the warranty. For service, contact your CAS Representative.

Maintenance Intervals

Preventive maintenance of the monitor is an important function that should be performed routinely to ensure safe and efficient monitor operation. The following maintenance intervals are recommended:

- **Battery Pack:** Replace the battery every 2 years.
- **SpO₂:** No user calibration is required. Perform a SpO₂ Simulator Check once per year or when there is doubt about the validity of the SpO₂ readings.
- **NIBP:** Perform a Pneumatic Pressure check every 6 months and a NIBP Calibration Check once a year or when there is doubt about the validity of the pressure readings.
- **CO₂:** Perform a CO₂ Calibration Check once per year, after 4000 operating hours (whichever comes first) or when there is doubt about the validity of the CO₂ readings. The monitor should be returned to CAS Medical Systems for periodic maintenance every 14,000 operating hours.

Maintenance Checks

Entering the Service Menu



Warning: Do not enter the SERVICE menu when the monitor is connected to a patient.

To enter the monitor's SERVICE menu, proceed as follows:

- Press and hold the PARAMETERS and HISTORY keys while the monitor is being turned on.

<u>Service</u>	
IrDA Test	OFF
EtCO ₂ Check	OFF
EtCO ₂ Cal	OFF
EtCO ₂ CAL Date	06-Feb-04
EtCO ₂ S/N	00234
Service EtCO ₂	xxxxx Hours
Cal EtCO ₂	xxxx Hours
Manometer Mode	OFF
Pneumatic Test	OFF
PIC Voltage	xx.xx v

Figure 17: Service Menu

The cursor is initially positioned at the first item in the left column.

- Use the UP or DOWN keys to select any other item.
- Press one of the NEXT keys to move the cursor to the next column to the right.
- Use the UP or DOWN keys to scroll through the available options.

Exiting the Service Menu

- Press the RETURN key to exit the menu and return to the Main screen.

Note: If no key is pressed for 15 minutes, the monitor will automatically return to the Main screen.

IrDA Test

The IrDA Test checks the alignment of the monitor and the printer to guarantee communication between their infrared windows.

To use the Infrared Test, proceed as follows:

- Enter the Service menu as described on page 86 and follow the general instructions given there.
- Change the “IrDA Test” setting from “OFF” to “ON”.

The message “Printing” will be displayed in the Equipment Message Window. When the Infrared windows are properly positioned, the printer will begin to print a series of “A” characters. The test will continue for 60 seconds or can be terminated by the user.

- Press one of the PREVIOUS keys to cancel the test and return to the IrDA Test menu selection.

CO₂ Calibration Check

The CO₂ Calibration Check is a tool to verify proper calibration of the CO₂ parameter.



Caution: Do not perform a CO₂ Calibration Check when the monitor is in normal measuring mode. This mode corrects the CO₂ value for BTPS (Body, Temperature, Pressure, Saturation), which assumes that alveolar gases are saturated with water vapor. The Calibration Check mode disables this correction.



Caution: The Calibration Check must be performed with a manufacturer authorized Calibration Kit containing (5% CO₂, 21% O₂, Balance N₂) gas, tubing adapter and a calibration filter line. A manufacturer approved Calibration Kit can be purchased from Scott Medical (P/N 0304653ORFBD).

Note: The Calibration Check should be performed only after the CO₂ module has been operating for at least 20 minutes in a normal operating mode.

To check CO₂ Calibration, proceed as follows:

- Verify that the FilterLine supplied with the Calibration kit is firmly attached to the gas canister.
- Enter the Service menu as described on page 86 and follow the general instructions given there.
- Press the DOWN key until the parameter “EtCO₂ Check” is selected.
- Press one of the NEXT keys to select the “OFF” setting.
- Change the EtCO₂ Check setting from “OFF” to “Connect Gas”.
- Connect the FilterLine to the monitor.
- The highlighted section should read “Open 5% Gas”.
- When the message “Open 5% Gas” appears, press and hold open for 15 seconds the gas valve until the reading stabilizes.
- Verify the reading on the display to be 5.0% ± 0.3 vol%.

Note: Calibration is not required if the measured value are the same as the concentration of the calibration gas ± 0.3 vol%.

- Press one of the PREVIOUS keys to cancel the test and return to the EtCO₂ Check menu selection.
- Disconnect the FilterLine from the monitor.

Should the monitor fail the EtCO₂ Check, it is recommended to perform an EtCO₂ Calibration or returning the monitor for service.

CO₂ Calibration



Note: Perform the Calibration only after performing the Calibration Check.

Caution: The Calibration must be performed with a manufacturer authorized Calibration Kit containing (5% CO₂, 21% O₂, Balance N₂) gas, tubing adapter and a calibration FilterLine. A manufacturer approved Calibration Kit can be purchased from Scott Medical (P/N 0304653ORFBD).

To perform a CO₂ Calibration, proceed as follows:

- Verify that the FilterLine supplied with the Calibration Kit is firmly attached to the gas canister.
- Enter the Service menu as described on page 86 and follow the general instructions given there.
- Press the DOWN key until the parameter “EtCO₂ Cal” is selected.
- Press one of the NEXT keys to select the “OFF” setting.
- Change the EtCO₂ Cal setting from “OFF” to “Connect Gas”.
- Connect the FilterLine to the monitor.
- The highlighted section should read “Open 5% Gas”.
- When the message “Open 5% Gas” appears, press and hold the gas valve until the message “Remove Gas” is displayed.
- The highlighted section should read “Passed” and two audible beeps should be heard when the calibration is completed.
- Verify the “EtCO₂ CAL Date” and the “Cal EtCO₂ Hours” have been updated.

The starting hours for Calibration are initially 1200 hours, and then 4000 after that. This timer is automatically reset to 4000 hours during the Calibration process as long as it has been more than 720 hours from the last calibration. If this was less than 720 hours, the timer does not reset.

- Press one of the PREVIOUS keys to cancel the test and return to the EtCO₂ Cal menu selection.
- Disconnect the FilterLine from the monitor.

Note: Should the monitor fail the EtCO₂ Calibration, it is recommended the procedure be repeated. Prior to repeating the Calibration procedure, carefully check all connections.

Should the monitor continue not to pass the Calibration procedure, the monitor should be returned for service.

NIBP Checks

To guarantee correct NIBP measurements, it is recommended to perform the following checks at least once per year. All checks can be done in one session beginning with the Pneumatic Pressure Check, then continue with the Calibration Check and finally perform the Over Pressure Test.

Note: The monitor must be in the Adult mode to perform the following pressure checks.

The following equipment is required to perform the NIBP pressure checks:

- A Calibration Kit (P9) which is included with the monitor.
- A Mercury manometer or equivalent whose accuracy meets the AAMI/ANSI Standard for Non-Automated Sphygmomanometers, 2002.



Warning: Do not activate the Service menu when a cuff is attached to a patient.

Pneumatic Pressure Check

To perform the Pneumatic Pressure Check, proceed as follows:

- Connect the NIBP inflation hose to the monitor without a cuff connected to the hose.
- Obtain the male luer plug from the Calibration Kit and use it to close the open end of the NIBP inflation hose. Twist the plug one-quarter turn to make sure the connection is not leaking. This is essential for the test to be performed.
- Enter the Service menu as described on page 86 and follow the general instructions given there.
- Press the DOWN key until the parameter “Pneumatic Test” is selected.
- Press one of the NEXT keys to move the cursor and begin the test.

The monitor starts to pressurize the pneumatic system to approximately 180 mmHg and attempts to hold this pressure. The pressure value will be displayed for about 15 seconds.

At the completion of a successful Pressure Check, the message “Passed” will display and the monitor will beep two times.

If the monitor fails the Pressure Check, the message "Failed" will be displayed and the monitor will beep three times.

- Press one of the PREVIOUS keys to exit and return to the Pneumatic Test selection.



Caution: Should the monitor fail the Pneumatic Pressure Check the test should be repeated. Instead of blocking the inflation hose with the male Luer plug, a 500 ml volume should be connected. (Obtain a fixed volume 500 ml Pressure Cylinder from CAS, part number 01-02-0248)

- Remove the LUER plug from the end of the air hose and connect the 500 ml Pressure Cylinder instead.

Should the monitor fail the Pneumatic Pressure Check with the 500 ml attached, it needs to be serviced.

Calibration Check and Overpressure Test

To perform a Calibration Check, proceed as follows:

- Connect the NIBP inflation hose to the monitor without a cuff connected to the hose.
- Enter the Service menu as described on page 86 and follow the general instructions given there.
- Press the DOWN key until the parameter "Manometer Mode" is selected.
- Press one of the NEXT keys to move the cursor and begin the test.

The pressure in the pneumatic system is being displayed. At this point it is 0 mmHg.

- Use the manometer inflation bulb to slowly inflate the system pausing for 30 seconds at the following points and verify calibration according to the following table:

0 mmHg:	± 1 mmHg
50 mmHg:	± 4 mmHg
100 mmHg:	± 4 mmHg
150 mmHg:	± 4 mmHg
200 mmHg:	± 5 mmHg

To test the Overpressure safety function, proceed as follows:

- Inflate the pressure slowly until 290 mmHg \pm 10 mmHg is reached. The monitor's NIBP display should stop updating and the message "OVER PRESS" will be displayed.

Note: The monitor needs to be serviced if it does not meet the test criteria.

PIC Voltage

The monitor displays the DC voltage level being received by the PIC processor from the Power Supply Board.

To view the PIC Voltage level, proceed as follows:

- Connect the monitor to an external power source (AC Line power).
- Enter the Service menu as described on page 86 and follow the general instructions given there.

Verify the voltage being displayed is 12.00V \pm 0.50V. If this is not the case, the monitor needs to be serviced.

Battery Replacement

Removing the Battery

- Turn the monitor off and disconnect the power cord.
- Push down on the battery latch to unlock the battery door from the rear panel of the monitor.
- Carefully remove the battery pack from the rear panel of the monitor.

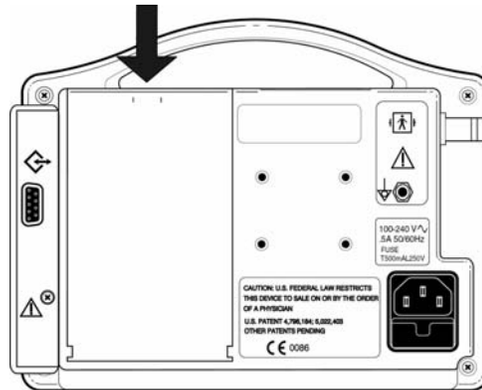


Figure 18: Removing the Battery Pack

Inserting the Battery

- Align the Battery Pack guides with the bottom of the monitor.
- Slowly close the battery door to ensure the connector in the monitor and the connector on the battery pack mate together.
- Lock the battery door closed.

Note: When the battery pack is re-inserted, the monitor will automatically turn on.



Warning: Use of unapproved batteries will invalidate the product's warranty and may result in serious safety consequences for the patient and user.



Warning: Do not disassemble the battery pack or batteries. The batteries contain electrolytes, which can cause injury to eyes, skin and clothing.

Note: This product contains a rechargeable battery that is recyclable. Under various state and local laws, it may be illegal to dispose of this battery into the municipal waste stream. Check with your local authorities for instructions on recycling options in your area.

Fuse Replacement

Note: The **CAS 750CM** monitor models have a single fuse located inside the monitor. This fuse is not user replaceable.

The **CAS 750C** monitor has a dual fuse power input receptacle. Both AC lines are fused.



Caution: For continued protection against fire hazard, replace only with identically rated fuses. Refer to paragraph "Power" on page 120.

A fuse may need to be replaced if the monitor is plugged into an electrical outlet but the Battery Power Visual Indicator is not illuminated green.



Warning: Before changing the fuse, unplug the power cord.

To replace fuses, proceed as follows:

- Turn the monitor off and disconnect the power cord.
- Press down on the locking tab, which holds the fuse holder in the power input receptacle. While holding down on the tab, pull the fuse holder out.
- Remove the fuses.
- Place new fuses directly into the fuse holder.
- Insert the fuse holder into the power input receptacle. There should be an audible click when it is secured.

Software Versions

The monitor displays the current software revision of its operating system and that of the internal modules being used inside.

To view the software versions, proceed as follows:

- Enter the monitor's Service menu as described on page 86.
- Press and hold the HISTORY key for 2 seconds to display the software versions.

<u>Versions</u>	
DSP Software	2.2
Lang Flash	2.2
BOOT	2.1
PIC	1.0
Masimo SpO ₂ ⁽¹⁾	1.1
Oridion EtCO ₂	2.00
CAS ND ⁽²⁾	1.0
Unit S/N	0503123

Figure 19: Software Versions

(1) Installed SpO₂ technology may be Masimo SpO₂ or Nellcor SpO₂.

(2) The NIBP module is optional, in the case when it is not installed, no text message is shown.

- Press the RETURN key to exit and return to the Main screen.

Note: If no key is pressed for 60 seconds, the monitor will automatically exit the Versions menu and return to the Main screen.

Storage

Refer to paragraph “Operating Environment” on page 119 for storage temperature information.



Warning: If it becomes necessary to store the monitor for longer than 6 months, remove the monitor’s battery pack and place the monitor in its original packing container if available.

Note: Batteries not charged and left in storage for more than six months could degrade and not recharge to full capacity.

10 Printer

For more detailed information on the Citizen Model CMP-10 Mobile Printer, refer to the User's Manual that was supplied with the printer.



Warning: The CAS 750C Monitor has been tested with the Citizen CMP-10 Mobile printer to comply with IEC 60601-1-1 and is the only printer that is recommended to be used with the monitor. If another printer is to be used, the user must read the Caution "*Leakage Current Test*" on page 15 and follow the guidance given.

The Citizen CMP-10 Mobile Printer interfaces to the monitor via an Infrared Port or by using the direct connect RS232 cable (supplied with printer). The Infrared ports are located on the top of the printer and on the bottom front panel of the monitor.



Caution: For safe and proper usage of the external printer, please observe the following:

- Avoid places where fluid may enter the printer opening accidentally.
- Avoid places subject to high or low temperature extremes.
- Avoid dusty places and where corrosive gasses are generated.
- Never attempt to dismantle or repair the printer mechanism.
- Do not drop or bump the printer.

When handling the thermal paper:

- Store in a dark, cool and dry place.
- Do not place near organic solvents.
- Avoid contact with vinyl chloride films erasers or adhesive tapes for extended periods.
- Avoid exposure to high temperature, humidity, liquid, or sunlight.
- Always use specified thermal paper. Refer to "Other Accessories and Options" on page 127 for order information.

Printer Controls and Indicators

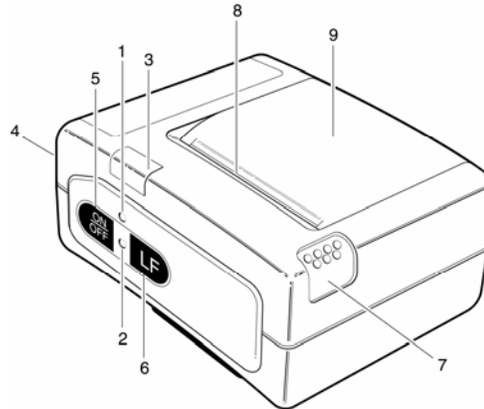


Figure 20: Printer Controls and Indicators

1. Charge LED (red = charging, green = fully charged)
2. Power LED (green = power on, red/green blinking fast = end of paper, red/green blinking slow = print head overheated)
3. Infrared (IrDA) port
4. RS 232 serial interface
5. Power ON/OFF key
6. Line Feed key
7. Paper Cover Release Button (press down to open cover)
8. Paper Cutting Edge
9. Paper Cover

Printer Operation

Direct Connection

The monitor uses the DB9 connector, located on the rear panel of the monitor, to interface to the RS232 port on the Citizen CMP-10 Mobile Printer. Use the cable, which is supplied with the printer.

Note: When using the direct connect method; connect the printer's serial cable to the printer before turning the printer on.

Infrared Connection

Note: When using the infrared port, disconnect the printer serial cable from the printer before turning the printer on.

Position the printer's infrared port window in-line with the infrared window of the monitor.

Note: When using the Infrared port, it is important to keep the two devices close together (less than three feet / one meter) and in-line to maintain proper communications. The monitor can perform a communication test to verify the Infrared connection. Refer to paragraph "IrDA Test" on page 87.

Note: When using the IrDA port for printing, it is recommended to keep all items clear of the communications path between the monitor and the printer.

- To turn the printer on, press and hold the ON/OFF key for 1 second. The Power LED illuminates green.
- To turn the printer off, press and hold the ON/OFF key. The Power LED will illuminate red and change back to green. When the LED illuminates green, remove your finger from the key.

Note: If the monitor is turned on and the printer is already on, the CAS logo is printed automatically.



The CMP-10 Mobile Printer has an Auto Power Off feature. If the monitor and printer become separated and during 10 minutes no data has been sent and the Line Feed (LF) key has not been pressed, the printer will automatically shutoff.

The monitor periodically sends a "wake-up" message to the printer that will disable the Auto Power Off feature.

Sample printouts of both, Trend and Alarm Histories, are shown below.

Header	<pre> 750X Series Monitor 27-Feb-04 15:48 Patient: _____ Notes: _____ Trend Hist: HR:MN %O2 PR CO2 RR 13:35 98 59 38 16 13:35 NIBP=120/80 (90) PR= 59 13:36 98 59 38 16 13:37 98 59 38 16 13:37 NIBP=112/78 (89) PR= 61 13:38 98 59 38 16 13:39* 98 59 38 16 13:39 NIBP=120/80 (90) PR= 58 13:40 98 59 38 16 </pre>	<pre> 750X Series Monitor 27-Feb-04 15:48 Patient: _____ Notes: _____ Alarm Hist: HR:MN Alarm 13:39 High %SpO2: 98% </pre>

Figure 21: History Sample Printouts

Note: The header information is printed each time the monitor is turned on.

Note: An asterisk (*) appears in the History printout to indicate an Alarm has occurred during that one-minute interval.

To print waveforms:

- Press and hold the RETURN key for 2 seconds while in the Main display screen.

The waveforms being displayed will print at the speed as previously selected. Refer to paragraph "Parameters Menu" on page 38.

When the Print On Alarm setting in the Parameters menu is set to "ON", the type of alarm is printed on the printout and the trace is annotated with an arrow marker to mark the point of the alarm.

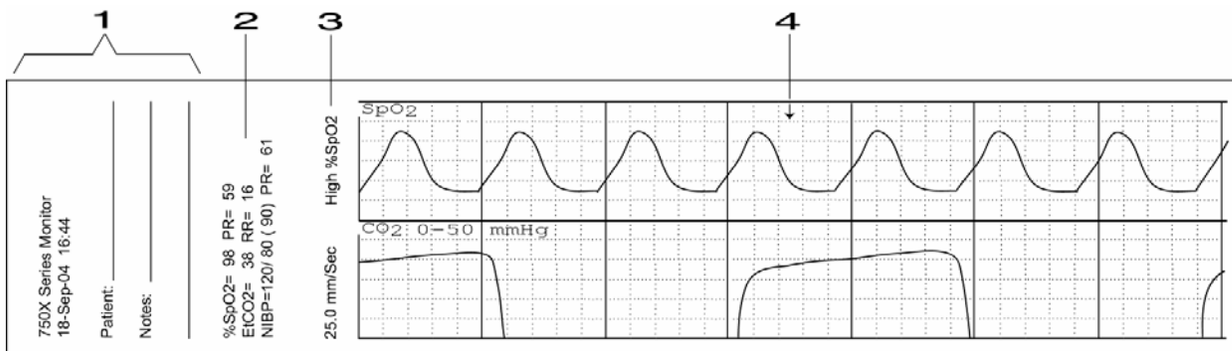


Figure 22: Waveform Sample Printout

1. Header
2. Numeric Information
3. Print Speed and Alarm Event Type (only printed when Print-on-Alarm is activated)
4. Alarm Event Marker

Charging the Printer Battery

The CMP-10 Mobile Printer is equipped with a rechargeable Lithium Ion (LiION) battery pack. When the printer detects a Low Battery condition, the message “Low Battery” is printed and an audio indicator sounds three times.



Warning: Charge the printer battery using the AC Adapter, Model TRC-09-1100-M from GROUP WEST, or equivalent, included with the printer.

- Plug the battery charger’s cord into the printer battery charger jack, located on the rear panel.
- Plug the charger into an AC wall outlet of the appropriate voltage.
- Verify the Charge LED is lit red.

Battery charge time is approximately 3 hours. Once the battery is fully charged, the Charge LED switches to green.

Installing Paper

Note: A red line appears when the remaining supply of thermal paper becomes low.

- Switch the printer off.
- Press the Cover Open button to access the paper compartment. Remove any remaining paper before installing the new roll.
- Place the new paper roll as shown on the illustration and pull out enough paper to reach out over the control panel of the printer.
- Close the paper door.

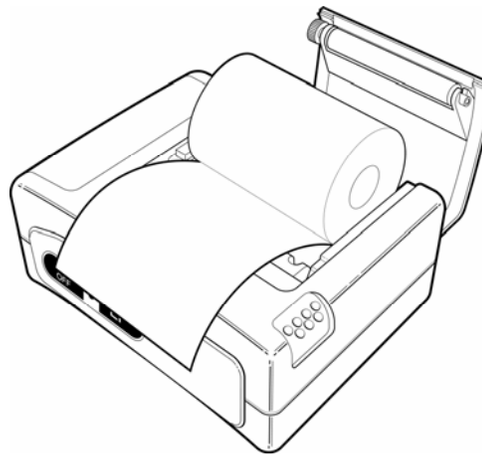


Figure 23: Paper Installation

Note: Make sure that the paper is correctly placed. If it is tilted in one or another direction and does not come out straight from under the cover, open the door and reposition the roll again.



Warning: Do not touch the print head or paper cutter while replacing the printer paper.

Removing the Battery Pack



Warning: Do not operate the printer or connect it to the monitor with the battery pack is removed.



Warning: Never change the battery pack while the battery charger is plugged in and/or the monitor is being operated.

- Switch the printer off.
- Disconnect the printer from the monitor and unplug the wall charger cord.
- Open the battery door by pressing in on the battery cover and pushing upward.

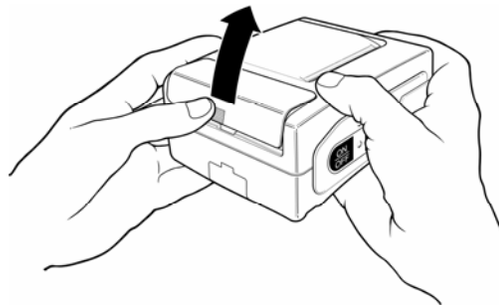


Figure 24: Opening the Battery Door

- Remove the battery cover.
- Remove the battery pack from the compartment and disconnect its connecting cable.

Installing the new Battery Pack:

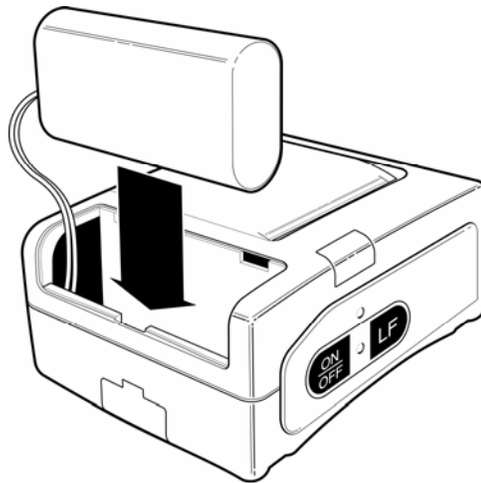


Figure 25: Installing the New Battery

- Connect the battery cable into the battery connector.
- Insert the battery and its connecting cable into the battery compartment.
- Replace the battery cover by sliding it in from the back of the printer and pushing down to lock it in place.



Caution: Be sure to place the battery cover firmly in its position after installing the new battery pack.



Warning: Do not disassemble the battery pack or batteries. The batteries contain electrolytes, which can cause injury to eyes, skin and clothing.

Note: Under various state and local laws, it may be illegal to dispose of this battery into the municipal waste stream. Check with your local authorities for instructions on recycling options in your area.

11 External Device Interfacing

RS232 Interface

The monitor uses the DB9 connector to interface to the RS232 port on the Citizen CMP-10 Mobile Printer. Use the cable, which is supplied with the printer.

The RS232 interface may be used for data collection or service purposes. Refer to the documentation provided with each application.

Note: The CAS 750C Monitor is not equipped with interfaces to remote equipment or network(s) to duplicate alarms.

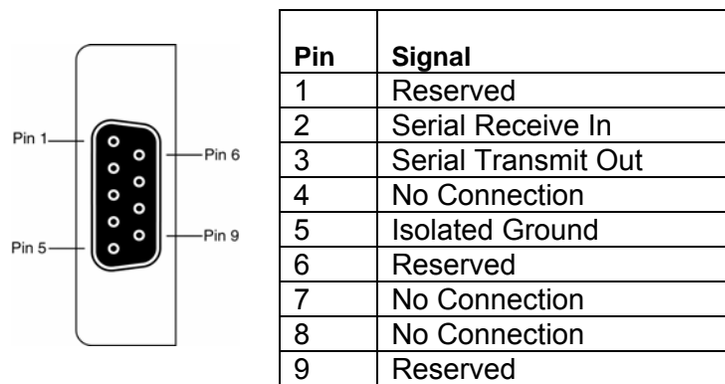


Figure 26: DB9 Connector

Mounting

There are several mounting devices (e.g. roll stand or swivel mount) available for the monitor. In all cases, the 4 screw threads at the back are used to attach the monitor to the mounting device.

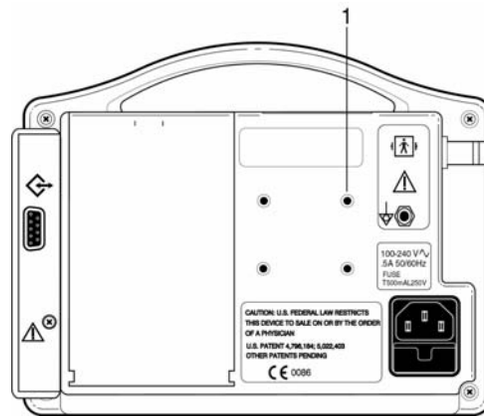


Figure 27: Mounting Threads

1. Screw Thread (one of four)

Note: Follow the instructions provided with each specific mounting solution.

12 Appendix

Symbols

The following is a summary of all symbols used on the monitor and accessories. Symbols may occur on the product or on its packaging.

Front Panel Symbols



Power Indicator.



POWER (On/Standby)



ALARM (Silence/Reset)



NIBP (Start, Stop, Menu)



RETURN (Main Screen, Freeze, Print)



PARAMETER



LIMITS



HISTORY



VOLUME (Audio/Visual)



UP



DOWN

Screen Indicators



Heart Beat Indicator



Respiration Breath Indicator



Battery Charge Level Indicator



Silence Indicator



Bar Graph Indicator

Symbols near Accessory Connections



Patient connections are Type BF



Communication Port RS232 Connector

SpO₂

Pulse Oximeter Probe Input Connector

MicroStream™ CO₂ Input Connector

CO₂ Scavenger Exhaust Port

MAXNIBP®

NIBP Hose and Cuff Connector

Symbols on Monitor or Printer



CAUTION, read instructions before using.



Potential Equalization Post

IPX1

Protection against ingress of water.



The CE Mark and Notified Body Registration Number signify the device has met all essential requirements of European Medical Device Directive 93/42/EEC.



Alternating Current



Symbol for DC power polarity



Symbol for DC power



Recycling suggested



Class II equipment



Read instructions in Printer Manual

Symbols on Packaging



Relative Humidity for Storage and Transport



Storage and Transport Temperatures

Warranty Policy

MONITORS (CAS 750C)

CAS Medical Systems, Inc. warrants the monitor, when new, to be free from defects in material and workmanship and to perform in accordance with manufacturer's specifications for a period of two (2) years from the date of original purchase from CAS or its authorized distributors or agents except as noted below.

The same warranty conditions are made for a period of one (1) year with respect to printer and battery and ninety (90) days on non-disposable accessories and certain components consisting of reusable SpO₂ sensors and other accessories provided by CAS as part of the original purchase. CAS warrants blood pressure cuffs and disposable or single-patient-use products for out-of-box failure only. Where the accessory is not a CAS manufactured product, the manufacturers own warranty conditions apply.

CAS reserves the right to perform warranty service operations in its own factory, at an authorized repair facility, or at the customers' site.

Our obligation under this warranty is limited to repairing or, at our option, replacing any defective parts or our equipment, without charge, if such defects occur in normal service and with prompt notification.

Damage to any part through misuse, neglect, or accident, or by affixing any accessories or attachments other than CAS, Masimo[®], Nellcor[®], and Oridion[®] manufactured accessories or attachments, is not covered by this warranty.

ACCESSORIES, BATTERIES, CUFFS, AND CERTAIN COMPONENTS

In all cases, policy applies from date of purchase from CAS or its authorized distributors or agents.

Batteries:	(1) Year
Chargers:	(1) Year (not including power cord: see other accessories).
Cuffs (all):	Out-of-box failure only.
SpO ₂ Sensors:	(90) Days for Masimo and Nellcor SpO ₂ Sensors.
CO ₂ Accessories:	Out-of-box failure only for Oridion Sample Lines.
Other Accessories:	Out-of-box failure only.
Certain Components:	(1) Year - Printer mechanism, but not including Thermal Print Heads.
Print Heads:	Out-of-box failure only.

THERE ARE NO WARRANTIES, WHICH EXTEND BEYOND THOSE EXPRESSLY DESCRIBED IN THIS AGREEMENT AND THE COMPANY MAKES NO WARRANTIES OF MERCHANTABILITY OR FITNESS FOR A PARTICULAR PURPOSE.

Monitor Error Messages

Clock Battery

The monitor's internal clock battery is almost discharged.

- The monitor needs to be serviced.

Dead Battery

The battery is fully discharged.

- Recharge the battery for at least 5 hours.

Low Battery

The battery is almost discharged. At least 30 minutes of operation is available from when the message first appears.

- Recharge the battery as soon as possible.

System Error

A failure has occurred on the Main Board.

- The monitor needs to be serviced.

Power Failure

Power was disconnected from the monitor.

- Press the ALARM key to clear the message.

OR

- Turn the monitor off and on again.

SET CLOCK

Time and date values are incorrect.

- Set the monitor's clock.
- If the condition persists after setting the monitors clock, then the monitor's internal clock battery needs to be replaced.

Monitor Configurations

CAS 750C Models with AC Power Supply

CAS 750C-2MS	Oridion CO ₂ , Masimo SpO ₂
CAS 750C-2NL	Oridion CO ₂ , Nellcor SpO ₂
CAS 750C-3MS	Oridion CO ₂ , Masimo SpO ₂ , MAXNIBP®
CAS 750C-3NL	Oridion CO ₂ , Nellcor SpO ₂ , MAXNIBP®

Table 2: CAS 750C Models with AC Power

CAS 750C Models with 12 VDC Power Input

CAS 750CM-2MS	Oridion CO ₂ , Masimo SpO ₂
CAS 750CM-2NL	Oridion CO ₂ , Nellcor SpO ₂
CAS 750CM-3MS	Oridion CO ₂ , Masimo SpO ₂ , MAXNIBP®
CAS 750CM-3NL	Oridion CO ₂ , Nellcor SpO ₂ , MAXNIBP®

Table 3: CAS 750C Models with DC Power

Monitor Configuration Record

Monitor Model

CAS 750C

CAS 750CM.....

Serial Number:

Installed Option Parameter

NIBP

Main Screen Defaults

Channel 1 : SpO₂

Channel 2 : CO₂

Channel 3 :

Alarm Limit Defaults

Parameters	Adult		Neonate	
	High	Low	High	Low
%SpO ₂				
Pulse Rate				
CO ₂				
Resp. Rate				
No Resp.				
Systolic				
Diastolic				
MAP				

Date: _____ Unit: _____

Specifications

Oximetry

Masimo SET®

Type:	Functional Oxygen Saturation																																
Wavelengths:	Red: 660 Nanometers Infrared: 905 Nanometers																																
Radiant Power:	0.79mW maximum at 50 mA, pulsed																																
Measurement Range:	%SpO ₂ : 0-100% Pulse Rate: 25-240 BPM																																
Accuracy:	%SpO ₂ : <table> <thead> <tr> <th>Sensor</th> <th>Accuracy</th> </tr> </thead> <tbody> <tr> <td>DC-195</td> <td>70 - 100%, +/-2 digits (1 S.D.)</td> </tr> <tr> <td>LNOP® Adt</td> <td></td> </tr> <tr> <td>LNOP Adt Long</td> <td></td> </tr> <tr> <td>LNOP DCI</td> <td></td> </tr> <tr> <td>LNOP DCSC</td> <td></td> </tr> <tr> <td>LNOP DC1P</td> <td></td> </tr> <tr> <td>LNOP DC150</td> <td></td> </tr> <tr> <td>LNOP Ear</td> <td></td> </tr> <tr> <td>LNOP Pdt</td> <td></td> </tr> <tr> <th>Sensor</th> <th>Accuracy</th> </tr> <tr> <td>LNOP Neo</td> <td>70 - 100%, +/-3 digits (1 S.D.)</td> </tr> <tr> <td>LNOP Neo PT</td> <td></td> </tr> <tr> <td>LNOP NeoPT-L</td> <td></td> </tr> <tr> <td>LNOP Neo-L</td> <td></td> </tr> <tr> <td>LNOP Inf-L</td> <td></td> </tr> </tbody> </table>	Sensor	Accuracy	DC-195	70 - 100%, +/-2 digits (1 S.D.)	LNOP® Adt		LNOP Adt Long		LNOP DCI		LNOP DCSC		LNOP DC1P		LNOP DC150		LNOP Ear		LNOP Pdt		Sensor	Accuracy	LNOP Neo	70 - 100%, +/-3 digits (1 S.D.)	LNOP Neo PT		LNOP NeoPT-L		LNOP Neo-L		LNOP Inf-L	
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LNOP Neo-L																																	
LNOP Inf-L																																	
	Pulse Rate: ± 3 BPM																																
Update of Numeric:	every 1 second																																
Waveform Display:	Plethysmogram, fixed for trace 1																																
Sweep Speeds:	12.5, 25 or 50 mm/sec																																
Patient Alarms:	High and low limits on %SpO ₂ , adjustable. High and low limits on Pulse Rate, adjustable.																																
Alarm Delay:	0 or 10 seconds																																

Nellcor® OxiMax®

Type: Functional Oxygen Saturation

Wavelengths:

Red: 660 Nanometers

Infrared: 890 Nanometers

Power: Not exceeding 15 mW

Measurement Range:

%SpO₂: 1-100%

Pulse Rate: 20-240 BPM

Accuracy:

<u>%SpO₂</u>	<u>Sensor</u>	<u>Accuracy</u>
	MAX-A	70 - 100%, +/-2 digits (1 S.D.)
	MAX-AL	
	MAX-N	
	MAX-P	
	MAX-I	
	MAX-FAST	
	<u>Sensor</u>	<u>Accuracy</u>
	OxiCliq A	70 - 100%, +/-2.5 digits (1 S.D.)
	OxiCliq I	
	OxiCliq N	
	OxiCliq P	
	<u>Sensor</u>	<u>Accuracy</u>
	DS-100A	70 - 100%, +/-3 digits (1 S.D.)
	D-YS	
	OXI-A/N	
	OXI-P/I	
	<u>Sensor</u>	<u>Accuracy</u>
	D-YS	70 - 100%, +/-3.5 digits (1 S.D.)
	D-YSE	
	D-YSPD	
	<u>Sensor</u>	<u>Accuracy</u>
	MAX-R	80 - 100%, +/-3.5 digits (1 S.D.)

Pulse: ± 3 digits

Updated of Numerics: every 1 second

Waveform Display: Plethysmogram, fixed for trace 1

Sweep Speeds: 12.5, 25 or 50 mm/sec

Patient Alarms: High and low limits on %SpO₂, adjustable.
High and low limits on Pulse Rate, adjustable.Alarm Delay: %SpO₂: SatSeconds®
Pulse: 0 or 10 seconds

Capnography

Method:	Sidestream Capnography (MicroStream [®])
Patient Range:	Adult - Neonate
Units:	mmHg, % or kPa
Sample Rate:	50 ml/min \pm 7.5 ml/min
System Response Time:	2.9 seconds typical, (includes delay and rise time)
	Rise Time: Adult and Neonate: 190 msec max.
	Delay Time: 2.7 seconds (typical)
Frequency Response:	EtCO ₂ accuracy is maintained up to 80 breaths/min. (For maintaining accuracy for respiration rate over 60 bpm, use Neonate mode.) From 81 to 150 bpm accuracy is \pm 12%, if the EtCO ₂ is higher than 18.8 mmHg in Neonate mode.
Ambient Pressure:	Compensated internally – automatic
Range:	0-99 mmHg (0-13.2 kPa and 0-13.0 vol% at sea level)
Initialization Time:	30 seconds (typical)
Accuracy:	
Before steady state:	0 – 38 mmHg \pm 4 mmHg 39 – 99 mmHg \pm 12% of reading
Steady state:	0 – 38 mmHg \pm 2 mmHg 39 – 99 mmHg \pm 5% of reading +0.08% for every 1 mmHg above 38 mmHg.
Resolution:	1 mmHg, 0.1 % or 0.1 kPa
Respiration Rate:	0-150 breaths/min
Waveform Display:	CO ₂ waveform, fixed for trace 2, Line or Filled
Sweep Speeds:	3, 6.25, 12.5 mm/sec
Patient Alarms:	High and low limits on EtCO ₂ , adjustable FiCO ₂ , fixed

NIBP Measurement

Technique:	Oscillometric (MAXNIBP Technology).	
Patient Range:	Adult – Neonate	
Self Test:	System self test is performed each time power is turned on.	
Auto Zero:	Zero pressure reference is automatically established after every reading.	
Inflation:	Initial inflation to 150 mmHg (Adult) or 100 mmHg (Neonatal) or user selectable. (100, 120, 140, 160, 180, 200) - Adult ; (60, 80, 100, 120) - Neonatal. Subsequent inflation to approximately 30 mmHg greater than previous Systolic pressure.	
Deflation:	Automatically after measurement. If cuff pressure exceeds 290 mmHg (Adult) If cuff pressure exceeds 145 mmHg (Neonate) If measurement time exceeds 120 seconds (Adult), 90 seconds (Neonate) If safety timer detects microprocessor failure	
Max Measurement Time:	Limited to 120 seconds (Adult), 90 seconds (Neonate)	
Overpressure:	If cuff pressure exceeds 290 mmHg (Adult), 145 mmHg (Neonate)	
Measurement Range:	<u>Adult</u>	<u>Neonate</u>
	Systolic: 30-255 mmHg	30-135 mmHg
	Diastolic: 15-220 mmHg	15-110 mmHg
	MAP: 20-235 mmHg	20-125 mmHg
	Pulse Rate Range: 30-240 BPM	40-240 BPM
Accuracy:		
Blood Pressure:	± 5 mmHg, Standard deviation no greater than 8 mmHg.	
Pulse Rate:	± 2% or ± 2 BPM, whichever is greater.	
Patient Alarms:	High and low limits for Systolic Pressure, adjustable. High and low limits for Diastolic Pressure, adjustable.	

Accuracy complies with that given in American National Standard for Electronic or Automated Sphygmomanometers, ANSI/AAMI SP10, 2002. Adult blood pressure measurements determined with this device are equivalent to those obtained by an auscultatory blood pressure measurement device and neonatal ones are equivalent to those obtained by an intra-arterial blood pressure device, within the limits prescribed by the American National Standard for Electronic or Automated Sphygmomanometers. The 4th Korotkoff sound was used to determine Diastolic pressure. Study findings are available.

Patient Alarms

Adjustable Alarms:	High and low alarms for %SpO ₂ , Pulse Rate, EtCO ₂ , Respiration Rate, Systolic Pressure, Diastolic Pressure, No Respiration.
Fixed Alarms:	Asystole, FiCO ₂
Alarm Indicators:	Audible Red Patient Alarm LED's Yellow Equipment Alarm LED's Text in Patient Message Window

Patient Parameter	Neonatal Limit Range		Adult Limit Range	
	Low	High	Low	High
%SpO ₂	70 – 95 %	80 – 99 %	70 – 95 %	80 – 99 %
SpO ₂ Pulse	25 – 235 BPM			
End Tidal CO ₂	1 to 99 mmHg .1 to 13.2 % .1 to 13.2 kPa	1 to 99 mmHg .1 to 13.2 % .1 to 13.2 kPa	1 to 99 mmHg .1 to 13.2 % .1 to 13.2 kPa	1 to 99 mmHg .1 to 13.2 % .1 to 13.2 kPa
Respiration Rate	5 – 145 BrPM			
No Respiration	5 to 30 Sec			
SYS	35 – 130 mmHg	35 – 130 mmHg	35 – 250 mmHg	35 – 250 mmHg
DIA	20 – 105 mmHg	20 – 105 mmHg	20 – 215 mmHg	20 – 215 mmHg

Each alarm limit may also be selected "OFF" individually or as a whole.
Low Limits cannot be set above the associated High Limit.
High Limits cannot be set lower than the associated Low Limit.

Operating Modes

Patient Modes:	Neonate or Adult
CO ₂ :	Continuous Monitoring
%SpO ₂ :	Continuous Monitoring
NIBP:	Manual, STAT or Automatic (at preset intervals).
History:	Trend: Review of previous measurements Alarm: Review of previous alarms
Other Modes:	Stand-by mode and Auto Dim

Display

Display:	LCD display of measurement results, instructions, troubleshooting messages, waveforms and signal strength bar.
Numerics:	%SpO ₂ , PR, NIBP (Systolic, Diastolic and MAP or PR), EtCO ₂ (mmHg, % or kPa), RR.
Waveforms:	Up to 3 traces; SpO ₂ , fixed as trace 1, CO ₂ , fixed as trace 2, 3 rd trace configurable
Sweep Speeds:	12.5, 25 or 50 mm/s for SpO ₂ . 3.0, 6.25 or 12.5 mm/s for CO ₂ .
Trend History:	480 1-minute entries of all parameters.
Alarm History:	25 most recent alarms

Physical Dimensions and Weight

(Fully configured unit with battery)

H x W x D:	170 mm x 215 mm x 102 mm (6.75 in x 8.5 in x 4.0 in)
Weight:	2.0 kg (4.4 lbs)

Operating Environment

Operating Temperature:	0 to 50°C (32°F to 122°F)
Storage Temperature:	-20 to 60°C (-4°F to 149°F)
Humidity:	15 to 95% RH, non-condensing
Altitude:	10,000 to -1,000 ft (690 to 1050 hPa)

Monitors may not meet performance specifications if stored or used outside temperature and humidity ranges. When moving the monitor from a storage location, wait at least one-hour prior to use to allow the monitor to adjust to room temperature.

Power

External Power:

AC: 100-240 VAC, 50/60 Hz, 0.5 A
Fuse Rating – T500mAL250V (two provided)

DC (optional): +12 VDC; 7W
Fuse Rating – Littelfuse Type 154, Slo-Blo,
5.0A, 125VAC or approved equivalent (one
provided)

Chassis Leakage Current: 100 μ A (maximum)

Battery: NiMH battery pack (user removable).

Charge Time: 3-5 hours

Operating Time: 3 hrs (minimum)

Serial Interface

Interface Type:	Bi-directional Serial Communication
Speed:	9600
Signal Level:	RS232C
Data Length:	8 bits
Start Bit:	1 bit
Stop Bit:	1 bit
Parity:	None
Flow Control:	None

Standards

Units comply with the following requirements:

ANSI/AAMI SP-10: 2002

EN 60601-1, EN 60601-1-2

EN 60601-2-27, EN 60601-2-30, EN 60601-20-49

EN 864, EN 865

UL classified – UL 60601-1, CAN/CSA C22.2 No. 601.1

CE marking according to Directive 93/42/EEC

All units covered by U.S. patent 4,796,184 and 5,022,403. Other patents pending.

Certificates

Electronic Emissions and Immunity

The Model 750C Monitor is intended for use in the electromagnetic environment specified below. The customer or the user of the Model 750C Monitor should assure it is used in such an environment.		
Emissions Test	Compliance	Electromagnetic Environment
RF emissions – CISPR 11	Group 1	The Model 750C Monitor uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.
RF emissions – CISPR 11	Class B	The Model 750C Monitor is suitable for use in all establishments, including domestic establishments and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.
Harmonic emissions IEC 61000-3-2	Class B	
Voltage fluctuations / flicker emissions	Complies	

Immunity Test	IEC 60601 Test Level	Compliance Level	Electromagnetic Environment Guidance
Electrostatic discharge (ESD) IEC 61000-4-2	±6 kV contact ±8 kV air	±6 kV contact ±8 kV air	Floors should be wood concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30%.
Electrical fast transient/burst IEC 61000-4-4	±2 kV for power supply lines ±1 kV for in-put/output lines	±2 kV for power supply lines ±1 kV for in-put/output lines	Mains power quality should be that of a typical commercial or hospital environment.
Surge IEC 61000-4-5	±1 kV differential mode ±2 kV common mode	±1 kV differential mode ±2 kV common mode	Mains power quality should be that of a typical commercial or hospital environment.
Voltage Dips, short interruptions and voltage variations on power supply input lines IEC 61000-4-11	< 5% UT (>95% dip in UT) for 0.5 cycle. 40% UT (60% dip in UT) for 5 cycles. 70% UT (30% dip in UT) for 25 cycles. < 5% UT (> 95% dip in UT) for 5 seconds.	< 5% UT (>95% dip in UT) for 0.5 cycle. 40% UT (60% dip in UT) for 5 cycles. 70% UT (30% dip in UT) for 25 cycles. < 5% UT (> 95% dip in UT) for 5 seconds.	Mains power quality should be that of a typical commercial or hospital environment. If user of the model 750C requires continued operation during power mains interruptions, it is recommended that the model 750C be powered from an uninterruptible power supply or a battery.
Power frequency (50/60 Hz) magnetic field IEC 61000-4-8	3 A/m	3 A/m	Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.
NOTE: U_T is the A.C. mains voltage prior to application of the test level.			

Guidance and Manufacturer's Declaration – Electromagnetic Immunity

The Model 750C Monitor is intended for use in the electromagnetic environment specified below. The customer or the user of the Model 750C Monitor should insure that it is used in such an environment.

Immunity Test	IEC 60601 Test Level	Compliance Level	Electromagnetic Environment - Guidance
Conducted RF IEC 61000-4-6	3 Vrms 150 kHz to 80 MHz	3 Vrms	Portable and mobile RF communications equipment should be used no closer to any part of the Model 750C Monitor, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter. Recommended separation distance: $d = 1.2\sqrt{P}$ $d = 1.2\sqrt{P}$ 80 MHz to 800 MHz $d = 2.3\sqrt{P}$ 800 MHz to 2.5 GHz Where P is the maximum output power rating of the transmitter in watts according to the transmitter manufacturer and d is the recommended separation distance in meters. Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey ^a , should be less than the compliance level in each frequency range. ^b Interference may occur in the vicinity of equipment marked with the following symbol: 
Radiated RF IEC 61000-4-3	3 V/m 80 MHz to 2.5 GHz	3 V/m	

NOTE 1 At 80 MHz and 800 MHz, the higher frequency range applies.

NOTE 2 These guidelines may not apply in all situations. Electromagnetic propagation is effected by absorption and reflection from structures, objects and people.

^a Field strengths from fixed transmitters, such as base stations for radio (cellular / cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the Model 750C Monitor is used exceeds the applicable RF compliance level above, the Model 750C Monitor should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as re-orienting or relocating the Model 750C Monitor.

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

Recommended Separation Distances Between Portable and Mobile RF Communications Equipment and the Model 750C Monitor

The Model 750C Monitor is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the Model 750C Monitor can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the Model 750C Monitor as recommended below, according to the maximum output power of the communications equipment.

Rated maximum output power of transmitter (Watts)	Separation distance according to frequency of transmitter (Meters)		
	150 kHz to 80 MHz $d = 1.2\sqrt{P}$	80 MHz to 800 MHz $d = 1.2\sqrt{P}$	800 MHz to 2.5 GHz $d = 2.3\sqrt{P}$
0.01	0.12	0.12	0.23
0.1	0.38	0.38	0.73
1	1.2	1.2	2.3
10	3.8	3.8	7.3
100	12	12	23

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

NOTE 1 At 80 MHz and 800 MHz, the separation distance for the higher frequency range applies.

NOTE 2 These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

CE Marking Information

Compliance

The CAS750C patient monitors bear the CE mark CE-0086 indicating conformity with the provisions of the Council Directive 93/42/EEC concerning medical devices and fulfills the essential requirements of Annex I of this directive.

Exceptions

None.

Accessories

Tuff-Cuff Blood Pressure Cuffs

Reusable Blood Pressure Cuffs

Catalog No.	Type	Description
01-01-0308	CR5216	Large Adult, 16 cm x 42 cm
01-01-0307	CR5214	Adult, 14 cm x 37 cm
01-01-0306	CR5212	Small Adult, 12 cm x 30 cm
01-01-0249	CR5209	Child, 9 cm x 27 cm
01-01-0248	CR5207	Small Child, 7 cm x 21 cm
01-01-0258	CR5206	Infant, 6 cm x 18 cm

Safe-Cuff Blood Pressure Cuffs

Single-Patient-Use Blood Pressure Cuffs

Catalog No.	Type	Description
01-01-0245	CD2060	X-Large Adult, 20 cm x 52 cm
01-01-0243	CD1642	Large Adult, 16 cm x 41 cm
01-01-0241	CD1437	Adult, 14 cm x 36 cm
01-01-0239	CD1230	Small Adult, 12 cm x 31 cm
01-01-0237	CD927	Child, 9 cm x 25 cm
01-01-0235	CD618	Infant, 6 cm x 16 cm

Pedisphyg Blood Pressure Cuffs

Single-Patient-Use Neonatal Blood Pressure Cuffs

Catalog No.	Type	Description
01-01-0267	C26	2.5 cm x 9.0 cm
01-01-0268	C39	3.0 cm x 11.5 cm
01-01-0269	C412	4.0 cm x 14.5 cm
01-01-0270	C515	5.0 cm x 17.75 cm

Inflation Hoses

Catalog No.	Type	Description
01-02-0185		Straight NIBP Hose for Neonatal and Infant applications, 6 feet
01-02-0131		Coiled NIBP Hose for Adult and Pediatric applications, 10 ten

Masimo SpO₂ Accessories

Catalog No.	Type	Description
01-02-0178	LNOP-DCI	Adult digit, reusable sensor
01-02-0190	LNOP-DCIP	Pediatric digit, reusable sensor
01-02-0191	LNOP-YI	Multisite reusable sensor
01-02-0312	LNOP DC-195	Adult digit, reusable sensor
01-02-0251	LNOP-Adt	Adult, single patient adhesive Sensor (box of 20)
01-02-0252	LNOP-Adt Long	Adult, single patient adhesive Sensor, long cable (box of 20)
01-02-0253	LNOP-Pdt	Pediatric/Slender digit single patient adhesive Sensor (box of 20)
01-02-0254	LNOP-Neo	Neonatal single patient adhesive sensor (box of 20)
01-02-0255	LNOP-NeoPt	Neonatal preterm single patient adhesive sensor (box of 20)
01-02-0182	PC 04	Patient Cable, 4 feet
01-02-0192	PC 08	Patient Cable, 8 feet

Nellcor SpO₂ Accessories

Catalog No.	Type	Description
01-02-0179	DS-100A	Durasensor DS-100A, adult finger-clip sensor
01-02-0183	DOC-10	OxiMax Patient Cable, 10 feet

Capnography Accessories

Catalog No.	Type	Description
Microstream CO₂ Consumables for intubated patients		
01-02-0338	006324	FilterLine H, infant/neonate, humid environments (box of 25)
01-02-0392	XS04624	FilterLine H, adult/pediatric, humid environments (box of 25)
01-02-0290	XS04620	FilterLine, adult/pediatric (box of 25)
Microstream CO₂ Consumables for non-intubated patients		
01-02-0329	007266	Smart CapnoLine, pediatric (box of 25)
01-02-0294	007269	Smart CapnoLine O ₂ , pediatric (box of 25)
01-02-0387	008177	CapnoLine H, adult (box of 25)
01-02-0388	008178	CapnoLine H, pediatric (box of 25)
01-02-0389	008179	CapnoLine H infant/neonate (box of 25)
01-02-0390	008180	CapnoLine H, O ₂ adult (box of 25)
01-02-0391	008181	CapnoLine H, O ₂ pediatric (box of 25)
01-02-0330	008174	NIV Line nasal, adult (box of 25)
01-02-0331	008175	NIV Line nasal, pediatric (box of 25)
Other		
01-02-0295		CO ₂ Gas Calibration Kit

Other Accessories and Options

Catalog No.	Description
01-02-0395	Replacement Power Cord, U.S.A.
01-02-0386	Replacement Power Cord, European
01-02-0385	Replacement Power Cord, Australian
01-02-0384	Replacement Power Cord, U.K.
03-08-0386	DC Power Cable
03-08-0450	Monitor Battery Pack
01-01-0047	P9 Calibration Kit
01-02-0248	500 ml Fixed Volume Cylinder
01-02-0172	Roll Stand with Basket, Domestic
01-02-0297	Roll Stand with Basket, International
01-02-0173	Swivel Mount Kit
01-02-0243	Universal Pole Mount
01-02-0174	Carry Bag
01-02-0189	Printer including Battery, RS232 Cable, Power Supply, Roll of Paper and Manual
01-02-0181	Printer Bracket Attachment for 01-02-0172 Roll Stand
01-02-0301	Printer Bracket Attachment for 01-02-0297 Roll Stand
01-02-0188	Printer Battery
28-02-0077	Printer Paper, one Roll
01-02-0266	Printer Power Supply Adapter Plug, European
01-02-0267	Printer Power Supply Adapter Plug, UK
01-02-0268	Printer Power Supply Adapter Plug, Australian
21-02-0191	CAS 750 Service Manual