Operator's Manual

Portable Bedside Capnograph

PN: 010678-A



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Safety Information

Warnings Symbols

To use the portable capnograph correctly and safely, carefully read this operator's manual and the *Directions for Use* for the Microstream EtCO₂ consumables. Use of the monitor requires full understanding and strict observance of these instructions, the precautionary information in boldface type, and the specifications.

Warnings

General

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: To ensure patient safety, do not place the monitor in any

position that might cause it to fall on the patient.

WARNING: Carefully route patient cabling (FilterLine) to reduce the

possibility of patient entanglement or strangulation.

WARNING: Do not lift the monitor by the FilterLine, as it could

disconnect from the monitor, causing the monitor to fall

on the patient.

WARNING: To ensure accurate performance and prevent device

failure, do not expose the monitor to extreme moisture,

such as rain.

WARNING: CO_2 readings and respiratory rate readings can be

affected by certain ambient environmental conditions

and certain patient conditions.



Safety Information

WARNING: The monitor is a prescription device and is to be

operated by qualified healthcare personnel only.

MRI Scanning

WARNING: During MRI scanning, the monitor must be placed

outside the MRI suite. When the monitor is used outside the MRI suite, EtCO₂ monitoring can be implemented using the FilterLine XL. (Refer to MRI Scanning on

page 44.)

Alarms

WARNING: Do not silence the audible alarm if patient safety may be

compromised.

WARNING: Always respond immediately to a system alarm since the

patient may not be monitored during certain alarm

conditions.

WARNING: Before each use, verify that the alarm limits are

appropriate for the patient being monitored.

WARNING: Check the audible alarm silence duration before

temporarily silencing the audible alarms.

Fire Hazard

WARNING: When using the monitor with anesthetics, nitrous oxide,

or high concentrations of oxygen, connect the gas outlets

to a scavenger system.

WARNING: The monitor is not suitable for use in the presence of

flammable anesthetic mixture with air, oxygen, or

nitrous oxide.

WARNING: The FilterLine may ignite in the presence of O_2 when

directly exposed to laser, ESU devices, or high heat. When performing head and neck procedures involving laser, electrosurgical devices, or high heat, use with caution to prevent flammability of the FilterLine or

surrounding surgical drapes.

Electrical

WARNING: To protect against electric shock hazard, the monitor's cover is to be removed only by qualified service

personnel. There are no user-serviceable parts inside.

WARNING: To ensure patient electrical isolation, connect only to

other equipment with circuits that are electrically

isolated.

WARNING: Use only the medical grade AC adapter provided by the

manufacturer. If in doubt about the integrity of the mains supply connection, operate the monitor from its internal

battery pack.

WARNING: Do not connect to a printer or to a PC unless using the

Communication Adapter provided by the manufacturer as an optional accessory. The printer and PC (when connected to the patient through the Communication Adapter) must be distanced from the patient

environment by at least 1.5 m.

Electro-magnetic Interference

This device has been tested and found to comply with the requirements for medical devices according to the standard EN60601-1-2/2001. These standards are designed to provide reasonable protection against harmful interference in a typical medical installation.

However, because of the proliferation of radio-frequency transmitting equipment and other sources of electrical noise in the healthcare environments (for example, cellular phones, mobile two-way radios, or electrical appliances), it is possible that high levels of such interference, due to close proximity or strength of a source, may result in disruption of performance of this device.

WARNING: Operating high frequency electrosurgical equipment in

the vicinity of the monitor can produce interference in

the monitor and give incorrect measurements.

WARNING: Do not use the monitor with nuclear spin tomography

(MRT, NMR, NMT) as the function of the machine may

be disturbed.

Symbols

The following symbols appear on the monitor and monitor LCD (liquid crystal display):

Table 1: Symbols on the Device

Symbol	Description
\triangle	See Directions for Use
GAS	Gas Outlet
(*)	Defibrillator-proof Type BF equipment (patient electrically isolated)
K	Audio Alarms Off
Â	Plug Icon
	Battery Icon
/min	Respiratory Rate (breaths per minute)
EtCO ₂	End tidal carbon dioxide value
<u> </u>	DC Input
Î	Refer to manual for connector interface and other information
Pump Off/On	Pump Off

Introduction

Monitor Features

This manual provides directions for setup and operation of the portable capnograph monitor.

The device is a portable capnograph that continuously monitors end tidal carbon dioxide (EtCO₂), fractional inspired carbon dioxide (FiCO₂) and respiratory rate (RR). The monitor is for monitoring only and must be used in the continuous presence of a qualified healthcare provider. It is intended for use in any environment where continuous, noninvasive monitoring of these parameters is desired, including hospital and hospital type facilities. The monitor is intended for use on adult, pediatric, and infant/neonatal patients.

Monitor Features

- Capnograph in a small, portable, lightweight monitor
- Measures and displays EtCO₂, FiCO₂, and respiration rate, in one graphic and two digital displays
- Displays CO₂ waveforms and trends
- Utilizes a wide range of Microstream EtCO₂ consumables for all applications
- Operates on mains line power or a rechargeable Nickel Metal Hydride battery pack
- Employs audible and visual alarm warnings for monitored parameters and instrument malfunctions
- Provides user selectable language options: English, French, German, Spanish, Italian, Dutch, Swedish, Norwegian, and Portuguese

Introduction

- Displays EtCO₂ and FiCO₂ values in mmHg, kPa or Vol%.
- Provides output for printer, PC, and Digital to Analog Converter
- Provides interface to hospital nurse call systems.

Overview

Principles of Operation
FilterLine
Displays, Controls, and Connectors

The monitor incorporates Oridion's Microstream capnography technology.

Principles of Operation

The monitor uses Microstream non-dispersive infrared (NDIR) spectroscopy to continuously measure the amount of CO₂ during every breath, the amount of CO₂ present at the end of exhalation (EtCO₂) and during inhalation (FiCO₂), and the Respiratory Rate.

Infrared spectroscopy is used to measure the concentration of molecules that absorb infrared light. Because the absorption is proportional to the concentration of the absorbing molecule, the concentration can be determined by comparing its absorption to that of a known standard.

The Microstream EtCO₂ consumables deliver a sample of the inhaled and exhaled gases from the ventilator circuit or directly from the patient (via an oral/nasal cannula) into the monitor for CO₂ measurement. Moisture and patient secretions are extracted from the sample while maintaining the shape of the CO₂ waveform.

The 50 ml/min. sampling flow rate reduces liquid and secretion accumulation, decreasing the risk of obstruction in the sample pathway in humid ICU environments.

Once inside the Microstream CO₂ sensor, the gas sample goes through a microsample cell (15 microliters). This extremely small volume is

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Overview

quickly flushed, allowing for fast rise time and accurate CO₂ readings, even at high respiration rates.

The Microbeam IR source illuminates the microsample cell and the reference channel. This proprietary IR light source generates only the specific wavelengths characteristic of the CO_2 absorption spectrum. Therefore, no compensations are required when different concentrations of N_2O , O_2 , anesthetic agents, and water vapor are present in the inhaled and exhaled breath. The IR that passes through the microsample cell and the IR that passes through the reference channel are measured by the IR detectors.

The microcomputer in the monitor calculates the CO_2 concentration by comparing the signals from both channels.

Microstream EtCO₂ Consumables

The following products comprise the Microstream EtCO₂ consumables:

Sample Lines and Airway Adapter Sets for Intubated Patients:

- FilterLine Set (for non-humid environments).
- FilterLine H Set (for humid environments).

Nasal and Oral/Nasal Cannulas for Non-intubated patients:

- Smart CapnoLine Plus for use in procedural sedation. Also available with O₂ delivery.
- CapnoLine H for patients receiving hi–flow oxygen by mask, on long term CPAP or Bi-PAP, or post-op pain management. Also available with O₂ delivery.
- NIV Line for use under oxygen, CPAP, Bi–PAP or NPPV mask.
- Smart products provide oral and nasal sampling.
 H products are for long term use.

Special Procedure FilterLines

• FilterLine XL – Provides extended length so that the monitor can be used safely during MRI (see MRI Scanning on page 44).

Note:

The generic term FilterLine, used in this manual, is interchangeable with any of the Microstream EtCO₂ consumables.

FilterLine

The FilterLine has five active elements that work together to offer a solution to the problems that have previously posed challenges to capnography in ICU, emergency, and intra-transport applications. These elements are described below.

* Hydrophobic filter

The hydrophobic filter is located at the end of the sample line that is closest to the capnograph. This filter strips the remaining water vapor from the gas sample while keeping a laminar flow of the gas. This laminar flow minimizes distortion of the CO_2 waveform.

This filter is made of a $0.2~\mu$ hydrophobic porous medium.

* Drying element

The drying element is a tube made of a synthetic material that is chemically stable and has high water absorption. This material allows the water vapor to pass outside the tube, thereby adjusting the humidity inside the FilterLine close to the level of humidity in the ambient air.

* Sample line

The sample line has low dead space due to its small internal diameter. This provides a sharp waveform and an accurate CO_2 reading at a high breath rate per minute. The sample line is not affected by gases and anesthetic agents in the operating room environment.

* FilterLine Recognition Safeguard

When the FilterLine is attached to the monitor, the FilterLine Recognition Safeguard (FRS) identifies the FilterLine and activates the monitor, thus enabling measuring.

* Airway Adapter

The airway adapter design provides multiple channels for the sampled air from the airway, minimizing the possibility of water infiltration or line blockage. These multiple channels allow uninterrupted monitoring for all adapter orientations and in all applications. The airway adapter provides optimal performance in all directions and is seldom disabled by secretions or liquids.

Displays, Controls, and Connectors

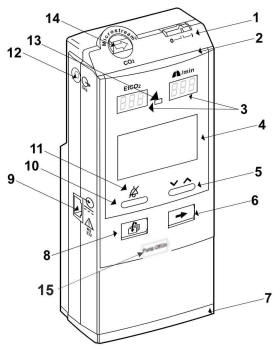


Figure 1: Monitor Front and Side View

The numbered labels in Figure 1: Monitor Front and Side View are described below.

- 1. On/Off Switch
- 2. Alarm Bar
- 3. Digital Display of EtCO₂ and Respiration Rate
- 4. Graphic Display
- 5. Contrast/Value Change Button
- 6. Next/Menu Button
- 7. Battery Pack
- 8. Event/Home Button

- 9. Port for AC Adapter or communication adapters
- 10. Alarm Silence/Alarm Silence Menu Button
- 11. Alarm Silence Indicator
- 12. Gas Outlet
- 13. Photo Resistor
- 14. FilterLine Input Connector
- 15. Pump Off/On Adhesive Label

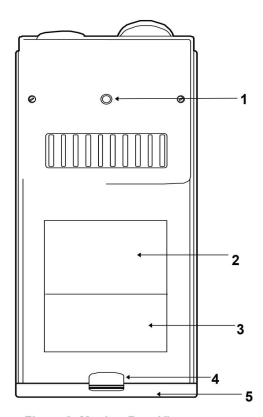


Figure 2: Monitor Rear View

The numbered labels in Figure 2: Monitor Rear View are described below.

- 1. Clamp Connector
- 2. Space for Quick Guide Adhesive Label
- 3. Serial Number Label
- 4. Battery Pack Release Button
- 5. Battery Pack

Initial Setup

Power Requirements
Unpacking and Inspection
Start-up and Self test
Measuring Mode
Quick Guide

Power Requirements

The monitor operates on batteries or on AC power. It is equipped with a rechargeable Nickel Metal Hydride battery pack. When a power outlet is available, use the medical grade AC adapter provided with the monitor.

Before using the monitor in the field, ensure that the battery pack is fully charged. At the Measuring mode, check that the battery icon at the right side of the graphic display is full.

Note: If the

If the battery is not fully charged, the icon may first show as full and after a short period of time will drop to indicate the real charge level.

A fully charged battery pack provides between four and seven operating hours, depending on power management (refer to Table 6: Instrument Settings Parameters (Menu 1) on page 38 for a description of the power management options).

WARNING:

Use only the medical grade AC adapter provided by the manufacturer. If unsure about the integrity of the line connection, operate the monitor from its internal battery pack.

WARNING:

To ensure patient electrical isolation, connect only to

other equipment with circuits that are electrically

isolated.

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Initial Setup

Battery and Power Usage

If power is lost when the monitor is operating from AC power, it automatically switches to its internal battery pack for power.

A plug-shaped icon

at the bottom right side of the graphic display is displayed when the monitor operates from an external power source and the battery pack is fully charged. A battery-shaped icon is displayed when the monitor operates from the battery pack. The battery icon will show the battery pack's approximate charge level. An advisory message, Battery

!, appears when approximately 40 minutes of battery charge remains. A caution message, Battery
!!, appears when approximately 15 minutes of charge time remains.

While the monitor is connected to AC power, the battery pack can be replaced without interrupting monitoring.

Battery Pack

Before using a new battery pack for the first time, charge and discharge the battery three times to ensure full battery capacity. For charging and discharging, the Microstream Capnograph Battery Charger is recommended (refer to the Microstream Capnograph Battery Charger *Directions for Use*).

Internal Recharge Function

CAUTION:

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Do not attempt to disassemble the battery pack. It is a sealed unit and has no serviceable parts inside.

When the monitor is connected to an external power source (even if the monitor is turned off), the battery pack charges automatically. If the instrument is on during charging, the battery-shaped icon displays a filling pattern. It takes approximately 4.5 hours to fully charge an empty battery pack. Additional battery packs can be purchased from your local representative.

The recommended temperature for battery charging is between 5°C and 45°C.

CAUTION:

Important! The following information relates to the safe handling, storage, and disposal of the monitor battery pack.

Battery Testing

The battery pack charge level should be tested before each use by observing the level on the battery icon after Self Test. For a correct reading, wait for the battery charge level to stabilize. Replace or recharge the battery pack when the advisory message, Battery ▶!, appears on the graphic display screen (refer to Troubleshooting on page 49).

Handling

- Do not immerse the battery pack in water; it may malfunction.
- Only recharge the battery pack in the monitor or use the Microstream Capnograph Battery Charger, provided by your local representative, to avoid possible overheating, burning or rupture of the battery pack.

Storage

- Short-term storage (one month or less): The battery pack
 has an automatic discharge feature. You must periodically
 check the charge level of the battery pack.
- Long-term storage (6 months or more): The battery pack must be stored in a cool, dry area. Its charge decreases over time. To restore the battery pack to full power, charge and discharge it three times before use. Long-term storage, without charging the battery, may degrade the battery capacity.

Disposal

- Do not dispose of the battery pack in fire; it may explode.
- Be sure to follow local governing ordinances and recycling instructions regarding disposal or recycling of batteries.

Unpacking and Inspection

Components

Remove the monitor and the accessories from the box carefully.

Check that the items listed on the back cover of this manual are included.

Inspect each component. If the package is damaged or any component is missing, contact your local representative.

Optional Accessories

The following items are available for use with the monitor:

- Protective Boot
- Carrying Case
- Clamp
- Rechargeable Battery Pack
- Microstream Capnograph Battery Charger
- Battery Pack Carrying Pouch
- 12 Volt Cable
- Communication Adapter Kit
- Calibration Gas Kit
- Service Manual
- Digital to Analog (D/A) Converter
- Seiko DPU 414 Printer
- Nurse Call Interface Kit
- MSM (Microstream Monitor) Interface Kit
- Vuelink Module & Interface Kit for use with Philips patient monitoring systems
- Flexport Module & Interface Kit for use with SpaceLabs patient monitoring systems
- Profox Respiratory Software

Start-up and Self test

Note: For information on operating the monitor with any

accessory, refer to the specific accessory's Directions

for Use.

CAUTION: To protect the unit, the manufacturer recommends using

the carrying case, the clamp, or the protective boot,

depending on the type of application.

Start-up and Self test

WARNING: Do not lift the monitor by the FilterLine as it could

disconnect from the monitor, causing it to drop on the

patient.

WARNING: To ensure patient safety, do not place the monitor in any

position that might cause it to fall on the patient.
Carefully route the FilterLine cable to reduce the possibility of patient entanglement or strangulation.

WARNING: When using the monitor with anesthetics, nitrous oxide,

or high concentrations of oxygen, connect the gas outlets

to a scavenger system.

CAUTION: The monitor is intended only as an adjunct in patient

assessment. It must be used in conjunction with clinical

signs and symptoms.

CAUTION: The monitor is a prescription device and is to be

operated by qualified healthcare providers only.

CAUTION: Only use Microstream EtCO₂ consumables to ensure

that the monitor functions properly.

Preparation

Prior to start-up:

- 1. Slide open the FilterLine input connector shutter and connect the appropriate FilterLine.
- 2. Connect the FilterLine to the patient as described in the *Directions for Use*.

Initial Setup

Note: When the monitor is used in stationary applications, secure it with the clamp (available as an optional

accessory).

Initialization

CAUTION: If any monitor response does not seem appropriate, do

not use the monitor. Instead, contact your local

representative.

CAUTION: Immediately after power-up, confirm that all display

segments and icons function.

3. Turn on the monitor by sliding the on/off switch to the on position.

- 4. Verify that the monitor is working properly. Proper working condition is verified by completing the power-on Self Test described below.
- 5. When turned on, the monitor automatically performs a Self Test. The display and alarm functions are tested activating the LCD, alarm bar, seven segment displays, alarm silence indicator, and buzzer. In this mode all alarms are disabled. The initialization screen displays for 5 seconds (refer to Figure 3:

Initialization Screen at

right).



Self Test

Figure 3: Initialization Screen

During Self Test, the $EtCO_2$ and Respiration Rate LEDs show dashes. When the monitor is ready and the FilterLine is connected, the dashes in the $EtCO_2$ and Respiration Rate LEDs are replaced by numeric values. If the FilterLine circuit is not connected to the monitor, dashes will appear on the LEDs.

Measuring Mode

In Measuring mode, the monitor measures, displays, and stores event data, or prints data that has been stored in its memory.

During measuring, the monitor shows EtCO₂ and respiration rate readings on the digital displays. Waveform, respiration rate, and other information, according to the selected screen (see Basic Operation on page 29), are shown on the graphic display.

The monitor begins measuring EtCO₂ after recognizing one breath (after monitor power-up or after exiting Standby). The monitor recognizes two breath measurement ranges:

Valid breath: values > 7.5 mmHg (for adult mode) or >5.0 mmHg (for neonatal mode)

Low readings breath: values <7.5 mmHg (for adult mode) or <5.0 mmHg (for neonatal mode)

Note:

If the first breath the monitor recognizes is a Low readings breath, the monitor will not display nor emit warning signals and a No Breath message will not appear. If the values go above 7.5 mmHg (for adult mode) or 5.0 mmHg (for neonatal mode), and then fall below these ranges, the monitor will display a No Breath message and emit warning signals (see Troubleshooting on page 49).

 $\rm EtCO_2$ readings between 3.0–7.0 mmHg (adult mode) or 3.0-5.0 mmHg (neonatal mode) appear as numerical values on the $\rm EtCO_2$ LEDs. Readings <3.0 mmHg show as 0 (zero) on the LEDs.

The monitor begins measuring Respiration Rate after two valid breaths.

The waveform appears on the graphic display for all EtCO₂ values.

Battery Pack and AC Operation

- Connect only Microstream EtCO₂ consumables to the monitor.
- 2. Battery pack operation: First, switch the monitor on and **AVOBUS** that the battery pack is charged (in Measuring mode,

check that the battery icon on the right side of the graphic display is full).

AC operation: Connect the AC adapter to the monitor, and plug the cord into the mains power supply. Switch the monitor on. Check that the battery icon displays a filling pattern or the plug icon appears.

(Refer to Figure 4: Quick Guide for all button functions.)

3. Adjust the parameters in the Alarm Limits menu, Instrument Setup menu, and Alarm Silence menu to the values appropriate to the patient's condition.

Quick Guide

The Quick Guide adhesive label is included in the monitor packaging. Apply the label to the monitor as shown in Figure 4: Quick Guide on page 18.

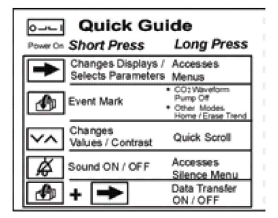


Figure 4: Quick Guide



Consumables

MICROSTREAM EtCO2 Consumables

MICROSTREAM EtCO₂ Consumables

- FilterLine Set
- FilterLine H Set
- Smart CapnoLine and Smart CapnoLine Plus (available with O₂ delivery)
- CapnoLine H
- NIV Line

(For a description of Microstream EtCO₂ Consumables see Microstream EtCO₂ Consumables on page 14.)

CAUTION: Before use, carefully read the Microstream EtCO₂

Consumables Directions for Use.

CAUTION: Only use Microstream EtCO₂ Consumables to ensure the

monitor functions properly.

CAUTION: Microstream EtCO₂ Consumables are designed for

single patient use, and are not to be reprocessed. Do not attempt to disinfect or flush the FilterLine as the monitor

can be damaged.

CAUTION: Dispose of Microstream EtCO₂ Consumables according

to standard operating procedures or local regulations for

the disposal of contaminated medical waste.

Consumables

WARNING:

The FilterLine may ignite in the presence of O_2 when directly exposed to laser, ESU devices, or high heat. When performing head and neck procedures involving laser, electrosurgical devices or high heat, use with caution to prevent flammability of the FilterLine or surrounding surgical drapes.

Basic Principles

When choosing Microstream EtCO₂ consumables, the following should be considered:

- Intubated versus non-intubated
- Whether the patient is on mechanical ventilation
- Duration of use
- Patient's size and weight

For further information, please contact your local representative.

Select the appropriate FilterLine and connect it to the monitor before attaching it to the patient's airway. Be sure to follow Microstream EtCO₂ Consumables' *Directions for Use* for proper connection.

Basic Operation

Data Display Screens
Displayed Data Options
Alarm Functions
Alarm Limits Menu
Alarm Silence/Standby Menu
Instrument Settings Menus
MRI Scanning
Standby
Pump Off Mode
Pump Off/On Label

Data Display Screens

In Measuring mode, the monitor constantly measures and displays the CO₂ waveform, EtCO₂ numerical value, respiratory rate (RR) and FiCO₂ (user selected) values.

Note:

For both neonatal and adult patients, the EtCO₂ displayed on the LED Numeric Displays represents the maximum value during the last 15 seconds (updated every 5 seconds). The EtCO₂ is displayed from the first breath. The EtCO₂ warning is according to the value in the 7-Segment display.

The respiratory rate and EtCO₂ values are constantly shown in the digital displays. Waveform or Trends are shown in the graphic display depending on the selected display screen (Figure 5: Monitor Display Screen and LEDs, below). Power icons, advisories, warnings, or cautions appear super-imposed over the data display.

At any time during the Measuring mode, the user can mark a special event by a short press of and a short duration tone sounds. The

Basic Operation

event is stored in the monitor's memory and will appear on the data printout marked by an asterisk (*) on the tabular trend printout and as a mark on the graphic trend printout (perpendicular to the trend graph).

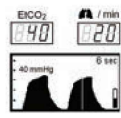


Figure 5: Monitor Display Screen and LEDs

There are four data display screens (Table 2: Display Screens on page 32):

- CO₂ Waveform
- CO₂ Trend, 30 minutes
- CO₂ Trend, 8 hours
- Meter Mode

CO₂ Waveform

The CO₂ waveform screen displays a real-time CO₂ waveform. The end tidal CO₂ and Respiration Rate values are shown simultaneously on the digital displays.

CO₂ Time Base

The time base is the period of time captured on the display. The time base default values are:

- 6 seconds for Adult Mode
- 3 seconds for Neonatal Mode

The instrument automatically changes the time base of the CO₂ waveform according to the actual respiration rate as follows:

Current Time Base		Time Base Change Condition	New Time Base	
	6 seconds	>35 bpm for 10 seconds	3 seconds	

Data Display Screens

3 seconds	<25 bpm for 10 seconds	6 seconds
Any	Initialization, "No Breath" or	6 seconds
	blockage	

During periods of high respiration rates, the display will automatically depict the shorter time base to avoid compression of the waveform.

The time base appears at the top right side of the graphic screen as a Temporary Silent Advisory and is shown for 5 seconds each time the instrument enters the CO₂ waveform screen or after every change of the time base. The instrument also automatically changes the time base when changing from Adult or Neonatal Mode.

CO₂ Trends

The trends graphs represent trend data of the last 30 minutes or 8 hours (15-second or 4-minute resolution respectively). The trends are shown in the CO₂ scale selected by the user. The tabular trend data for 14 hours (5-second resolution) is relevant for the print/PC option only.

During the 14-hour tabular trend period, the data of (up to) the last 100 patients is stored. A new patient is defined each time the monitor is turned off and on or enters Standby.

Note: In the case of the "Autoscale Option," the CO₂ scale is that of the maximum range.

The FiCO₂ value shows as light pixels (a light area) at the bottom of the trends graph.

When the monitor is turned on, a trend data border will mark the end of the previous trend. A trend data border will also appear after exiting Standby and returning to Measuring mode. The trend data border is a vertical line on the graph. An event will appear on the tabular trend printout marked by an asterisk (*) and as a mark on the graphic trend printout (perpendicular to the trend graph).

When you enter a trend display, a temporary message Press To Erase appears for 3 seconds. You now have the option to erase all old trends as follows: Press (the message begins to flash) and hold it until the message disappears. This message will not appear during an alarm.



Basic Operation

The real-time EtCO₂ and respiration rate values are shown on the real-time digital display.

Meter Mode

The Meter Mode screen displays the EtCO₂ value on the left side of the screen and the respiration rate on the right side of the screen.

This mode is recommended in the following cases:

- When the power management is Low (see Table 6: Instrument Settings Parameters (Menu 1) on page 38).
- When the monitor display is exposed to direct sunlight affecting the digital display reading.

Graphic Display Screen Contrast

The LCD contrast intensity can be adjusted during Measuring mode.

To adjust the contrast, press on the contrast button ; press the right side for a darker contrast and the left side for a lighter contrast.

The photo resistor senses the ambient light intensity and accordingly switches the backlight on or off during Normal power management setting.

Displayed Data Options

Table 2: Display Screens

To View	To View Press Result	
CO ₂ waveform	Appears automatically	ERCOz / min BB0 40 mmHg 6 sec

To View	Press	Result
CO ₂ Trend – 30 min	1st short press	EICO2
CO ₂ Trend - 8 hr.	2nd short press	
Meter Mode	3rd short press	40 20 RADIN
CO ₂ waveform	4th short press	EICO ₂ / min 40 mmHg

Note: To return to CO₂ waveform (Home) from any screen

or menu, long press the button.

Alarm Functions

WARNING: Do not turn off the audible alarm if patient safety could

be compromised. Pressing the alarm silence button turns the audio alarms OFF and turns the alarm silence icon LED indicator ON. In this condition, there will be no audible alarms in the event of adverse patient conditions.

WARNING: When exiting Standby mode, the monitor reverts to the

Factory default of "All Alarms On."

The monitor has four levels of alarms. For full details on alarms, see Troubleshooting on page 49.

Alarms

Warnings are the highest level of alarms, used to alert the user that the patient's condition is beyond predefined limits. Alarms can be set from the *Alarm Limits* menu (see Alarm Limits Menu on page 35). The monitor has the following alarms with adjustable level settings:

- No Breath (alerting the user when no valid breath is detected after a predetermined time)
- EtCO₂ high and low levels
- Respiration rate (RR) high and low levels
- FiCO₂ high level

The following alarms alert the user of the instrument's status or malfunction:

- Caution messages (audible and visual)
- Advisory messages (audible and visual)
- Silent advisory messages (visual)
- Pump Off two minute warning (audible and visual)

Factory Default Alarm Range Values

Table 3: Factory Default Alarm Range Values on page 35 lists the default values of the various alarm ranges. These values can be changed from the *Alarm Limits* menu.

CAUTION: Make sure that the monitor default alarm settings are

appropriate for the specific patient being monitored.

CAUTION: The monitor will revert to its default alarm limit settings

at power on, power interruption, and when changing the

patient mode.

Note: The user can have the factory default values of the

alarm range permanently changed (see Institutional Settings on page 43). For further information, call your

local representative.

The CO₂ values in the table are shown in mmHg. The values in brackets correspond to the kPa and Vol% (at sea level).

Table 3: Factory Default Alarm Range Values

Parameter	Adult Default	Neonate Default	Maximum	Minimum
EtCO ₂ high	60 [8.0]	60 [8.0]	100 [13.0]	5 [0.5]
EtCO ₂ low	0	0	99 [12.9]	0 [0.0]
FiCO2 high	8 [1.1]	8 [1.1]	99 [12.9]	2 [0.1]
RR high	150	150	150	1
RR low	3	12	149	0
No Breath delay*	30	20	60	10

No Breath appears in the *Alarm Limits* menu as "No Resp." See Instrument Settings Menus on page 38 for a list of parameters that are set by the user and stored in the memory.

Alarm Limits Menu

Table 4: Alarm Limits on page 35 explains how to access the *Alarm* Limits menu and to change the parameters and values.

Important! "No Resp" will appear as "No Breath" on Note: the monitor display.

Table 4: Alarm Limits

Objective	Action	Result		
To access the Alarm Limits menu from any measuring display*	long press	<u>Patient Adolt</u> EtC02 末 50 坐 0 FiC02 末 8 NoResp© 30 RR 末150 坐 3		
To change the patient mode**	short press	Patient Neonatal EtC02		

Basic Operation

 ore operation		
To access any displayed parameter	short press	Patient Neonatal EtCO2
To change the parameter's value	short press/long press***	Patient Neonatal EtCO2
To exit and return to Measuring mode (at any point in the Alarm Limits menu)****	long press	40 minHg

- If after 15 seconds no action is taken, the display returns to Measuring mode.
- The Neonatal mode is recommended when a patient's breath rate is >50 breaths per minute.
- *** Long press: the value advances quickly.
- **** Display does not necessarily return to the wave form shown in the Results column; it returns to the screen active prior to entering the Alarm Limits menu

Alarm Silence

Alarms can be temporarily silenced. A short press of the alarm silence button will temporarily disable the audible alarm for a pre-set period of time and the alarm silence indicator will be lit. The audible alarm can be reactivated with a short press of the alarm silence button. The default setting is 2 minutes. You can change this setting from the Alarm Silence/Standby Menu (Table 5: Alarm Silence/Standby on page 37).

From the Alarm Silence menu, you can choose to permanently disable a specific audible alarm or all audible alarms. Whenever an alarm is disabled indefinitely, the alarm silence indicator \checkmark will be lit on the front panel and the Alarm Silence icon will appear on the right side of the graphic display with the appropriate label.

• ALL: All the audible alarms are turned off.

• CO2: CO₂ audible alarms (including No Breath

message) are turned off.

Note: When any alarm is disabled, a single caution burst

can sound once every three minutes if this option is selected through Institutional Settings (refer to Institutional Settings on page 43). If an alarm condition occurs when any corresponding alarm is disabled, a message is generated on the monitor

display.

Note: If either ALL or CO₂ audible alarms are off and the

alarm silence button is pressed, the screen will remove the ALL or CO₂ message next to the alarm icon while all alarms are silenced temporarily. When the time limit for alarm silence is reached, the ALL or

CO₂ message returns.

Alarm Silence/Standby Menu

Table 5: Alarm Silence/Standby

Objective	Action	Result	
To access the Alarm Silence/Standby menu from any measuring display*	long press	Alarm Silence 2min All Alarms On CO2 Alarms On Standby Off	
To change the silence period**	short press	Alarm Silence 1min All Alarms On CO2 Alarms On Standby Off	
To access any displayed parameters	short press	Alarm Silence 1 min All Alarms On CO2 Alarms On Standby Off	

Basic Operation

Objective	Action	Result	
To change the setting of the selected parameter	short press	Alarm Silence 1 min All Alarms Off CO2 Alarms On Standby Off	
To exit and return to measuring display (at any point in the Alarm Silence/Standby	long press	40 mmHg	

^{*} If after 15 seconds no action is taken, the display returns to Measuring mode.

Instrument Settings Menus

Instrument Settings Menu Parameters

Table 6: Instrument Settings Parameters (Menu 1) and Table 7: Instrument Settings Parameters (Menu 2), both below, explain the user-defined parameters that can be set from the Instrument Settings menus.

Table 6: Instrument Settings Parameters (Menu 1)

Parameter		User options
CO ₂ units	mmHg, kPa, Vol%	
Power Mgmt	Full -	Display backlight on and 7 segment
		LEDs at high intensity.
	Normal	Display backlight on and 7 LEDs
	segments at normal intensity.	
	Low -	Backlight and 7 LEDs segment off.
	Note: During AC power, power management	
	appears as full.	

^{**} Alarm silence limits are from 1-2 minutes.

	ge	
Parameter	User options	
Print	Screen – the current display is printed	
	Graphic Trend – real time trend is printed in a	
	graphic form	
	Trend History – stored trend is printed in graphic	
	and tabular form	
	Tab. Trend (5s) Real time trend data is printed in	
	tabular form (every 5 seconds)	
	Tab. Trend (1m) Real time trend data is printed in	
	tabular form (every minute)	
	Tab. Trend (14H) – with a resolution of 5	
	seconds. Stored trend is communicated in tabular	
	form	
CO ₂ scale	0-50 mmHg (0-7 kPa or Vol%)	
	0-99 mmHg (0-14 kPa or Vol%)	
	Autoscale	
FiCO ₂	On: display FiCO ₂	
	Off: do not display FiCO ₂	
	Default: Off	

Make sure the patient type and CO₂ scale are appropriate **WARNING:** for each patient. An error in the patient type can cause incorrect alarm limits or incorrect CO₂ readings. If the CO₂ scale is not appropriate, the waveform will be either incomplete or small.

CO₂ Scale: Autoscale

When Autoscale is selected, the CO₂ scale changes as follows:

- From lower to higher scale after 12 consecutive breaths with EtCO₂ values greater than low level scale limit.
- From higher to lower scale after 12 consecutive breaths with EtCO₂ values less than low level scale limit.

Basic Operation

Whenever autoscale is selected, the trend scale (and printed graphic scale) will be the high-level scale limit.

The Factory Default CO₂ is 0-50 mmHg. The CO₂ scale option will not return to the Factory Default after being changed by the user. See User-defined Parameters on page 40.

Table 7: Instrument Settings Parameters (Menu 2)

Parameter	User options
Language	English, French, German, Spanish, Italian, Dutch,
	Swedish, Norwegian and Portuguese.
Check Cal.	Off/Start
	See CO2 Calibration Check on page 59
Factory	Off/Start
Default	This option will reset the device to the factory
	default settings.

User-defined Parameters Stored as Defaults

The following parameters will not return to their defaults after being changed by the user. These parameters are stored in the memory of the monitor until the next time they are changed by the user.

CO ₂ Scale CO ₂ Units	
CO ₂ Mode (Patient)	Print
Language	Power Management

Note:

When changing any of these parameters, wait approximately 10 seconds before turning the monitor off. If you turn off the monitor immediately after changing the parameter, the new setting may not be saved.

Changing Instrument Settings

Table 8: Changing Instrument Settings (Menu 1) and Table 9: Changing Instrument Settings (Menu 2), both below, describe how to change the instrument settings.

Note: If after 15 seconds no action is taken, the display returns to Measuring mode.

Table 8: Changing Instrument Settings (Menu 1)

Objective	Action	Result
To access the Instrument Settings menu 1. (From any measuring display, the 1st long press accesses the Alarm Limits menu. The 2nd long press accesses the Instrument Settings menu 1.)	long press (x2)	CO2 Scale 0 – 50 CO2 Units mmH9 FiCO2 Off Power Mgmt Normal Print Screen
To change parameter setting	short press	CO2 Scale 0 - 99 CO2 Units mmH9 FiCO2 Off Power M9mt Normal Print Screen
To access the next displayed parameter	short press	CO ₂ Scale Ø – 99 CO ₂ Units mmHg FiCO ₂ Off Power M9mt Normal Print Screen
To change the parameter setting	short press	CO ₂ Scale 0 – 99 CO ₂ Units VolX FiCO ₂ Off Power Mgmt Normal Print Screen

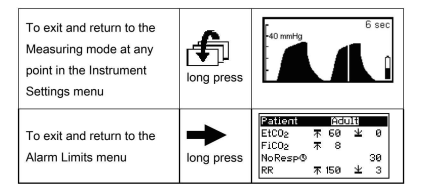


Table 9: Changing Instrument Settings (Menu 2)

Objective	Action	Result
To access the Instrument Settings menu 2. (From any measuring display, the 1st long press accesses the Alarm Limits menu. The 2nd long press accesses the Instrument Settings menu 1. The 3rd long press accesses the Instrument Settings menu	long press (x3)	Check Cal. Off Language English
To change the Check Calibration option To access the language	short press	Check Cal. Start Language English Check Cal. Off Language English
option	short press	

Objective	Action	Result
To switch languages	short press (until desired language appears)	Check Cal. Off Language Français

Institutional Settings

The factory default parameter settings in Table 10: Institutional Settings, below, can be changed by your local service representative.

Table 10: Institutional Settings

Parameter	Factory Default Setting
Alarm Default Settings*	See Table 3: Factory Default
	Alarm Range Values. on
	page 35
3 Min Alert	OFF
(to remind user that alarms are set	
to off)	
BTPS (body temperature,	ON
pressure, saturation assumed	
37°C, 47mmHg)**	
Pump Off	15 minutes

* Calculations are made according to:

$$PCO_2 = FCO_2 \times (Pb - 47)$$

Where

FCO₂ is the Fractional concentration of CO₂ in dry gas,



Where $FCO_2 = \% CO2/100$ Pb = the ambient pressure $PCO_2 =$ the partial pressure of CO_2 at BTPS

MRI Scanning

WARNING: Do not use the FilterLine H Set Infant/Neonatal

consumable during magnetic resonance imaging (MRI) scanning. Using the FilterLine H Set Infant/Neonatal during MRI scanning could harm the patient.

during MRI scanning could harm the patient.

CAUTION: During MRI scanning, the monitor must be placed outside the MRI suite. When the monitor is used outside

the MRI suite, EtCO₂ monitoring can be implemented by using the FilterLine XL, to provide extended length.

Non-invasive EtCO₂ monitoring during magnetic resonance imaging (MRI) can be accomplished with the monitor and a FilterLine XL as follows:

1. Place the monitor outside the MRI suite. There must be a hole in the wall of the suite (approximately 10cm diameter).

Note: A small hole at the base of the wall does not affect the integrity of the MRI shielding (shielding of a 1.5 Tesla magnet).

2. Connect the FilterLine XL to the monitor and guide the FilterLine XL through the hole in the wall of the MRI suite. Attach the FilterLine XL to the patient.

Note: Due to the extended length of the FilterLine XL, there may be a slower response and a decreased

fraguency response time

frequency response time.

To purchase the FilterLine XL, contact your local representative.

Standby

The Standby mode is an automatic or selectable function designed to reduce power consumption and to avoid unnecessary alarms.

To set the monitor manually in the Standby mode, choose the Standby ON option from the *Alarm Silence/Standby* menu (Table 5: Alarm

Silence/Standby on page 37). The Standby screen appears. A long press of any key restores the Measuring mode. (The *Alarm Limits* menu appears briefly before the Measuring mode but the alarms cannot be changed at this time).

The monitor will automatically enter Standby mode if, after Power ON, no signal is registered for 10 minutes.

Note: When exiting Standby mode, the monitor reverts to

the Factory default of "All Alarms On."

Note: The Alarm Limits settings are not changed (do not

revert to defaults) when moving to and from Standby

mode.

Pump Off Mode

The Pump Off mode is a selectable function designed to prevent liquids from entering and saturating the filter. During Pump Off mode, pump activity is suspended to facilitate drug delivery, suctioning and equipment changes while avoiding the need to replace the consumable due to blockage.

WARNING: If at any time the device displays the Blockage!! message, replace the consumable.

1. From the CO₂ Waveform screen, select Pump Off mode by long pressing once.



Figure 6: Pump Off

Note: During Pump Off mode the CO₂ parameter is dashed

and the SpO₂ function operates normally.

Note: The time range for Pump Off is 5–30 minutes. The

factory default time for Pump Off mode is 15 minutes. To change the factory default, contact a qualified

service technician.

2. Exit Pump Off mode by one long press except during the last two minutes.

Basic Operation

3. During the last two minutes an alarm sounds indicating there are two minutes left before the device exits Pump Off mode automatically. This alarm cannot be disabled. One long press resets Pump Off mode to the default time.



Figure 7: Pump Off Additional
Time

Another long press will exit the Pump Off mode.

Pump Off/On Label

The Pump Off adhesive label is included in the monitor packaging. Adhere the label to the monitor as shown in Figure 1: Monitor Front and Side View on page 17.



Figure 8: Pump Off/On Label

Communication Interface

Communication Interface

Communication Interface

The monitor can integrate data to the following devices:

- Communication Adapter
- Printer (Seiko DPU-414)
- PC
- Digital to Analog Converter
- Nurse Call systems (using the Nurse Call Interface Kit)
- Patient monitoring systems (Philips and SpaceLabs systems only; available for Microcap devices with software versions of 3.15 and up)

Note: The monitor interfaces to the PC or Printer with the Communication Adapter Kit.

For interface directions to the different devices, refer to the *Directions* for *Use* of the appropriate devices and/or the *Communication Interface Manual*.

WARNING: When connecting the monitor to another instrument,

verify its proper operation before clinical use. Refer to the other device's manual for full instructions. For further questions, contact your local representative.

WARNING: Do not connect the monitor to a printer or to a PC unless

using the Communication Adapter Kit provided by the

manufacturer, as an optional accessory.

Communication Interface

WARNING:

When using the printer/PC with mains line power, it is recommended to use a medical grade power supply complying with the following standards: EN60601-1, UL 60601-1, CSA C22.2 No. 601.1-M90. If the power supply is not medical grade, the printer/PC must be placed at least 1.5 meters from the patient environment to comply with standard EN60601-1-1.

Troubleshooting

Alarms and Messages Troubleshooting Guide

This section lists the alarms and messages and the corresponding actions the operator should take. The Troubleshooting section discusses potential problems and suggestions for resolving them. If the problem persists and the message remains, contact qualified service personnel or your local representative.

Alarms and Messages

The monitor displays the following four types of alarms and messages in order of priority:

- Warnings
- Cautions
- Advisories
- Silent Advisories

Alarm and Message Priorities

The messages in the following tables (Table 11: Warning Messages, Table 12: Caution Messages, Table 13: Advisory Messages, Table 14: Silent Advisory Messages) are listed in order of priority.

In the event that several problems occur simultaneously, the higher priority will appear first on the display. After each problem is resolved, the next message is displayed in order of priority.



Warnings

WARNING:

Always respond immediately to a system alarm since the patient may not be monitored during certain alarm conditions.

Warnings refer to either patient or alarm limit settings problems. They are serious and require immediate operator attention. The message appears on the screen followed by !!!, the numerical parameter associated with the alarm blinks, the alarm bar flashes red, and a special, repetitive warning tone is heard.

If one of the following warning messages appears, first check the patient, then check the ventilation equipment (if used), and then check the alarm limits settings (Table 11: Warning Messages, below).

Table 11: Warning Messages

Message	Possible Causes	Action
No Breath xxx !!!*	No valid breath has	
	been detected for xxx	
	seconds.	
EtCO₂ ↑!!!	The EtCO ₂ exceeded	First check the
	the EtCO ₂ high alarm	patient, then the
	limit.	connections from
EtCO ₂ ↓ !!!	The EtCO ₂ fell below the	patient to the
	EtCO ₂ low alarm limit.	monitor, then
RR ↑ !!!	The RR exceeded the	ventilation
	RR high alarm limit.	equipment (if used),
		and then the alarm
		settings (refer to the
		Alarm Limits Menu
		on page 35).
RR ↓ !!!	The RR fell below the	
	RR low alarm limit.	
FiCO ₂ !!! = xx**	The FiCO ₂ exceeded the	
	FiCO ₂ high alarm limit.	

- * xxx= the number of seconds elapsed since the last valid breath has been detected.
- ** The FiCO₂ value is displayed if selected in the Instrument Settings menu 1 (Refer to Instrument Settings Menus on page 38).

Cautions

Caution messages appear during Measuring mode and indicate that a problem has occurred requiring the operator's attention. The message appears on the screen followed by !!, the alarm bar will flash yellow and a special repetitive caution tone is heard (see Table 12: Caution Messages, below).

Table 12: Caution Messages

Message	Possible Causes	Action
Check Unit !!	Instrument fault.	Contact authorized
		service
		representative.
Battery √ !!	Message appears when	Prepare to replace
	battery charge level is very	or recharge battery
	low (approximately 15	or connect monitor
	minutes left).	to AC power.
FilterLine !!	FilterLine is disconnected	Connect FilterLine to
	or not securely connected	CO ₂ input connector
	to the monitor.	or tighten
		connection.
Blockage !!	FilterLine is twisted or	Check the FilterLine
	clogged. The message	and, if necessary,
	appears after 30 seconds	replace it.
	of unsuccessful clearing of	Check the airway
	the FilterLine.	adapter and, if
	FilterLine airway	necessary, replace
	connection is clogged.	the FilterLine.

Advisory Messages

Troubleshooting

Advisories are informative messages appearing at start-up before any patient input has been detected by the monitor, or during operation. The message appears on the screen followed by !. The alarm bar will light yellow and a special one-time advisory tone is heard (see Table 13: Advisory Messages, below).

Table 13: Advisory Messages

Message	Possible Causes	Action
Check Unit!	Instrument fault.	Contact authorized
	service	
		representative.
Battery Empty!	Battery pack is	Replace or
	discharged.	recharge the
		battery, or connect
		to AC power.
Pump-Off xxx	*The pump is currently	Restart pump-off
	off.	timer by one long
		press 🗗
Battery ↓ !	Message appears	Prepare to replace
	when battery charge	or recharge the
	level is low	battery, or connect
	(approximately 40	to AC power.
	minutes left).	

^{*} xxx is the remaining time in seconds until the pump turns back on.

Silent Advisories

Silent Advisories are instrument status messages indicating the operational state of the monitor or consumables. Silent advisories are low priority signals and only a message appears (with no exclamation marks, nor any other visual or audible indicator) (see Table 14: Silent Advisory Messages, below).

Table 14: Silent Advisory Messages

Message	Possible Causes	Action
Pump-Off	The pump is currently off.	Activate the pump
		by one long press .
Clearing	FilterLine tube twisted or	Check the FilterLine
FilterLine	clogged.	and, if necessary,
		untwist it or replace
		it.
FilterLine	FilterLine is not	Connect FilterLine
	connected to the	to input connector.
	instrument.	
Autozero	Monitor automatically	No action required.
	performs a zero-point	
	calibration.	
CO ₂	CO ₂ module is preparing	Wait for "Ready"
Warm-up	itself for operation.	message before
		measuring for
		EtCO ₂ .
		No action required.
Calibration	Monitor requires	Calibrate Unit
Required	calibration.	

Demo	User mistakenly activated	Reset the monitor
	Demo mode	by sliding the on/off
		switch to off position
		and then to on
		position
BTPS On	BTPS setting is on.	No action required.
Ready	CO ₂ module is	
	operational but breath is	
	not detected.	
	Note: If BTPS is set to	
	OFF, only Ready	
	appears.	
FiCO ₂ = xx	The FiCO ₂ value (xx	No action required.
	mmHg or x.x Vol% or	
	kPa). Activated by user.	
6 sec	Patient setting for Adult	No action required.
	mode, or respiration rate	
	is low.	
3 sec	Patient setting for	No action required.
	Neonate Mode, or	
	respiration rate is high.	
Press To	Trend screen displayed	No action required.
Erase	(CO ₂ Trend-8 hrs and	(To erase trends,
	CO ₂ Trend-30 min.)	press and hold
		until message
		disappears.)

Troubleshooting Guide

In Table 15: Troubleshooting Guide on page 55 are potential problems you may experience while using the monitor and suggestions for

resolving them. If you are unable to correct the problem, contact qualified service personnel or your local representative.

Table 15: Troubleshooting Guide

Problem	Cause	Action
Monitor does not	Power cable	Check power cable
turn on.	improperly attached	connection and check
	or disconnected, or	that on/off switch is on.
	cable has faulty	
	electrical connection.	
	Battery pack may be	Replace or recharge
	discharged.	the battery pack, or
		connect to AC power.
	Battery pack may not	Be sure the battery
	be inserted properly	pack is in the monitor
	or missing.	and inserted properly.
Monitor switches	Electrical connection	Check connections
on but then	is faulty, or the AC	and correct problem.
switches off	wall outlet has no	
automatically.	power.	
	The battery pack is	Replace or recharge
	almost discharged.	battery pack, or
		connect to AC power.
	One of the monitor	If previous actions are
	subsystems is out of	not effective, contact
	order.	authorized service
		representative.

EtCO ₂ values read	Mechanically	No action needed.
erratically.	ventilated patient who	
	breathes	
	spontaneously.	
	A leak in the airway.	Check for connection
		and line leaks to patient
		and correct if necessary.
EtCO ₂ values are	Physiological cause.	Check patient.
consistently higher	Ventilator malfunction.	Check ventilator and
or lower than		patient.
expected.	Improper calibration.	Check calibration. See
		CO2 Calibration Check
		on page 59.
EtCO ₂ values are	BTPS setting ON or	Check BTPS setting on
consistently higher	OFF.	the graphic display after
or lower than	Note: When BTPS is	Power on. Contact your
expected.	on the correction	local service
	lowers the EtCO ₂	representative.
	reading to	
	compensate for Body,	
	Temperature,	
	Pressure, and	
	Saturation.	





Maintenance

Periodic Maintenance

Service

Cleaning

Calibration

CO2 Calibration Check

Returning the Monitor

Technical assistance

Periodic Maintenance

Periodic Maintenance is recommended according to operating hours:

The Pump and Flow System should be replaced every 7,000 operating hours.

The monitor should be returned to the manufacturer for periodic maintenance every 14,000 operating hours.

As part of routine preventative maintenance, a calibration check should be performed with safety checks as outlined by hospital protocol.

To check the monitor's operating hours, go to the information screen in the Service mode. Table 16: Accessing the Service Mode on page 58 describes how to access the information screen in the Service mode.

The battery pack should be replaced once every two years.

Maintenance

Table 16: Accessing the Service Mode

Objective	Action	Result
To access the Service mode	During Self Test, press and hold simultaneously	Microstream [®]
	■ and △	Self Test Service Mode Board No 12345 Operating Hours 5,783 Firmware V X.XX

Note:

Contact your local distributor to order spare parts, calibration kits, or to answer any questions regarding periodic maintenance.

Service

The monitor requires no routine service other than any performance testing mandated by the operator's institution. Troubleshooting Guide on page 54 discusses potential difficulties, their possible causes, and suggestions for resolving them. Contact your local distributor for service instructions, and performance tests and checks.

CAUTION: The monitor must be returned for repair if the "Check Unit" message appears.

Cleaning

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To clean the monitor's surfaces, dampen a cloth with a commercial, nonabrasive cleaner and wipe the top, bottom, and front surfaces lightly.

CAUTION: Do not spray or pour any liquid directly on the monitor,

accessories, or consumables.

CAUTION: Do not use caustic or abrasive cleaners.

Cal			

CAUTION:	Microstream EtCO ₂ consumables are designed for single
	patient use and are not to be reprocessed. Do not attempt
	to disinfect or flush the FilterLine as the monitor can be
	damaged.

Calibration

Calibrate after 1,400 hours of initial use. After that, calibration should be performed any time the monitor displays the advisory message Calibration Required. Calibration should be performed annually or after 4,000 hours, whichever comes first, by qualified service personnel.

Note: It is recommended that you calibrate the monitor

within two weeks of the message appearing on the

monitor.

Note: The monitor is calibrated when it leaves the factory.

CO₂ Calibration Check

The process should be performed only after the device has been operating for at least 20 minutes in a normal operating mode and connected to a FilterLine.

The calibration check must be performed with a manufacturer authorized Calibration Kit containing 5% CO₂ gas and the connecting means. A manufacturer approved Calibration Kit can be purchased from your Technical Services representative or from Scott Medical (part number 0304653ORFBD). The kit consists of:

- Calibration Gas containing 5% CO₂, 21% O₂
- Tubing Adapter
- Calibration Line

CAUTION: Do not check CO₂ values from the 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.

Maintenance

CAUTION:	The instrument should not be in Standby Mode before beginning the calibration check process. To prevent the device from entering Standby Mode, measure at least two breaths. The device will then remain in normal operating mode with an active Apnea (for software versions prior to 2.7) or No Breath (from software version 2.7) alarm.
Note:	If this process is performed while a battery powers the device, make sure that the battery is fully charged.
Note:	Prior to calibration, verify that the FilterLine supplied with the Calibration Kit is firmly attached.

Start the process from the Setup menu as described in Table 17: CO₂ Calibration Check, below.

Table 17: CO₂ Calibration Check

Objective	Action	Result
Access Instrument Settings menu 2. Change option to start.	long press (x3) (long press x2 for software versions prior to 2.7)	Check Cal. Off Language English Check Cal. Start Language English
Start Check Cal. (An Autozero process begins.)	short press	Autozero Please Wait

Returning the Monitor

Start Cal. Check process.	Connect the calibration gas via the connecting means.	Connect Gas Cal, Check
Check the measured values (shown in Vol% in the EtCO ₂ digital display).*	Press the gas valve for 15 seconds until the readings stabilize.	Connect Gas Cal. Check
	required if the measured ration of the calibration	
To return to Measuring mode if calibration is not required.	long press	40 mmHg 5 sec
If calibration is required, contact your local service representative.		

Returning the Monitor

If it is necessary to return the monitor for repairs, call Technical Services or your local representative for shipping instructions.

To repack the monitor, disconnect the consumable from the instrument and wrap each item separately. Pack it in the original shipping carton. If the original carton is unavailable, use a suitable box filled with the appropriate amount of packing material. It is not necessary to return the Microstream $EtCO_2$ consumables or power cords.

If the monitor malfunctions, carefully package the monitor with the consumable used at the time of malfunction and return it with the monitor for inspection.



Technical assistance

For technical information, contact Technical Services or your local representative.

The *Service Manual* includes information that is required by qualified personnel to service the monitor.

Specifications

Physical

Environmental

Safety Standards

Compliance

Manufacturer's Declaration

Performance

Power Specifications

Electrical

Components and User Interface

Physical

Size

206 mm H x 88 mm W x 52.5 mm D (8.11"H x 3.46" W x 2.06"D)

Weight

750 grams (1.66 lb.) (including battery pack)

Noise Emission

Maximum 45 dB(A)

Environmental

Temperature

Operating	0°C to 45°C (32°F to 113°F)
Relative Humidity	10 to 95% (noncondensing)
Storage	-35°C to 70°C (-31°F to 158°F)

Specifications

Pressure and Altitude (for operating and storage)

Pressure	430 mmHg to 795 mmHg	
Altitude	-380m to 4,570m (-1,250 ft. to 15,000 ft.)	

Transport and Storage

Parameter	Value	
Temperature	For Monitor: -35°C to 70°C (-31°F to 158°F) not	
	in shipping container	
	For Microstream Accessories:	
	-20°C to 70°C (-4°F to 158°F)	
	in shipping container	
Altitude	-380m to 4,570m	
	(-1,250 ft. to 15,000 ft.)	
Atmospheric Pressure	50 kPa to 106 kPa	
	(14.7 in Hg. To 31.3 in. Hg)	
Relative Humidity	10% to 95% non-condensing	

Safety Standards

The monitor was designed to comply with EN60601-1, UL 60601-1 and CSA C22.2 No. 601.1-M90, ISO 21647.

Compliance

Item	Compliant With	
Equipment	Safety Standards: IEC 60601-1 (same as	
classification	EN60601-1), CSA 601.1, UL 60601-1, ISO	
	21647, and EN/IEC 60601-1-2.	
Type of protection	Class I or II (on AC power)	
	Internally powered (on battery power)	
Degree of protection	Type BF – Applied part	
Mode of operation	Continuous	

	Сотрна
Item	Compliant With
Resistant to liquid	IEC 60601-1, sub-clause 44.6 for class
ingress	IPX1 Drip-Proof equipment
Degree of safety in	UL 60601-1, sub-clause 5.5, Not suitable
presence of a	
flammable anesthetic	
Applied sensor label	IEC 60601-1 Symbol 2 of Table DII of
to indicate Type BF	Appendix D
applied part	
Attention symbol,	IEC 60601-1 Symbol 14 of Table DI of
consult accompanying	Appendix D
documentation	
External case made	IEC 60601-1, sub-clause 16(a)
with non-conductive	
plastic	
No holes in case top	IEC 60601-1, sub-clause 16(b)
Rigid case	IEC 60601-1, sub-clause 21(a)
Case mechanically	IEC 60601-1, sub-clause 21(b)
strong	
Resistant to rough	IEC 60601-1, sub-clause 21.6
handling	
Tip/tilt test	IEC 60601-1, sub-clause 24.1
Resistant to liquid	IEC 60601-1, sub-clause 44.3 as modified
ingress due to spills	by ISO 9919 clause 44.6
Environmental	IEC 60601-1, sub-clause 44.5
Cleaning	IEC 60601-1, sub-clause 44.7
Case surface made of	IEC 60601-1, sub-clause 48
non-toxic materials	
Case resistant to heat	IEC 60601-1, sub-clause 59.2 (b)
and fire	
Exterior markings	IEC 60601-1, sub-clause 6.1., 6.3, and 6.4;
	ISO 9919, clause 6

Decinications	
ltem	Compliant With
Front panel and case	IEC 60878, EN 980, ISO 7000, EN 60417-
labeling	1, EN 60417-2
Button spacing	ISO 7250
Year of manufacture	EN 980
symbol	
Conductive coating	UL 60601-1, clause 55
and polymeric	
materials	
Operation during	IEC 60068-2-27 at 100 g
physical shock	
Operation during	IEC 60068-2-6 and IEC 60068-2-34
vibration	
Electromagnetic	IEC 60601-1, sub clause 36, IEC/EN
compatibility	60601-1-2 (second edition)
Radiated and	EN 55011, Group 1, Class B
conducted emissions	
Harmonic emissions	IEC 61000-3-2
Voltage	IEC 61000-3-3
fluctuations/flicker	
emissions	
Electrostatic	EN 61000-4-2, level 3 table top equipment
discharge immunity	
Radiated radio-	IEC 61000-4-3 at 3 V/m
frequency	
electromagnetic field	
immunity	
Electrical fast	IEC 610004-4-4
transient/burst	
immunity	
Surge immunity	IEC 61000-4-5

Manufacturer's Declaration

Item	Compliant With	
Conducted EMI	IEC 61000-4-6 at 3 V/m	
susceptibility		
Power frequency	IEC 61000-4-8 at 3A/m	
magnetic fields		
Laser Safety	The sensor LED light output falls within	
	Class I level, according to 60825-1:2001.	
	No special safety precautions are required.	
Operation with line	IEC 61000-4-11	
voltage variations		

Manufacturer's Declaration

WARNING:	The use of accessories and cables other than those
	specified may result in increased emission and/or

decreased immunity of the equipment and/or system.

Performance

<u> </u>		
Sampling Rate	50 ml/min.	
CO ₂ Range	0-99 mmHg (0-13.2 kPa and 0-13.0 Vol%)	
	at sea level	
Accuracy		
EtCO ₂ readings	From power-up until steady state is reached,	
	the CO ₂ reading accuracy is:	
	0 - 38 mmHg: (±4 mmHg)	
	39 - 99 mmHg: (±12% of reading)	
	The CO ₂ reading reaches its steady state	
	accuracy 20 minutes after power up.	
	0 - 38 mmHg: (±2 mmHg)	
	39- 99 mmHg: (±5% of reading + 0.08% for	
	every 1 mmHg above 40mmHg)	
	Equivalent values for kPa and Vol%	



Specifications

Joindations		
Respiration Rate	0-150 breaths/min.	
Warm-up Time	30 seconds (typical)	
Frequency	EtCO ₂ accuracy is maintained up to 80	
Response	breaths/min. (For maintaining accuracy for	
	respiration rate over 60 bpm, use the	
	neonatal mode.) From 81 to 150 bpm	
	accuracy is ±12%, if the EtCO ₂ is higher	
	than 18.8 mmHg in neonatal mode.	
System Response	2.45 seconds (typical), 2.9 seconds	
Time	maximum (includes delay and rise time)	
Rise Time		
Neonate	190 msec with low dead space endotracheal	
	tube adapter	
Adult	240 msec with FilterLine airway adapter	
Ambient Pressure	Compensated internally - automatic	
Alarms	EtCO ₂ high, EtCO ₂ low, RR, FiCO ₂ high, No	
	Breath	

Display Update Interval

2 seconds

Power Specifications

External Power Source

12V DC Medical Grade Adapter

Internal Power Source

Ni-MH Rechargeable Battery Pack 7.2V 2.1 A/h (intended for continuous operation)

Operating Time	Between 4 and 7 hours, depending on power	
(fully charged)	management. These values reflect the	
	performance of a new battery; age and usage	

	2.00
	will decrease capacity.
	Note: If the battery pack is stored for 6
	months or longer, you must charge and
	discharge it (leave unit on, not connected to
	AC power, until battery is empty) three times
	before use in order to ensure full capacity.
Recharging Period	Approximately 4.5 hours internal recharging
Charger Type	Internal

Electrical

Instrument

Rated 100-250VAC, 50/60HZ, 0,5A

Electromagnetic Emissions

The monitor is suitable for use in the specified electromagnetic environment. The user of the monitor should ensure that it is used in an electromagnetic environment as described below:

Emissions Test	Compliance	Electromagnetic Environment Guidance
RF emission	Group 1	The monitor uses RF energy
		only for its internal function.
CISPR 11		Therefore, its RF emissions
		are very low and are not
		likely to cause any
		interference in nearby
		electronic equipment.

Specifications

Emissions Test	Compliance	Electromagnetic Environment Guidance
RF emissions	Class B	The monitor is suitable for
		use in all establishments,
CISPR 11		including domestic
		establishments and those
		directly connected to the
		public low-voltage power
		supply network that supplies
		buildings used for domestic
		purposes.
Harmonic	Class A	
emissions		
IEC 61000-3-2		
Voltage	Complies	
fluctuations/ flicker		
emission		
IEC 61000-3-3		

Electromagnetic Immunity

The monitor is suitable for use in the specified electromagnetic environment. The user of the monitor should assure that it is used in an electromagnetic environment as described below.

Immunity Test	IEC 60601- 1-2 Test Level	Compliance Level	Electromagnetic Environment Guidance
Electrostatic	±6 kV	±6 kV	Floor should be
discharge	contact	contact	wood, concrete or

			Eleci
Immunity Test	IEC 60601- 1-2 Test Level	Compliance Level	Electromagnetic Environment Guidance
(ESD)	±8 kV air	±8 kV air	ceramic tile. If
IEC 61000-4-2			floors are covered
			with synthetic
			material, the
			relative humidity
			should be at least
			30%.
Electric fast	±2 kV for	±2 kV for	Mains power
transient/burst	power	power supply	quality should be
IEC 61000-4-4	supply lines	lines	that of a typical
	±1 kV for	±1 kV for	commercial
	input/output	input/output	and/or hospital
	lines	lines	environment
Surge	±1 kV	±1 kV	Mains power
IEC 61000-4-5	differential	differential	quality should be
	mode	mode	that of a typical
			commercial
			and/or hospital
			environment
	±2 kV	±2 kV	
	common	common	
	mode	mode	
Voltage dips,	<5% UT	<5% UT	Mains power
short			quality should be
nterruptions	(>95% dip in	(>95% dip in	that of a typical
and voltage	UT) for 0.5	UT) for 0.5	commercial
variations on	cycle	cycle	and/or hospital

Immunity Test	IEC 60601- 1-2 Test Level	Compliance Level	Electromagnetic Environment Guidance
power supply input lines. IEC 61000-4- 11	40% UT (60% dip in UT) for 5 cycles	40% UT (60% dip in UT) for 5 cycles	environment. If the user of the monitor requires continued operation during power mains interruption, it is recommended
	70% UT (30% dip in UT) for 25 cycles <5% UT (95% dip in UT) for 5 sec.	70% UT (30% dip in UT) for 25 cycles <5% UT (95% dip in UT) for 5 sec.	that the monitor be powered from an uninterruptible power supply or 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.

 U_T is the AC mains voltage prior to application of the test level.

Recommended Separation Distances

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

Rated maximum	Separation distance according to frequency of transmitter		
output power of transmitter	150 kHz to 80	80 MHz to	800 MHz to 2.5
W W	MHz	800 MHz	GHz
	d=1.2√ P	d=1.2√ P	d=2.3√ P
0.01	0.12	0.12	0.23
0.1	0.38	0.38	0.73
1	1.2	1.2	2.3
10	3.8	3.8	7.3
100	12	12	23

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

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

Note 2: These guidelines may not apply in all situations.

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

Electric and Communication Cables

Cables	Maximum Length	Complies with:
791001, Power cord	10 ft. (3 m)	RF emissions, CISPR
North America		11, Class B/ Group 1
RJ11	10 ft. (3 m)	Harmonic emissions.
communication		IEC 61000-3-2
cable (contained in		Voltage
048127		fluctuations/flicker
communication		emission. IEC 61000-3-
adapter kit)		3
RJ45	1.7 ft. (0.5 m)	Electrostatic discharge
communication		(ESD), IEC 61000-4-2
cable (contained in		Electric fast
048127		transient/burst, IEC
communication		61000-4-4
adapter kit)		Surge, IE 61000-4-5
15 pin D-type	10 ft. (3 m)	Conducted RF IEC
output connector		61000-4-6
cable (contained in		Radiated RF, IEC
063755 D/A		61000-4-3
Converter kit)		
RS232 monitor	1.7 ft. (0.5 m)	
cable (contained in		
063755 D/A		
Converter kit		
060606, 12 VAC	2.3 ft (0.7 m) not	
adapter cable	extended	



Components and User Interface

Displays

Graphic LCD	(128 x 64 dots) with LED backlight dimension	
display	75 mm x 53 mm.	
Two numeric fields	3 digits each, using 7-segment LED	
	dimension 22 mm x 14 mm.	
Alarm bar	yellow, red	

Controls and Indicators

Front Panel	On/Off switch; Alarm Silence/Alarm Silence
	Menu button; Contrast/Value change button;
	Event/Home button; Next/Menu button.

Connections

Front Panel	CO ₂ Input connector	
Rear Panel	Clamp connector	
Side Panel	Power supply/Communication Adapter port,	
	Gas outlet	

