R Series®
End Tidal Carbon Dioxide
(EtCO₂)
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Indication of use

The R Series system is indicated for the continuous noninvasive monitoring of end tidal carbon dioxide (EtCO₂) and respiration rate in patients requiring ventilator support, in-hospital transport, or anesthesia. EtCO₂ Monitoring is indicated for in patients from newborn (neonate) to adult.

Contraindications for Use

There are no known contraindications for the use of the EtCO2 monitor.

General Information

Federal (U.S.A.) law restricts this defibrillator to sale by or on the order of a physician.
End-Tidal Carbon Dioxide (EtCO₂)

Product Description

R Series® units equipped with software revision 12.xx or higher support two End Tidal Carbon Dioxide (EtCO₂) monitoring options for the continuous measurement of respiratory carbon dioxide (CO₂) and respiration rate. These options use the same connector on the R Series unit and may be used interchangeably.

For R Series BLS/Plus models, these EtCO₂ options can only be used while the unit is in Manual mode.

The first option uses a unique, mainstream, solid-state, infrared sensor called the CAPNOSTAT® 5 Mainstream CO₂ Sensor. The CAPNOSTAT 5 CO₂ sensor is attached to an airway adapter that connects to an endotracheal (ET) tube or other airway and measures gases flowing through these breathing circuit components. A disposable mouthpiece may be connected to the adapter for monitoring non-intubated patients. A CAPNO₂mask™ is also available for use with non-intubated patients. This option provides for O₂ delivery while monitoring expired CO₂.

The second option is a sidestream sampling system called the LoFlo™ CO₂ Module. The LoFlo module contains a gas sampling pump, which draws small samples of gas from the patient’s airway via a nasal/oral cannula or airway adapter, and passes these gases through a solid state infrared sensor (located away from the patient’s airway) that measures CO₂. While the sidestream system is typically used on non-intubated patients, it can also be used for EtCO₂ measurement on intubated infant, pediatric and adult patients. The sidestream system should not be used, however, on patients who cannot tolerate the 50ml/min removal of the sample gases from their breathing circuit. The sidestream module uses specially designed cannulas and airway adapters for sampling airway gases and passing them through an integrated sample cell, which connects to the LoFlo module’s CO₂ sensor. These cannulas incorporate a filter and sample cell, providing maximum filtration of fluids and contaminants, and protecting the system from aspiration of these fluids.

In both systems, the CO₂ sensor generates infrared light and beams it through the airway adapter or sample cell to a detector on the opposite side. CO₂ from the patient, flowing through the mainstream airway adapter or sample cell, absorbs some of this infrared energy. The R Series unit determines CO₂ concentration in the breathing gases by measuring the amount of light absorbed by gases flowing through the airway or sample cell.

The R Series unit displays EtCO₂ (the concentration of carbon dioxide detected at the end of each exhalation) as a numerical value in millimeters of mercury (mmHg), percent (%), or kilopascals (kPa). In addition, the unit can display a capnogram. This capnogram is a valuable clinical tool that can be used to assess patient airway integrity and proper endotracheal (ET) tube placement. The unit calculates respiration rate by measuring the time interval between detected peaks of the CO₂ waveform. The technology differentiates between waveforms caused by breathing and those caused by cardiogenic oscillations and artifact.
How to Use This Manual

This section explains how to set up and use the R Series End Tidal Carbon Dioxide option. Important safety information relating to general use of the R Series End Tidal Carbon Dioxide monitor appears in the “Safety Considerations” section of this manual.

The R Series Operator’s Guide provides information operators need for the safe and effective use and care of the R Series unit. It is important that persons using this device read and understand all the information contained therein.

Please read both safety considerations and warnings sections thoroughly before operating your R Series unit.

All CAPNOSTAT 5 sensor, LoFlo module, airway adapter and cannula questions with regards to the Declaration of Conformity with European Union Directives should be directed to the authorized ZOLL representative:

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The Netherlands
+31 (0) 481 366410 Telephone
+31 (0) 481 366411 Telefax

Safety Considerations

Warnings

General

Carefully read the R Series Operator’s Guide and these operating instructions before operating the EtCO₂ monitoring option.

Ensure that the R Series EtCO₂ option is operated by qualified personnel only.

Do not use the R Series EtCO₂ option as an apnea monitor.

Do not immerse the R Series unit, patient cables, or sensors in water, solvents, or cleaning solutions.

If the accuracy of any reading is suspect, first check the patient’s vital signs by alternate means and then check the R Series EtCO₂ option for proper operation.

If an alarm condition occurs while the alarms are suspended, the suspended alarm indications will only be visual displays and symbols. No audio alarm indications will occur.

Elevated oxygen levels, nitrous oxide, or halogenated agents contained in the breathing gases may degrade the accuracy of measurements made with the R Series EtCO₂ option. Activate oxygen compensation if O₂ levels in excess of 60% are introduced. Activate N₂O compensation if nitrous oxide is introduced into the airway circuit.

Do not use the LoFlo module on patients who cannot tolerate the removal of 50ml/min of breathing gases from the airway.

The presence of Desflurane beyond 5% may positively bias the carbon dioxide reading by up to 3 mmHg. The presence of Xenon in the exhaled breath may negatively bias the reading by up to 5 mmHg.
Carefully route patient cabling to reduce the possibility of patient entanglement or strangulation.

Do not touch the bed, patient, or any equipment connected to the patient during defibrillation. A severe shock can result. Do not allow exposed portions of the patient’s body to come in contact with metal objects, such as a bed frame, as unwanted pathways for defibrillation current may result.

**CAPNOSTAT 5 and Accessories**

Always ensure the integrity of the patient breathing circuit after insertion of the airway adapter by verifying a proper CO₂ waveform (capnogram) on the monitor display.

Do not use CAPNOSTAT 5 or LoFlo sensors in the presence of flammable anesthetics or other flammable gases.

Do not attempt to open the sensor. An electrical shock hazard exists internally. Refer servicing to qualified personnel.

Do not operate the sensor when it is wet or has exterior condensation.

**Cautions**

CAUTION: Federal (U.S.A.) law restricts this device to sale, or use by or on the order of a licensed medical practitioner.

Use only ZOLL/Respironics Novametrix CAPNOSTAT 5 sensors and LoFlo modules, airway adapters, nasal and nasal/oral cannula sets with the R Series EtCO₂ option.

The device is protected against interference from radio frequency emissions typical of two-way radios and cellular phones (digital and analog) used in emergency service/public safety activities. Users should assess the device’s performance in their typical environment of use for the possibility of radio frequency interference from high-power sources. Radio Frequency Interference (RFI) may be observed as shifts in monitor baseline, trace compression, display brightness changes or transient spikes on the display.

Do NOT sterilize or immerse the CAPNOSTAT 5 CO₂ sensor or LoFlo module.

Do NOT reuse, disassemble, sterilize, disinfect, or clean the disposable airway adapter, airway adapter with mouthpiece, CAPNO₂ mask, nasal or nasal/oral sampling cannula sets, as system performance will be compromised. These items are intended for single patient use only.

Do NOT use a damaged sensor or airway adapter.

Do NOT use the device if it fails to operate properly.

Do NOT place the mainstream or sidestream airway adapters between the ET tube and the breathing circuit elbow, as this may cause patient secretions to accumulate in the adapter.

Position airway adapters with windows in a vertical, NOT a horizontal, position. This helps keep patient secretions from pooling on the windows.

Do NOT insert any object other than the sample cell into the sample cell receptacle on the LoFlo module.

Remove the LoFlo sample cell from the sample cell receptacle when not in use.

Clean or replace the airway adapter if excessive secretions are observed.
ZOLL Medical Corporation recommends that the airway adapter be removed from the circuit whenever aerosolized medication is delivered. The increased viscosity of the medications may contaminate the adapter windows, requiring premature cleaning or replacement of the adapter.

In order to eliminate the potential build up of CO₂ inside the storage bag, ensure that the LoFlo module exhaust tube vents gases away from the module environment.

To avoid injury to the patient, remove the nasal/oral cannula from the patient before cutting the oral cannula tip.

Do NOT apply tension to the sensor cable.

Periodically inspect the sampling tubing for the absence of kinks.

Monitor the capnogram for an elevated baseline. If an elevated baseline is observed, verify patient condition first. If the caregiver determines that the patient condition is not contributing to the elevated baseline, follow the instructions for zeroing the sensor or module detailed in this manual.

Do NOT store sensors, modules, airway adapters, or cannula sets at temperatures less than -40° C or greater than 70° C.

Do not operate CAPNOSTAT sensors at temperatures less than 0° C or greater than 45° C. Do not operate LoFlo modules at temperatures less than 0° C or greater than 40° C.

Do not use the LoFlo module on R Series units that have a software version lower than 12.xx.
EtCO₂ Intended Use

The ZOLL R Series EtCO₂ option with Respironics Novametrix technology is intended to be used for continuous noninvasive monitoring of end tidal carbon dioxide (EtCO₂) and respiration rate in patients requiring ventilator support, in-hospital transport, or anesthesia. The R Series EtCO₂ option with Respironics Novametrix technology supports two methods for continuous measurement of end tidal carbon dioxide (EtCO₂) and respiration rate.

The first method uses the CAPNOSTAT 5 Mainstream CO₂ sensor attached to an airway adapter that connects to an endotracheal tube, mask or disposable mouthpiece.

The second method uses the LoFlo CO₂ module to monitor both non-intubated and intubated patients using specially designed sampling cannulas and airway adapters.

The R Series EtCO₂ option is designed to monitor adult, pediatric, and neonatal patients.

The following substances can influence CO₂ measurements made with the CAPNOSTAT 5 CO₂ sensor:

- elevated oxygen levels
- nitrous oxide
- halogenated agents

The R Series EtCO₂ option provides settings for high oxygen and/or nitrous oxide compensation. Halogenated anesthetic agents alter CO₂ readings, but the R Series unit will monitor CO₂ within specifications when these agents are present at normal clinical levels. The presence of Desflurane in the exhaled breath beyond normal values (5%) may positively bias measured carbon dioxide values up to an additional 3 mmHg.

The R Series EtCO₂ option is intended for use only with the ZOLL/Respironics Novametrix CAPNOSTAT 5 Mainstream CO₂ Sensor and mainstream airway adapters, the LoFlo CO₂ Module, nasal and nasal/oral sampling cannula sets, and sidestream on-airway adapters. The R Series EtCO₂ option can be used on adult patients (21 years of age and older) and on pediatric patients, as described in the following table:

<table>
<thead>
<tr>
<th>Pediatric Subpopulation</th>
<th>Approximate Age Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Newborn (neonate)</td>
<td>Birth to 1 month of age</td>
</tr>
<tr>
<td>Infant</td>
<td>1 month to 2 years of age</td>
</tr>
<tr>
<td>Child</td>
<td>2 to 12 years of age</td>
</tr>
<tr>
<td>Adolescent</td>
<td>12-21 years of age</td>
</tr>
</tbody>
</table>

Mainstream EtCO₂ Setup

There are several steps involved with mainstream EtCO₂ setup. These steps include:

- Attaching the CAPNOSTAT sensor cable.
- Selecting a mainstream airway adapter.
- Attaching the airway adapter to the CAPNOSTAT sensor.
- Zeroing the CAPNOSTAT sensor/airway adapter.
- Attaching the airway adapter to the airway circuit.
- Applying an airway adapter with mouthpiece.
Attaching the CAPNOSTAT 5 CO₂ Sensor Cable

To attach the CAPNOSTAT 5 CO₂ sensor cable, plug the cable’s connector into the yellow CO₂ connector at the back of the R Series unit by matching the key on the cable to the key on the connector (Figure 1).

![Proper Grasp vs. Improper Grasp](image)

**Figure 1**

**Note:** To remove the sensor cable from the R Series unit, grasp the collar surrounding the cable’s R Series connector and pull up.

Selecting a Mainstream Airway Adapter

Select an airway adapter based on the patient’s ET tube diameter and monitoring situation. For more information refer to the following table or contact ZOLL Medical Corporation.

<table>
<thead>
<tr>
<th>Airway Adapter Type</th>
<th>ET Tube Diameter</th>
</tr>
</thead>
<tbody>
<tr>
<td>SPU* Pediatric/Adult</td>
<td>&gt; 4.0 mm</td>
</tr>
<tr>
<td>Adult Reusable</td>
<td>&gt; 4.0 mm</td>
</tr>
<tr>
<td>SPU* Neonatal/Pediatric</td>
<td>≤ 4.0 mm</td>
</tr>
<tr>
<td>Neonatal Reusable</td>
<td>≤ 4.0 mm</td>
</tr>
</tbody>
</table>

*SPU = Single Patient Use

Attaching the Airway Adapter to the CAPNOSTAT 5 CO₂ Sensor

Before attaching the airway adapter to the CAPNOSTAT 5 CO₂ sensor, verify that the airway adapter windows are clean and dry. Clean or replace the adapter if necessary.

**Caution**

The disposable (SPU) Pediatric/Adult and the Neonatal/Pediatric airway adapters are intended for single patient use. Do NOT reuse or sterilize these adapters as system performance will be compromised.

Attach the airway adapter to the CAPNOSTAT sensor, as follows:

1. Align the arrow on the bottom of the airway adapter with the arrow on the bottom of the sensor.
2. Press the sensor and airway adapter together until they click.
3. Turn the Selector switch on the R Series unit to **MONITOR (ON** for BLS/Plus units, and select **Manual** mode).
4. Wait for the airway adapter and sensor to warm up.
   The unit will display the **CO2 WARM UP** message for approximately one minute while the sensor and adapter warm to operating temperature. The message disappears when the sensor is ready to use.
   **Note:** Warm up time varies with ambient temperature of the sensor.
5. If the unit displays the **CHECK CO2 ADAPTER** message, follow steps a through c.
   a. Verify proper connection of the adapter to the sensor.
   b. Verify that the airway adapter windows are clean and dry.
   c. If the adapter is properly connected, and the windows are clean and dry, then zero the adapter as described in the next section, “Zeroing the Mainstream CAPNOSTAT 5 CO₂ Sensor/Airway Adapter.”

### Zeroing the Mainstream CAPNOSTAT 5 CO₂ Sensor/Airway Adapter

**Note:** Do not zero the CAPNOSTAT without an airway adapter installed.

Adapter zeroing compensates for the optical differences between airway adapters and should be performed at certain times. Zeroing is recommended after switching between single patient use and reusable airway adapters, in order to obtain accurate readings. It is also recommended the first time a particular CAPNOSTAT 5 CO₂ sensor is connected to the unit.

1. Place the sensor with the adapter installed away from all sources of CO₂ (including the patient’s – and your own – exhaled breath and ventilator exhaust valves).
2. Press the **Param** softkey, then the **CO2** softkey.
3. Press the **Zero** softkey.
   The unit zeroes the adapter and displays the **ZEROING CO2 ADAPTER** message for 15 to 20 seconds.
   The unit displays the message **ZERO DONE** upon completion of the zeroing.
   **Note:** Do not attempt zeroing for 20 seconds after removing the adapter from the patient’s airway. This time allows any CO₂ remaining in the adapter to dissipate before zeroing. Do not attempt to zero the adapter while it is connected to the patient’s airway. Zeroing with CO₂ in the adapter can lead to inaccurate measurement and/or other error conditions. If you attempt zeroing while CO₂ remains in the adapter, the time required to zero the adapter may be increased. If zeroing cannot be completed, the message **ZERO FAILED** will be displayed. If this occurs, clear any occlusion in the adapter, remove any source of CO₂, wait 20 seconds, and try zeroing again.

### Attaching the Airway Adapter to the Airway Circuit

If you have not yet done so, you must attach the airway adapter to the CAPNOSTAT 5 CO₂ sensor before attaching the breathing adapter to the airway circuit. Refer to “Attaching the Airway Adapter to the CAPNOSTAT 5 CO₂ Sensor” on page 6 if necessary.

Attach the airway adapter to the breathing circuit as follows:
1. Place the CAPNOSTAT 5 CO₂ sensor/airway adapter assembly between the elbow and the ventilator circuit wye, as shown in Figure 2.

**Note:** Do NOT place the airway adapter between the ET tube and the elbow, as this may allow patient secretions to accumulate in the adapter.

Position the airway adapter with its windows in a vertical, NOT a horizontal, position. This helps keep patient secretions from pooling on the windows. If pooling does occur, the airway adapter may be removed from the circuit, rinsed with water and reinserted into the circuit. To prevent moisture from draining into the airway adapter, do NOT place the airway adapter in a gravity dependent position.

2. Check that connections have been made correctly by verifying the presence of a proper CO₂ waveform on the R Series display.

3. The sensor cable should face away from the patient.

### Applying an Airway Adapter with Mouthpiece

The disposable Pediatric/Adult airway adapter with mouthpiece can be used for spot checking CO₂ on non-intubated adult or pediatric patients.

**Caution** The disposable Pediatric/Adult Airway Adapter with mouthpiece is intended for single patient use. Do NOT reuse or sterilize the adapter, as system performance will be compromised.

1. Remove adapter with mouthpiece from the package. Verify that the adapter and mouthpiece are intact and securely fastened to each other.

2. Attach the airway adapter to the CAPNOSTAT 5 CO₂ sensor, as follows:
   a. Align the arrow on the bottom of the airway adapter with the arrow on the bottom of the CAPNOSTAT sensor.
b. Press the sensor and airway adapter together until they click (see Figure 3).

![Figure 3](image)

3. If the unit displays the **CHECK CO2 ADAPTER** message, follow steps a through c, then go to step 4.
   a. Verify proper connection of the adapter to the sensor.
   b. Verify that the airway adapter windows are clean and dry.
   c. If the adapter is properly connected, and the windows are clean and dry, then zero the adapter as described in “Zeroing the Mainstream CAPNOSTAT 5 CO2 Sensor/Airway Adapter” on page 7.

4. Ensure the patient seals his or her mouth completely around the mouthpiece and breathes normally.
   A nose clip may be needed if the patient is exhaling through the nose. It is important that all, or most, of the exhalation be routed through the airway adapter.

**Sidestream EtCO2 Setup**

There are several steps involved with sidestream EtCO2 setup. These steps include:

- Attaching the LoFlo Module Cable
- Selecting a Sidestream Airway Adapter Kit or Cannula
- Inserting the Sample Cell
- Zeroing the LoFlo CO2 Module/Sample Cell
- Applying a Sidestream Airway Adapter Kit
- Applying a Nasal or Nasal/Oral Cannula

**Attaching the LoFlo Module Cable**

To attach the LoFlo module cable, plug the cable into the yellow CO2 connector at the back of the R Series unit by matching the key on the cable to the key on the connector.
To remove the sensor cable from the R Series unit, grasp the collar surrounding the cable’s R Series connector and pull up.

Selecting a Sidestream Airway Adapter Kit

Select an airway adapter kit based on the patient’s size, ET tube diameter, and monitoring situation. Airway adapter kits are disposable and single patient use.

<table>
<thead>
<tr>
<th>Airway Adapter Kit</th>
<th>ET Tube Diameter</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adult/Pediatric Airway Adapter Kit</td>
<td>&gt; 4.0 mm</td>
</tr>
<tr>
<td>Adult/Pediatric Airway Adapter Kit with Nafion® tubing</td>
<td></td>
</tr>
<tr>
<td>Pediatric/Infant Airway Adapter Kit</td>
<td>≤ 4.0 mm</td>
</tr>
<tr>
<td>Pediatric/Infant Airway Adapter Kit with Nafion tubing</td>
<td></td>
</tr>
</tbody>
</table>

**Note:** For monitoring times exceeding 6 hours, Nafion tubing is recommended.
Selecting a Sidestream Cannula

Select a sidestream cannula based on the patient’s size and monitoring situation. Nasal and nasal/oral cannulas are disposable and single patient use.

<table>
<thead>
<tr>
<th>Cannula</th>
<th>Application</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nasal CO₂ Sampling Cannula, Adult</td>
<td>Nasal CO₂ sampling only</td>
</tr>
<tr>
<td>Nasal CO₂ Sampling Cannula, Pediatric</td>
<td></td>
</tr>
<tr>
<td>Nasal CO₂ Sampling Cannula, Infant</td>
<td></td>
</tr>
<tr>
<td>Oral/Nasal CO₂ Sampling Cannula, Adult</td>
<td>Oral/Nasal CO₂ sampling only</td>
</tr>
<tr>
<td>Oral/Nasal CO₂ Sampling Cannula, Pediatric</td>
<td></td>
</tr>
<tr>
<td>Nasal CO₂ Sampling with O₂ Delivery Cannula, Adult</td>
<td>Nasal CO₂ sampling with oxygen delivery</td>
</tr>
<tr>
<td>Nasal CO₂ Sampling with O₂ Delivery Cannula, Pediatric</td>
<td></td>
</tr>
<tr>
<td>Oral/Nasal CO₂ Sampling with O₂ Delivery Cannula, Adult</td>
<td>Oral/Nasal CO₂ sampling with oxygen delivery</td>
</tr>
<tr>
<td>Oral/Nasal CO₂ Sampling with O₂ Delivery Cannula, Pediatric</td>
<td></td>
</tr>
</tbody>
</table>

Inserting the Sample Cell

Follow these steps:

1. Remove the LoFlo sampling cannula or airway adapter kit from the package.
2. Insert the LoFlo sample cell into the LoFlo sample cell receptacle and ensure that it clicks into place.
3. Ensure that the LoFlo module exhaust tube vents gases away from the module environment.
4. Turn the selector switch on the R Series to MONITOR (ON for BLS/Plus units).
5. Wait for the CO₂ module to warm up.
   The unit will display the *WARM UP* message for approximately one minute while the module warms up to operating temperature. The message disappears when the module is ready for use.

**Note:** Warm up time varies with ambient temperature of the module.

Zeroing the LoFlo CO₂ Module/Sample Cell

The module/sample cell zero allows the LoFlo CO₂ module to adjust to the optical characteristics of the sample cell. While zeroing is recommended the first time a particular
LoFlo module is connected to the unit, it is only absolutely necessary when the message ZERO CO2 MODULE is displayed.

**Caution** Always ensure that the sample cell is properly connected to the LoFlo module before zeroing.

1. Ensure that the nasal cannula or on-airway adapter is not connected to the patient or close to any source of CO2 (including the patient’s—and your own—exhaled breath and ventilator exhaust valves).
2. Press the **Param.** softkey and then the **CO2** softkey, then press **Enter**.
3. Press the **Zero** softkey.

   The unit zeroes the module and displays the **ZEROING CO2 MODULE** message for approximately 15-20 seconds.

   The units displays the message **ZERO DONE** upon completion of the zeroing.

**Note:** Do not attempt zeroing for 20 seconds after removing the adapter or cannula from the patient’s airway. This time allows any CO2 remaining in the adapter or cannula to dissipate before zeroing. Do not attempt to zero the module while the adapter or cannula is in the patient’s airway. Zeroing with CO2 in the adapter or cannula can lead to inaccurate measurements and/or other error conditions. If you attempt zeroing while CO2 remains in the adapter or cannula, the time required to zero the module may be increased. If zeroing cannot be completed, the message “ZERO FAILED” will be displayed. If this occurs, clear any occlusion in the adapter or cannula, remove the source of CO2, wait 20 seconds, and try zeroing again.

### Applying a Sidestream Airway Adapter Kit

The sidestream airway adapter kit is intended for monitoring the EtCO2 of intubated patients.

Before attaching the airway adapter to the breathing circuit, verify that the adapter is clean, dry, and undamaged. Replace if necessary.

**Caution** The disposable (SPU) Adult/Pediatric and Pediatric/Infant airway adapter kits are intended for single patient use. Do NOT reuse or sterilize these adapters as system performance will be compromised.

1. Attach the airway adapter kit’s sample cell to the sample cell receptacle on the LoFlo module, and ensure that it clicks into place.
2. If the unit displays either of the following messages take the appropriate action.
3. Place the airway adapter assembly at the proximal end of the airway circuit between the elbow and the ventilator circuit wye. Do NOT place the airway adapter between the ET tube and the elbow as this may allow patient secretions to accumulate in the adapter. If pooling does occur, the airway adapter may be removed from the circuit, rinsed with water and reinserted into the circuit. To prevent moisture from draining into the sample tubing, ensure that the sampling tube exits from the top of the airway adapter, not its bottom or sides. See Figure 6.

4. Check that connections have been made correctly by verifying the presence of a proper capnogram on the R Series display.

Applying a Nasal or Nasal/Oral Cannula

The nasal and nasal/oral cannulas are intended for monitoring EtCO₂ in non-intubated patients. Oral/nasal sampling cannulas should be used on patients who are prone to mouth breathing, since most (if not all) of the CO₂ is exhaled through the mouth. If a standard nasal CO₂ sampling cannula is used on such patients, the EtCO₂ values and capnogram displayed will be substantially lower than the actual CO₂ levels present in the patient’s expired breath.

Caution

The disposable nasal and nasal/oral cannula sets are intended for single patient use. Do NOT reuse or sterilize the cannula, as system performance will be compromised.
1. Remove the cannula from the package. Verify that the cannula is clean, dry, and undamaged. Replace if necessary.

2. Attach the cannula’s sample cell to the sample cell receptacle on the LoFlo module, and ensure that it clicks into place.

3. If the unit displays either of the following messages take the appropriate action.

<table>
<thead>
<tr>
<th>If you see this message:</th>
<th>Take this action:</th>
</tr>
</thead>
<tbody>
<tr>
<td>CHECK CO2 LINE</td>
<td>Verify that the sample cell is plugged into the module and seated properly.</td>
</tr>
<tr>
<td></td>
<td>Verify that neither the sample line nor the exhaust tube are blocked, kinked, or pinched. If the sample line,</td>
</tr>
<tr>
<td></td>
<td>or exhaust tube is blocked or otherwise compromised for 15 seconds, this message will appear. The pump will</td>
</tr>
<tr>
<td></td>
<td>shut off after 2 minutes if the condition that caused the message is not cleared. To restart the pump,</td>
</tr>
<tr>
<td></td>
<td>correct the blockage, then remove and reinsert the sample cell into the sample cell receptacle.</td>
</tr>
<tr>
<td></td>
<td>If the problem persists, replace the sample line.</td>
</tr>
</tbody>
</table>

| CHECK CO2 MODULE                | Check that module cable is plugged in and seated properly.                                                  |
|                                | Check that module is not exposed to excessive heat.                                                         |
|                                | If problem persists, replace module.                                                                          |

4. Place the nasal cannula onto the patient as shown in Figure 7.

![Figure 7](image)

5. Place the oral/nasal cannula onto the patient as shown in Figure 8.

![Figure 8](image)
Cleaning the CAPNOSTAT 5 CO₂ Sensor and LoFlo Module

The outside of the sensor or module may be cleaned and disinfected by wiping with 70% isopropyl alcohol, a 10% bleach solution, or mild soap.

After cleaning, wipe with a clean, water-dampened cloth to rinse. Dry before use.

Cleaning Reusable Airway Adapters

Reusable airway adapters may be cleaned by rinsing in a warm soapy solution, followed by soaking in a liquid disinfectant such as 70% isopropyl alcohol, a 10% bleach solution, Cidex® or System 1® (refer to the disinfectant manufacturer’s instructions for use). Adapters should then be rinsed with sterile water and dried.

The adult reusable airway adapter may be autoclaved at 121° C (250°F) for 20 minutes, unwrapped.

Note: The Neonatal reusable airway adapter cannot be autoclaved.

Before reusing the adapter, ensure the windows are dry and residue-free, and that the adapter has not been damaged during handling or by the cleaning process.

How EtCO₂ is Displayed

The R Series unit displays the numeric EtCO₂ value in units of mmHg, unless configured for percent or kPa. Refer to the R Series Configuration Guide (Part No. 9650-1201-01) for instructions on how to configure alternate units of measure. The unit also displays the number of breaths per minute, labeled “RR” for respiration rate. In addition, a capnogram may be displayed using the Trace 2 or Trace 3 softkey.

Displaying the Capnogram Waveform

The R Series unit can display 1, 2 or 3 waveforms in Monitor, Defib, or Manual (BLS/Plus models) mode, as long as the defibrillator is not charging or ECG analysis is not in progress. The unit displays only 1 or 2 waveforms in Pacer mode.

Note: If you don’t see the CO₂ display box on the monitor, check the sensor cable connection to the R Series unit. The CO₂ display box is not displayed if the sensor is not connected to the unit. Once the box is displayed, after power-on, it remains displayed even if the sensor is disconnected from the unit.
With EtCO₂ monitoring, the unit can display a capnogram below the ECG trace for a visual indicator of the moment-by-moment CO₂ values. The unit displays the capnogram at half the speed of the ECG display, and provides 8 or 10 seconds of data, depending on the setup.

The unit removes the third waveform from the display when the user presses the CHARGE, ANALYZE, or ENERGY SELECT buttons, or the Sync On/Off softkey.

To cycle the display from the capnogram to other waveforms (SpO₂, CPR, or Filt ECG) press the Options softkey, then the Traces softkey. Select Trace 2 or Trace 3 and then the desired waveform. Press Off to remove any additional waveforms.

Use the Zoom softkey from the EtCO₂ submenu to adjust the waveform display size. Numbers shown on the left side of the capnogram display indicate the scaling.

**Physiological Monitoring**

The physiological monitoring menu includes the following softkeys: Options, Param, Code Marker, Report Data, and Alarms.

**Param Softkey**

When you press the Param softkey, the following softkeys are displayed: ECG, SpO₂, NIBP, CO₂, and Return.

*Note:* SpO₂, NIBP, and CO₂ will only appear if these options are installed on your unit.

Press the Return softkey to return to the physiological monitoring menu.
Selecting the CO2 parameter causes the following softkeys to appear: Zero, Average, Comp., Zoom, Disable EtCO2 (or Enable EtCO2), and Return.

<table>
<thead>
<tr>
<th>Zero</th>
<th>RR Filter/Average</th>
<th>Comp.</th>
<th>Zoom</th>
<th>Disable EtCO2</th>
<th>Return</th>
</tr>
</thead>
</table>

The Zoom softkey will only appear if a capnogram is currently displayed.

Pressing the Disable EtCO2 softkey puts the sensor into the power saving sleep mode. Pressing Enable EtCO2 softkey takes the sensor out of sleep mode and turns the heater on for normal operation.

Press the Return softkey to return to the physiological monitoring menu.

### Zero Softkey

Adapter zeroing should be performed whenever you switch between reusable and disposable adapters, or if a CAPNOSTAT 5 CO2 sensor is being connected to the unit for the first time. Module zeroing may be necessary if the unit displays the message ZERO CO2 MODULE. Adapter zeroing may also be necessary if the unit displays ZERO CO2 ADAPTER.

**Note:** Do not attempt zeroing for 20 seconds after removing the adapter or cannula from the patient’s airway. This time allows any CO2 remaining in the adapter or cannula to dissipate before zeroing. Do not attempt to zero while the adapter or cannula is in the patient’s airway. Zeroing with CO2 in the adapter or cannula can lead to inaccurate measurements and/or other error conditions. If you attempt zeroing while CO2 remains in the adapter or cannula, the time required to zero may be increased. If zeroing cannot be completed, the message ZERO FAILED will be displayed. If this occurs, clear any occlusion in the adapter or cannula, remove any source of CO2, wait 20 seconds, and try zeroing again.

Press the Zero softkey to initiate adapter or module zeroing.

The unit displays the ZEROING CO2 ADAPTER or ZEROING CO2 MODULE message during the zeroing process, which is typically finished in 15-20 seconds.

The unit displays the ZERO DONE message when the zeroing process is complete.

The unit displays the ZERO FAILED message if the zeroing process did not complete successfully. If this occurs, clear any occlusion in the adapter or sample line, remove any source of CO2, and try zeroing again.

Press the Return softkey to return to the main menu.

### Zoom Softkey

The Zoom softkey will only appear if a capnogram is currently displayed. Select the full scale range for the displayed capnogram by scrolling among the different zoom levels. Zoom levels change with each press of the Zoom softkey. They are as follows:

- 0-12.5 mmHg
- 0-25 mmHg
- 0-50 mmHg
- 0-75 mmHg
• 0-100 mmHg
• 0-150 mmHg

If the unit is using kPa or %, the scales are 0-1.7, 0-3.3, 0-6.6, 0-10, 0-13.3, and 0-20.0. (Refer to the *R Series Configuration Guide* for instructions on how to configure alternate units of measure.)

**Average Softkey**

The R Series unit provides 3 different time periods over which EtCO₂ values are averaged: 1 breath, 10 seconds (default), and 20 seconds.

The user can select the averaging period by pressing the *Average* softkey. When the *Average* softkey is pressed, the unit displays the selections.

| 1 breath | 10 secs | 20 secs | RR Filter | Enable | Return |

Press the *1 breath*, *10 secs* or *20 secs* softkey for the desired time period.

**RR Filter Enable/Disable**

The respiration rate filter makes respiration rate counting more accurate in the presence of artifact, and is only valid when the LoFlo module is in use. Pressing the:

• *RR Filter Enable* softkey turns the respiration rate filter on.
• *RR Filter Disable* softkey turns the respiration rate filter off.

Note that when the RR filter is in use, the R Series adapts more slowly to sudden changes in respiratory rate.

Press the *Return* softkey to return to the main menu.

**Comp. Softkey**

The R Series unit can compensate for elevated levels of oxygen and/or the presence of nitrous oxide. Oxygen compensation should be activated when oxygen levels in excess of 60% are present in the airway circuit. Nitrous oxide compensation should be activated when nitrous oxide is present in the airway circuit. If the concentration of oxygen in the breathing circuit exceeds 60% and nitrous oxide is in use, both O₂ and N₂O should be activated.

When the *Comp* softkey is pressed, the unit displays the *None*, *O₂*, *N₂O*, *O₂&N₂O*, and *Return* softkeys.

| None | O₂ | N₂O | O₂&N₂O | Return |

The O₂ selection turns oxygen compensation on and displays an asterisk in the upper left of the CO₂ box. The N₂O selection turns nitrous oxide compensation on and displays an asterisk (*) to the right of the O₂ asterisk. The None selection turns all compensations off and eliminates the asterisks from the display.

The O₂&N₂O selection turns oxygen and nitrous oxide compensation on. The unit displays two asterisks (**) in the upper left of the CO₂ box to indicate compensation for both oxygen and
Alarms

The R Series EtCO₂ option provides user programmable “out-of-range” alarms for both EtCO₂ and respiration rate.

**Note:** The EtCO₂ and respiration rate alarms cannot be enabled or disabled separately. Enabling the EtCO₂ alarms enables both EtCO₂ and respiration rate alarm functions; disabling EtCO₂ or respiration rate alarms disables both functions. See the *R Series Operator's Guide* for details on enabling, disabling, and suspending alarm functions on the R Series unit.

When the EtCO₂ and respiration rate alarm states are set to AUTO, the unit automatically sets the lower and upper limits for EtCO₂ and respiration rate. The unit sets the upper and lower alarm limits to +/- 25% of the patient’s currently measured CO₂ value. If the EtCO₂ value is greater than 40 mmHg (equivalent to 5.3 kPa or 5.3% at a barometric pressure of 760 mmHg), then 10 mmHg (1.3 for kPa or %) is added and subtracted from the current reading to set the upper and lower limits. The auto alarm limits are set only if valid measurements are present for the vital sign.

For the automatic respiration rate alarm limits, the unit sets the upper and lower limits for respiration by adding and subtracting the values shown in the following table to/from the patient's current breath rate.

<table>
<thead>
<tr>
<th>Respiration Limits (Auto)</th>
<th>High Limit</th>
<th>Low Limit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Respiration Rate Average</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1-15 breaths/min.</td>
<td>+7 breaths/min.</td>
<td>-50% value</td>
</tr>
<tr>
<td>16-40 breaths/min.</td>
<td>+10 breaths/min.</td>
<td>-7 breaths/min.</td>
</tr>
<tr>
<td>&gt; 40 breaths/min.</td>
<td>+15 breaths/min.</td>
<td>-10 breaths/min.</td>
</tr>
</tbody>
</table>

See “Default Settings” on page 21 for low and high alarm limit default values and ranges.

Recorder Operation

If EtCO₂ measurements have been taken, press the RECORDER button to print a stripchart that includes the following values across the top part of the paper:

- Date and time
- ECG lead and size
- Heart rate
- EtCO₂ value
- Respiration rate

The recorder runs continuously until the button is pressed again. If selected, the capnogram is also printed at a fixed scale of 40 mmHg/cm or 5.3 kPa/cm. All waveforms printed by the recorder are delayed by 6 seconds relative to their occurrence.
Check Out Procedures

The following procedures verify that the EtCO₂ option is functioning properly.

Mainstream EtCO₂ (CAPNOSTAT 5 CO₂ Sensor)

1. Connect the CAPNOSTAT 5 CO₂ sensor cable to the yellow EtCO₂ connector at the back of the R Series unit.
2. Connect an airway adapter to the CAPNOSTAT 5 CO₂ sensor.
3. Turn the selector switch to **MONITOR** mode (ON for BLS/Plus units and select **Manual** mode).
   If the CO₂ box displays DISABLED, enable the sensor by pressing the **Param** softkey, then the **CO₂** softkey, then the **Enable EtCO₂** softkey.
4. Wait for the CO₂ sensor to warm up. The message **CO₂ WARM UP** is displayed for approximately one minute.
5. Perform a zero procedure if necessary (see “Zeroing the Mainstream CAPNOSTAT 5 CO₂ Sensor/Airway Adapter” on page 7).
6. Breath normally into the adapter.
7. Verify that the unit displays appropriate readings in the EtCO₂ display area of the monitor.
8. With alarms enabled, verify that the patient alarms are functional by adjusting the high and low limits until the unit:
   - Emits a continuous audio tone.
   - Highlights the alarming parameter’s value and flashes the alarm symbol on the display.

Sidestream EtCO₂ (LoFlo Module)

Use an Adult/Pediatric Airway Adapter kit when performing this procedure.

1. Connect the LoFlo module cable to the EtCO₂ connector at the back of the R Series unit.
2. Insert the sample cell into the LoFlo module sample cell receptacle.
3. Turn the selector switch to **MONITOR** mode (ON for BLS/Plus units and select **Manual Mode**), and wait approximately one minute while the module warms to operating temperature (unit displays **WARM UP** message).
4. Perform a zero procedure if necessary (see “Zeroing the LoFlo CO₂ Module/Sample Cell” on page 11).
5. Breath normally into the adapter.
6. Verify that the unit displays EtCO₂ readings in the EtCO₂ display area of the monitor.
7. Verify the capnogram is displayed by pressing the **Options** softkey, then the **Traces** softkey.
8. With alarms enabled, verify that the patient alarms are functional by adjusting the high and low limits until the unit:
   - Emits a continuous audio tone.
   - Highlights the alarming parameter’s value and flashes the alarm symbol on the display.
Default Settings

When the unit is turned on, the following default EtCO₂ settings are automatically selected and remain in operation until changed.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Default Setting</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Averaging Mode</td>
<td>10 seconds</td>
<td>1 breath, 10 seconds, 20 seconds</td>
</tr>
<tr>
<td>High EtCO₂ Alarm Limit</td>
<td>55 mmHg</td>
<td>5 - 100 mmHg, OFF, 0.6 - 13.1%, OFF, 0.6 - 13.3 kPa, OFF</td>
</tr>
<tr>
<td>Low EtCO₂ Alarm Limit</td>
<td>25 mmHg</td>
<td>0 - 95 mmHg, OFF, 0 - 12.5%, OFF, 0 - 12.6 kPa, OFF</td>
</tr>
<tr>
<td>High Respiration Rate Alarm Limit</td>
<td>120 respirations per minute</td>
<td>5 - 150 respirations per minute, OFF</td>
</tr>
<tr>
<td>Low Respiration Rate Alarm Limit</td>
<td>5 respirations per minute</td>
<td>0 - 100 respirations per minute, OFF</td>
</tr>
</tbody>
</table>

**Note:** The power-on default settings for the capnogram waveform scale and CO₂ compensation are set in System Configuration, as are the power-on default settings for alarm limits. See the *R Series Configuration Guide* for more information.

EtCO₂ Accessories

The following table lists the accessories available for the R Series mainstream EtCO₂ monitoring option.

Table 1-1. CAPNOSTAT 5 Mainstream CO₂ Accessories

<table>
<thead>
<tr>
<th>Accessory</th>
<th>REF</th>
<th>Application</th>
</tr>
</thead>
<tbody>
<tr>
<td>CAPNOSTAT 5 CO₂ Sensor and Cable</td>
<td>8000-0312-01</td>
<td>Reusable</td>
</tr>
<tr>
<td>SPU* Pediatric/Adult Airway Adapter</td>
<td>8000-0260-01</td>
<td>Single patient use, for ET tube sizes &gt; 4.0 mm</td>
</tr>
<tr>
<td>SPU* Neonatal/Pediatric Airway Adapter</td>
<td>8000-0261-01</td>
<td>Single patient use, for ET tube sizes ≤ 4.0 mm</td>
</tr>
<tr>
<td>Reusable Adult Airway Adapter</td>
<td>8000-0262-01</td>
<td>Reusable, for ET tube sizes &gt; 4.0 mm</td>
</tr>
<tr>
<td>Reusable Neonatal/Pediatric Airway Adapter</td>
<td>8000-0263-01</td>
<td>Reusable, for ET tube sizes ≤ 4.0 mm</td>
</tr>
<tr>
<td>SPU* Pediatric/Adult Airway Adapter with Mouthpiece</td>
<td>8000-0265-01</td>
<td>Single patient use, for non-intubated patients</td>
</tr>
<tr>
<td>CAPNO₂ mask, Large Adult</td>
<td>8000-0761</td>
<td>SPU, for non-intubated large adults</td>
</tr>
<tr>
<td>CAPNO₂ mask, Standard Adult</td>
<td>8000-0760</td>
<td>SPU, for non-intubated adults</td>
</tr>
<tr>
<td>CAPNO₂ mask, Pediatric</td>
<td>8000-0762</td>
<td>SPU, for non-intubated adults pediatric patients</td>
</tr>
</tbody>
</table>

* SPU = Single Patient Use
### Table 1-2. LoFlo Sidestream CO₂ Accessories

<table>
<thead>
<tr>
<th>Accessory</th>
<th>REF</th>
<th>Application</th>
</tr>
</thead>
<tbody>
<tr>
<td>LoFlo Module and Cable</td>
<td>8000-0367</td>
<td></td>
</tr>
<tr>
<td>Nasal CO₂ Sampling Cannula, Adult</td>
<td>8000-0351</td>
<td>SPU, Nasal CO₂ sampling only (adult)</td>
</tr>
<tr>
<td>Nasal CO₂ Sampling Cannula, Pediatric</td>
<td>8000-0352</td>
<td>SPU, Nasal CO₂ sampling only (pediatric)</td>
</tr>
<tr>
<td>Nasal CO₂ Sampling Cannula, Infant</td>
<td>8000-0353</td>
<td>SPU, Nasal CO₂ sampling only (neonate)</td>
</tr>
<tr>
<td>Oral/Nasal CO₂ Sampling Cannula, Adult</td>
<td>8000-0354</td>
<td>SPU, Oral/Nasal CO₂ sampling only (adult)</td>
</tr>
<tr>
<td>Oral/Nasal CO₂ Sampling Cannula, Pediatric</td>
<td>8000-0355</td>
<td>SPU, Oral/Nasal CO₂ sampling only (pediatric)</td>
</tr>
<tr>
<td>Nasal CO₂ Sampling with O₂ Delivery Cannula, Adult</td>
<td>8000-0356</td>
<td>SPU, Nasal CO₂ sampling with O₂ delivery (adult)</td>
</tr>
<tr>
<td>Nasal CO₂ Sampling with O₂ Delivery Cannula, Pediatric</td>
<td>8000-0357</td>
<td>SPU, Nasal CO₂ sampling with O₂ delivery (pediatric)</td>
</tr>
<tr>
<td>Oral/Nasal CO₂ Sampling with O₂ Delivery Cannula, Adult</td>
<td>8000-0358</td>
<td>SPU, Oral/Nasal CO₂ sampling with O₂ delivery (adult)</td>
</tr>
<tr>
<td>Oral/Nasal CO₂ Sampling with O₂ Delivery Cannula, Pediatric</td>
<td>8000-0359</td>
<td>SPU, Oral/Nasal CO₂ sampling with O₂ delivery (pediatric)</td>
</tr>
<tr>
<td>Adult/Pediatric Airway Adapter Kit</td>
<td>8000-0362</td>
<td>SPU, for ET tube sizes &gt; 4.0 mm</td>
</tr>
<tr>
<td>Adult/Pediatric Airway Adapter Kit with Nafion tubing</td>
<td>8000-0363</td>
<td>SPU, for ET tube sizes &gt; 4.0 mm</td>
</tr>
<tr>
<td>Pediatric/Infant Airway Adapter Kit</td>
<td>8000-0361</td>
<td>SPU, for ET tube sizes ≤ 4.0 mm</td>
</tr>
<tr>
<td>Pediatric/Infant Airway Adapter Kit with Nafion tubing</td>
<td>8000-0364</td>
<td>SPU, for ET tube sizes ≤ 4.0 mm</td>
</tr>
</tbody>
</table>

* SPU = Single Patient Use

**Note:** Components of this product and its associated EtCO₂ accessories that make patient contact are free of latex.

**Note:** The CAPNOSTAT 5 and its accessories are covered by the following US patents:

- 5,146,092
- 5,153,436
- 5,369,277
- 5,616,923
- 5,793,044
- D490,332
- D501,802
- 5,693,944
- 6,935,338
- 6,954,702
- 7,004,168
- 7,059,322
- 7,121,134
- 7,291,851
- 7,294,839

Other patents pending.
The following three tables list the messages that may appear on the R Series unit, possible causes, and the action(s) to take if the message indicates a problem. You should become thoroughly familiar with this information before monitoring patients.

### COMMON MESSAGES

<table>
<thead>
<tr>
<th>Message/Symptom</th>
<th>Possible Cause(s)</th>
<th>Recommended Action(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>--- (dashed lines in EtCO₂ field)</td>
<td>After a defibrillation discharge, the numeric value displays as &quot;---&quot; for up to 10 seconds. When the respiration rate is zero, the numeric CO₂ value will display &quot;---&quot;. When the respiration rate is greater than zero, the actual CO₂ numeric value will be displayed. The EtCO₂ module is warming up (up to 1 minute). Cable not plugged in. Cable defective.</td>
<td>None, normal operation. Check/replace sensor or module.</td>
</tr>
<tr>
<td>--- (dashed lines at top of capnograph waveform)</td>
<td>The scale value setting is incorrect. CO₂ higher than scale limits was detected.</td>
<td>Adjust to higher scale setting using the Zoom softkey.</td>
</tr>
<tr>
<td>CAL BARO. PRESSURE</td>
<td>Barometric pressure reading is out of range.</td>
<td>Calibrate the barometric pressure as described in the R Series Service Manual.</td>
</tr>
<tr>
<td>CO₂ COMM ERROR</td>
<td>There is no communication from the EtCO₂ module or sensor.</td>
<td>Turn R Series unit off, then on again to reset. Try another EtCO₂ sensor. If the problem persists, return sensor and/or unit for service.</td>
</tr>
<tr>
<td>CO₂ DEVICE NOT READY</td>
<td>The zero operation cannot be initiated because: • The sensor or module is still warming up. • No sensor or module is attached to the unit. • The sample cell is not plugged into the module. • Zeroing was attempted within 20 seconds of a detected breath.</td>
<td>Wait for sensor or module to warm up. Attach sensor or module to the unit. Plug sample cell into sample cell receptacle on module.</td>
</tr>
<tr>
<td>CO₂ OUT OF RANGE (dashed lines for CO₂)</td>
<td>The calculated CO₂ value is greater than 150 mmHg.</td>
<td>If error persists, perform a mainstream airway adapter or module zero, as described in &quot;Zeroing the Mainstream CAPNOSTAT 5 CO₂ Sensor/Airway Adapter&quot; on page 7 and &quot;Zeroing the LoFlo CO₂ Module/Sample Cell&quot; on page 11.</td>
</tr>
<tr>
<td>CO₂ UNIT ERROR</td>
<td>The EtCO₂ sensor or module has detected a hardware error.</td>
<td>Check that the sensor is properly plugged in. Reinsert the sensor. Turn R Series unit off, then on again to reset. Perform a mainstream airway adapter or module zero, as described in &quot;Zeroing the Mainstream CAPNOSTAT 5 CO₂ Sensor/Airway Adapter&quot; on page 7 and &quot;Zeroing the LoFlo CO₂ Module/Sample Cell&quot; on page 11. If the problem persists, contact Technical Support.</td>
</tr>
</tbody>
</table>
### COMMON MESSAGES (continued)

<table>
<thead>
<tr>
<th>Message/Symptom</th>
<th>Possible Cause(s)</th>
<th>Recommended Action(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>CO2 WARM UP</strong></td>
<td>The mainstream sensor or LoFlo module is warming up. This may take up to 5 minutes.</td>
<td>Wait for sensor or module to warm up. If the message persists more than 5 minutes, replace the sensor.</td>
</tr>
<tr>
<td><strong>ZERO DONE</strong></td>
<td>The sensor/adapter zero or the LoFlo module zero is finished.</td>
<td>No action required.</td>
</tr>
<tr>
<td><strong>ZERO FAILED</strong></td>
<td>The zero operation did not complete successfully.</td>
<td>Clear the occlusion, remove any source of CO₂, and try zeroing again. If problem persists, contact Technical Support.</td>
</tr>
</tbody>
</table>

### MAINSTREAM MESSAGES

<table>
<thead>
<tr>
<th>Message/Symptom</th>
<th>Possible Cause(s)</th>
<th>Recommended Action(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>CHECK CO2 ADAPTER</strong></td>
<td>This is usually caused when the airway adapter is removed from the CAPNOSTAT 5 CO₂ sensor, or when there is an optical blockage on the windows of the airway adapter. It may also be caused by not having performed an adapter zero after changing the adapter type (single patient use vs. reusable).</td>
<td>Clean the airway adapter and reattach it. If the problem persists or the adapter type was changed, perform a mainstream airway adapter zero as described in “Zeroing the Mainstream CAPNOSTAT 5 CO₂ Sensor/Airway Adapter” on page 7.</td>
</tr>
<tr>
<td><strong>CHECK CO2 SENSOR</strong></td>
<td>The CAPNOSTAT 5 CO₂ sensor cable is not properly plugged in or is over temperature.</td>
<td>Check that the sensor cable is plugged in and properly seated in the connector. Check that the sensor is not exposed to excessive heat. If the problem persists, replace the sensor.</td>
</tr>
<tr>
<td><strong>CO2 DEVICE NOT READY</strong></td>
<td>There is CO₂ in the airway adapter when attempting to zero. Zeroing was attempted within 20 seconds of previous zero operation.</td>
<td>Remove airway adapter from CO₂ source including the patient's, and your own exhaled breaths, and ventilator exhaust valves. Wait up to 20 seconds before retrying a mainstream airway adapter zero, as described in “Zeroing the Mainstream CAPNOSTAT 5 CO2 Sensor/Airway Adapter” on page 7.</td>
</tr>
<tr>
<td><strong>ZERO CO2 ADAPTER</strong></td>
<td>Negative CO₂ detected. May be caused by a sensor that was zeroed with CO₂ in the airway, or by an optical blockage of the airway adapter.</td>
<td>Check the airway adapter and clean if necessary. Perform a mainstream airway adapter zero as described in “Zeroing the Mainstream CAPNOSTAT 5 CO2 Sensor/Airway Adapter” on page 7.</td>
</tr>
<tr>
<td><strong>ZEROING CO2 ADAPTER</strong></td>
<td>Adapter zeroing is in progress.</td>
<td>Wait for the adapter zeroing to finish.</td>
</tr>
</tbody>
</table>
## SIDESTREAM MESSAGES

<table>
<thead>
<tr>
<th>Message/Symptom</th>
<th>Possible Cause(s)</th>
<th>Recommended Action(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>CHECK CO2 LINE</strong></td>
<td>Sample line blockage or damage. Sample line is kinked or pinched. Exhaust tube is blocked.</td>
<td>Verify that the sample line is plugged into the module and seated properly. Verify that neither the sample line nor the exhaust tube are blocked, kinked, or pinched. Verify that the airway adapter is not blocked. If the sample line, exhaust tube, or the airway adapter is blocked or otherwise compromised for 15 seconds, this message will appear. The pump will shut off after 2 minutes if the condition that caused the message is not cleared. To restart the pump, correct the blockage, then remove and reinset the sample cell into the sample cell receptacle. If the problem persists, replace the sample line.</td>
</tr>
<tr>
<td><strong>CHECK CO2 MODULE</strong></td>
<td>Module not properly plugged in. Module over temperature.</td>
<td>Check that module cable is plugged in and seated properly in the connector. Check that module is not exposed to excessive heat. If problem persists, replace module.</td>
</tr>
<tr>
<td><strong>CO2 IN LINE: WAIT</strong></td>
<td>CO2 in cannula/adapter when attempting to zero. Sample line disconnected while zero in progress.</td>
<td>Wait up to 20 seconds before retrying module zero. Remove adapter or cannula tip from CO2 source including the patient’s - and your own - exhaled breaths, and ventilator exhaust valves.</td>
</tr>
<tr>
<td><strong>ZERO CO2 MODULE</strong></td>
<td>Negative CO2 detected. May be caused by a module that was zeroed with CO2 in the sample line.</td>
<td>Perform module zero as described in “Zeroing the LoFlo CO2 Module/Sample Cell” on page 11.</td>
</tr>
<tr>
<td><strong>ZEROING CO2 MODULE</strong></td>
<td>Module zeroing in progress.</td>
<td>Wait for module zeroing to finish.</td>
</tr>
</tbody>
</table>
## Specifications

This section summarizes the specifications of the R Series End Tidal Carbon Dioxide (EtCO\(_2\)) option.

<table>
<thead>
<tr>
<th>CAPNOSTAT 5® CO(_2) Sensor</th>
<th>LoFlo™</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Transducer Type</strong></td>
<td>Mainstream</td>
</tr>
<tr>
<td><strong>Principle of Operation</strong></td>
<td>Non-Dispersive Infrared (NDIR) single beam optics, dual wavelength, no moving parts.</td>
</tr>
<tr>
<td><strong>Warm Up Time</strong></td>
<td>Full specifications within 2 minutes at an ambient temperature of 25°C. Capnogram in 15 seconds.</td>
</tr>
<tr>
<td><strong>EtCO(_2) Measurement Range</strong></td>
<td>0 - 150 mmHg</td>
</tr>
<tr>
<td><strong>EtCO(_2) Accuracy</strong> (at 760 mmHg, ambient temperature of 25°C)</td>
<td>0 - 40 mmHg, ±2 mmHg</td>
</tr>
<tr>
<td>(At respiration rates &gt; 80 breaths per minute, all ranges are ±12% of actual.)</td>
<td></td>
</tr>
<tr>
<td><strong>EtCO(_2) Resolution</strong></td>
<td>0.1 mmHg 0 - 69 mmHg</td>
</tr>
<tr>
<td><strong>EtCO(_2) Stability</strong></td>
<td>Short-Term Drift: Drift over four hours ≤ 0.8 mmHg. Long-Term Drift: Accuracy specification will be maintained over a 120 hour period after zeroing.</td>
</tr>
<tr>
<td><strong>EtCO(_2) Noise</strong></td>
<td>RMS noise of the sensor ≤ 0.25 mmHg at 7.5% CO(_2).</td>
</tr>
<tr>
<td><strong>EtCO(_2) Rise Time (10-90%)</strong></td>
<td>&lt; 60 ms (Adult/pediatric adapters)</td>
</tr>
<tr>
<td><strong>Respiration Rate Range</strong></td>
<td>2 - 150 breaths per minute</td>
</tr>
<tr>
<td><strong>Respiration Rate Accuracy</strong></td>
<td>±1 breath per minute. Verified using a solenoid test setup to deliver a square wave of known CO(_2) concentration to the device. 5% and 10% CO(_2) concentrations were used and respiration rate was varied over the range of the device. Pass/Fail criteria was comparison of the respiratory rate output from the sensor to the frequency of the square wave. EtCO(_2) measurements at those rates were compared to the CO(_2) readings under static flow conditions.</td>
</tr>
<tr>
<td><strong>Sample Flow Rate</strong></td>
<td>N/A</td>
</tr>
<tr>
<td><strong>Compensations</strong></td>
<td>Barometric pressure 400 - 850 mmHg (automatic). Operator selectable O(_2)/N(_2)O compensation.</td>
</tr>
<tr>
<td><strong>EtCO(_2) Alarm Limits</strong></td>
<td>User selectable, High 5 - 100 mmHg, Low 0 - 95 mmHg, OFF.</td>
</tr>
<tr>
<td><strong>Respiration Rate (RR) Alarm Limits</strong></td>
<td>User selectable, High 5 - 150 respirations per minute, Low 0 - 100 respirations per minute, OFF.</td>
</tr>
<tr>
<td><strong>Halogenated Agents</strong></td>
<td>Specification allows for halogenated anesthetic agents that may be present at normal clinical levels. The presence of Desflurane in the exhaled breath beyond normal values (5-6%) may positively bias carbon dioxide values by up to an additional 2-3 mmHg.</td>
</tr>
<tr>
<td><strong>Airway Adapter Deadspace</strong></td>
<td>Adult &lt; 5 cc</td>
</tr>
<tr>
<td>Specifications</td>
<td>CAPNOSTAT 5® CO₂ Sensor</td>
</tr>
<tr>
<td>------------------------------------</td>
<td>------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Environmental (sensor or module)</td>
<td>Operating Temperature: 0º C to 40º C</td>
</tr>
<tr>
<td></td>
<td>Storage and Shipping Temperature: -40º C to 70º C</td>
</tr>
<tr>
<td></td>
<td>The R Series unit may not perform to specifications when stored at the upper or lower</td>
</tr>
<tr>
<td></td>
<td>extreme limits of storage temperature and immediately put into use.</td>
</tr>
<tr>
<td>Electromagnetic Immunity</td>
<td>AAMI DF-80:2003, IEC 60601-1-2, 10 V/m</td>
</tr>
<tr>
<td>Software Hazards</td>
<td>Minimized by compliance with EN60601-1-4</td>
</tr>
<tr>
<td>Operating Time with EtCO₂ and SpO₂</td>
<td>For a new fully charged battery pack at 20º C (68º F):</td>
</tr>
<tr>
<td>Options</td>
<td>• 90 defibrillator discharges at maximum energy (200 J), or</td>
</tr>
<tr>
<td></td>
<td>• 3.0 hours minimum of continuous ECG monitoring, or</td>
</tr>
<tr>
<td></td>
<td>• 2.5 hours of continuous ECG monitoring/pacing at 60 mA, 70 beats per minute.</td>
</tr>
</tbody>
</table>