Using the LI-830 and LI-850 Gas Analyzers
Using the LI-830 and LI-850 Gas Analyzers
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Notes on Safety

This LI-COR product has been designed to be safe when operated in the manner described in this manual. The safety of this product cannot be assured if the product is used in any other way than is specified in this manual. The product is intended to be used by qualified personnel. Read this entire manual before using the product.

<table>
<thead>
<tr>
<th>Equipment markings:</th>
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<td><strong>WARNING</strong></td>
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<tr>
<td><strong>CAUTION</strong></td>
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<table>
<thead>
<tr>
<th>Manual markings:</th>
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<tr>
<td><strong>Warning</strong></td>
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<tr>
<td><strong>Caution</strong></td>
</tr>
<tr>
<td><strong>Note</strong></td>
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</tbody>
</table>
CE Marking:
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WARNING: This product contains chemicals known to the State of California to cause cancer and birth defects or other reproductive harm.

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WARNING: This equipment generates, uses, and can radiate radio frequency energy and if not installed in accordance with the instruction manual, may cause interference to radio communications. It has been tested and found to comply with the limits for a Class A computing device pursuant to Subpart J of Part 15 of FCC rules, which are designed to provide a reasonable protection against such interference when operated in a commercial environment. Operation of this equipment in a residential area is likely to cause interference in which case the user, at his own expense, will be required to take whatever measures may be required to correct the interference.

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This symbol indicates that the product is to be collected separately from unsorted municipal waste. The following applies to users in European countries: This product is designated for separate collection at an appropriate collection point. Do not dispose of as household waste. For more information, contact your local distributor or the local authorities in charge of waste management.
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Section 1.
Introduction to the instruments

The LI-830 and LI-850 gas analyzers both measure CO₂ in air at concentrations from 0 to 20,000 ppm. The instruments differ in that the LI-850 measures water vapor in air, but the LI-830 does not. As a consequence of the water vapor measurement, the LI-850 is able to measure CO₂ concentrations with greater accuracy than the LI-830. Regardless of this difference, operating both instruments is essentially the same. This document provides basic operating instructions for the LI-830 CO₂ analyzer and LI-850 CO₂/H₂O analyzer.

Online resources

In addition to the contents of your box, you may be interested in the following resources. All are available from www.licor.com/env/support. Select the LI-830 and LI-850 Analyzer.

Application software

The software used to interact with the instrument is available for both Windows® and macOS® operating systems. It is used to configure the instrument, read measurements, and configure data logging to a computer. For Windows 7 and 8, the device drivers must be installed manually before you can connect with the instrument. Software is available from licor.com/env/support/LI-850/software.html.

Integrator's guide

The Integrator's Guide describes how to control the instrument using a terminal program and read data in a command-line interface.
Instruction manual

A .pdf version of this document can be downloaded to your computer.

What's what

The LI-830 and LI-850 include a core set of components and each may include optional components. These are described below.

Gas analyzer

This is the gas analyzer in an enclosure. It may include an optional built-in pump and display.

- **Power** indicator: Illuminates when the instrument is powered on.
- **Low Battery** indicator: Illuminates if the power supply voltage drops below 10.5 volts. The instrument will continue to operate with a low battery, but expect a corresponding decline in performance.
- **Alarm** indicators illuminate if an alarm threshold has been crossed. See *Using alarms* on page 3-9.
- **RS-232 Serial** and **USB** connectors to connect with a computer or serial device.
- **Air Inlet** and **Air Outlets** are shipped with covers in place. Keep the covers so you can put them back over the ports when the analyzer is not in use.
- **Power In**: Compatible with 12 to 30 VDC power supplies.
- The **Terminal Strip** has another power connector and outputs for alarms and digital-to-analog converters (DAC).

### Accessories kit

The instrument includes a standard accessories kit, which has accessories and some replacement parts for your gas analyzer. The accessories kit includes the following components:

<table>
<thead>
<tr>
<th>Description</th>
<th>Quantity</th>
<th>Part Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Universal power supply; Input: 100 to 240 VAC, 50 to 60 Hz; Output: 12 VDC, 2.5 amps</td>
<td>1</td>
<td>591-13032</td>
</tr>
<tr>
<td>Outlet adapter kit for universal power supply</td>
<td>1</td>
<td>591-13033</td>
</tr>
<tr>
<td>USB cable; Standard 1.8 m USB-A to USB-B&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1</td>
<td>392-06652</td>
</tr>
<tr>
<td>RS-232 null modem, cross-over cable kit&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1</td>
<td>9975-016</td>
</tr>
<tr>
<td>14-pin terminal block; Includes label sticker (250-05340)</td>
<td>1</td>
<td>331-05273</td>
</tr>
<tr>
<td>Tube fitting nuts for air inlet and outlet</td>
<td>2</td>
<td>9861-036</td>
</tr>
<tr>
<td>Bev-a-line IV plastic tubing; 3.6 meter roll</td>
<td>1</td>
<td>222-01824</td>
</tr>
<tr>
<td>Air filters</td>
<td>2</td>
<td>9967-008</td>
</tr>
<tr>
<td>Quick-connect straight unions</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cleaning kit</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3&quot; Source and detector cleaning swabs</td>
<td>5</td>
<td>610-05314</td>
</tr>
<tr>
<td>5&quot; Optical path cleaning swabs</td>
<td>5</td>
<td>610-05315</td>
</tr>
<tr>
<td>O-Rings (AS-018 Nitrile 70)</td>
<td>4</td>
<td>192-00226</td>
</tr>
<tr>
<td>Black polyurethane feet</td>
<td>4</td>
<td>234-02268</td>
</tr>
</tbody>
</table>

### Air pump (optional)

An optional air pump is available for the gas analyzer. The pump (if applicable) is installed in the gas analyzer case. If you want to add a pump, you can order one later.

### Display (optional)

An optional display is available. The display shows CO<sub>2</sub> concentration, H<sub>2</sub>O concentration (LI-850 only), bench temperature, and pressure.

<sup>a</sup>The use of cables other than those provided may result in improper electrical performance.
Section 1. Introduction to the instruments
Section 2.
Connecting with the analyzer

The following tutorial describes the basic steps you'll follow when you turn on the LI-830 or LI-850 for the first time.

1 Install the software on your computer

The software is used to configure data communication, analog and digital outputs, alarms, data logging, and to verify the calibration. Software is available from licor.com/env/support/LI-850/software.html.

2 Install an air filter on the air-in port

**Caution:** Always install an air filter on the input air before operating the LI-830 or LI-850. Failure to use a filter will cause contamination of the optical bench.

Retrieve an air filter and tubing from the accessories kit. Cut a 5 to 10 cm piece of tubing. Attach the filter and tubing to the air-in port. Observe the arrow on the filter and be sure that the air flow is in the proper direction.
3 Install the USB cable

The USB connection is the simplest way to connect the analyzer with a computer. If you want to use the RS-232 serial connection instead, connect the cable to the serial port, and install a serial-to-USB adapter on the serial cable, if needed.

4 Power on the gas analyzer

The LI-830 and LI-850 will turn on when power is supplied, so plug in the power cable to turn it on. If you use an alternative power supply, it must be able to source a minimum of 1.2 A at 12 VDC while the instrument warms up. After the warm-up, the instrument will draw about 0.3 A at 12 VDC.

5 Connect with the gas analyzer

Start the software.

Click the Connection button in the upper right of the display. Select your instrument from the list, set the data rate, then click Connect. Data will populate the window after connecting. You can enter the COM port manually by checking Manual Port and setting the port number (COM#).

If you are using Windows® 7 or 8, install the drivers before you connect, as described in Installing device drivers on page 2-5.
The terminal strip

The terminal strip connects to the front panel of the instrument. The pin assignments are given on the top of the instrument, and there is a sticker in the accessories kit that you can use to help identify which terminal is which.

<table>
<thead>
<tr>
<th>Terminal</th>
<th>Label</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>12-30 VDC</td>
<td>Voltage In, 12-30 VDC</td>
</tr>
<tr>
<td>2</td>
<td>GND</td>
<td>Ground</td>
</tr>
<tr>
<td>3</td>
<td>High Alarm</td>
<td>High Alarm</td>
</tr>
<tr>
<td>4</td>
<td>GND</td>
<td>Ground</td>
</tr>
<tr>
<td>5</td>
<td>Low Alarm</td>
<td>Low Alarm</td>
</tr>
<tr>
<td>6</td>
<td>GND</td>
<td>Ground</td>
</tr>
<tr>
<td>7</td>
<td>V OUT 2</td>
<td>Voltage output channel 2</td>
</tr>
<tr>
<td>8</td>
<td>GND</td>
<td>Ground</td>
</tr>
<tr>
<td>9</td>
<td>V OUT 1</td>
<td>Voltage output channel 1</td>
</tr>
<tr>
<td>10</td>
<td>GND</td>
<td>Ground</td>
</tr>
<tr>
<td>11</td>
<td>4-20 mA 2</td>
<td>Current output channel 2</td>
</tr>
<tr>
<td>12</td>
<td>GND</td>
<td>Ground</td>
</tr>
<tr>
<td>13</td>
<td>4-20 mA 1</td>
<td>Current output channel 1</td>
</tr>
<tr>
<td>14</td>
<td>GND</td>
<td>Ground</td>
</tr>
</tbody>
</table>
Section 2. Connecting with the analyzer

Powering the LI-830 and LI-850

A power supply—either a battery or the universal power adapter—can be connected to the LI-830 and LI-850 power jack or to pins 1 and 2 on the terminal strip. The instruments require 12 to 30 VDC. The power supply must be able to provide 14 watts (1.2 A at 12 V) during warmup and about 3.6 watts (0.3 A at 12 V) during normal operation.

Using the universal power adapter

The included power supply is compatible with mains power in most localities globally (input 100 to 240 VAC, 50 to 60 Hz; output 12 VDC, 2.5 amps). You may need to install different blades from the outlet adapter kit to fit your wall outlets.

Using the terminal strip power connectors

The terminal strip has connectors for any power supply that has bare leads. Pin 1 is the positive (+) terminal, and pin 2 is the negative (-) terminal.
Installing device drivers

If you are using Windows 7 and 8, you'll need to install device drivers. After connecting the LI-830 or LI-850 to your computer with the USB cable, follow these steps:

1. Download the drivers and unzip the file.
   Drivers are available at licor.com/env/support/LI-850/software.html or directly at licor.com/documents/pamrykdwsl8c61wj6y815gnf4tukh3t

2. Press the Windows key ( ), type Device Manager, and press Enter to open your device manager.

3. Under Other Devices, select the LI-830/850.

4. Right-click LI-830/850 and select Update Driver Software.

5. Browse and select the folder with the driver files.

6. Click Next to finish installing the drivers.
   When they are installed, you should see LI-COR 8x0 under Ports (COM & LPT).

Storing the gas analyzer

When you are done using the gas analyzer, follow these practices to be sure that it is ready to use the next time you need it. The instrument can be stored in non-condensing conditions. If the instrument is stored in high humidity or moderate humidity for a long time while powered off, some optical components may temporarily become saturated with humidity. If this occurs, the instrument may need to run for several days before readings return to normal. When storing the instrument:

- Install the covers over the air inlet and outlet. This will keep dust, bugs, and other contaminants out of the optical cell.
- Disconnect the power supply and power wires.
- If possible, store the instrument in an air conditioned environment.
Section 2. Connecting with the analyzer

Using the LI-830 and LI-850 Gas Analyzers
Section 3. Configuring the gas analyzer

This section describes the basic operating procedures for the LI-830 and LI-850 gas analyzers. This section depends upon the instrument software, which can be downloaded from the LI-COR technical support web site: www.licor.com/env/support.

Overview of the software

After connecting, the software presents you with live data and graphs. With the LI-830, you'll have CO₂; with the LI-850, you'll have both CO₂ and H₂O.
Configuring graphs

The software will always display a graph on the main page (two with the LI-850), and you can configure the graphs by clicking the **Charting** button or any of the **Chart Settings** buttons beside a particular chart. The options available are the same, regardless of how you get to them.

The graphs display a fixed variable: CO₂ and H₂O with the LI-850. You can configure the settings for each graph and add a second variable. The options are:

**Y-Scaling:** Choose **Autoscale** or **Fixed Scale**. With **Fixed Scale**, you can set the minimum and maximum value for each variable displayed.

**X-Axis:** With **Continuous** checked, the graph will display all of the data from the time you connect to the instrument, rescaling the x-axis when needed. With **Continuous** cleared, the graph can be configured to display the most recent data for a time period.

**Series 2** is an option to plot a second variable on the graph with scaling on the right axis. When checked, you can choose from CO₂ concentration, cell temperature, cell pressure, CO₂ absorption, or input voltage. With the LI-850, the H₂O concentration, H₂O dewpoint, and H₂O absorption will also be available.
Logging data to a PC

The instrument can log data as a text file, which will be stored to the connected computer. Configure the log file under Settings > Logging Options. To log data:

1. Create a file.
   Click Select File, then choose the directory where you want to store the file, and name the file.

2. Configure the log parameters.
   - **File headings**: When checked, the Date and Time and Field Labels for data columns will be included in the data file header.
   - **Field delimiter**: Choose between a space, tab, or semi-colon delimited text file.
   - **Log rate (seconds)**: How often a value is recorded. This can be one measurement per 0.5, 1, 2, 3, 4, 5, 10, or 20 seconds. It is independent of the Filter setting (described in Other options on page 3-8).
   - **Log values**: Select the values you want in the log file.

3. Click Start logging.
   The instrument will create a text file according to the parameters you set. You can open the logged data in a text editor or most spreadsheet programs.
Instrument settings

Under Instrument > Settings, you'll find the DAC (digital-to-analog) Outputs (0 to 2.5 V, which also enables the 4 to 20 mA output, and 0 to 5 V, which disables the 4 to 20 mA output), and Other Options, including the heater controls, pressure compensation control, pump control, and the digital signal filter. The 4 to 20 mA output is enabled only when the 0 to 2.5 V range is selected.

![Instrument Settings](image)

Configuring the DAC outputs

This section describes how to configure the voltage and current outputs. Both voltage and current outputs are configured at the same time, in the same place:

- **DAC 1** configures `V Out 1` (pin 9) and `4-20 mA 1` (pin 13; only when the 0 to 2.5 V range is selected) on the terminal strip
- **DAC 2** configures `V Out 2` (pin 7) and `4-20 mA 2` (pin 11; only when the 0 to 2.5 V range is selected) on the terminal strip

Optimizing the DAC resolution

When configuring the analog outputs, keep in mind that the selected ppm range will affect the resolution of data that is sent over the analog outputs.

Think of it like this: The 16-bit DACs can output a fixed number of values (65,536 to be exact). If a DAC is configured to output 0 to 5000 mV, you will have resolution of 0.076 mV per count.

\[
\frac{5000 \text{ mV}}{65536 \text{ counts}} = 0.076 \text{ mV/count}
\]
With the 0 to 5 volt output range corresponding with a 0 to 20,000 ppm CO₂ concentration range, the finest change in concentration that can be resolved is equal to 0.31 ppm.

\[
\frac{20000 \text{ ppm range}}{65536 \text{ counts}} = 0.31 \text{ ppm/count}
\]

If you are measuring concentrations between 0 and 10,000 ppm, the range is 10,000 ppm. By configuring the DAC range for 0 V = 0 ppm CO₂ and 5 V = 10,000 ppm, the DAC output will have twice the resolution as it would in the previous configuration.

\[
\frac{10000 \text{ ppm range}}{65536 \text{ counts}} = 0.15 \text{ ppm/count}
\]

Figure 3-1. The resolution of the analog outputs depends upon the range. If the DACs are configured to output data over the full 0 to 20,000 ppm range, the data output over the DACs will have lower resolution (left). If the DACs are configured to output data over a narrower range (for example, 0 to 10,000 ppm as shown on the right), the data output over the DACs will have higher resolution.

Also keep in mind that the instrument will not output meaningful information over the DACs if the measured value is outside of the configured output range.

**Computing readings from the DAC output**

Here we give some example computations.

**Example 1: Computing CO₂ from a voltage output; 0 to 500 ppm range**

The CO₂ concentration is calculated from the DAC output. In this example, let's configure the output range for 0 V = 0 ppm and 5 V = 500 ppm. The concentration is computed from:
\[ CO_2 = V_{output} \left( \frac{CO_2_{range}}{V_{range}} \right)\]  

If the output voltage is 2.9 V, then

\[ CO_2 = 2.9 \, V \left( \frac{500 \, ppm}{5 \, V} \right) = 290 \, ppm\]  

Example 2: Computing CO\(_2\) from a voltage output; 300 to 500 ppm range

In this example, let's configure the output range for 0 V = 300 ppm and 5 V = 500 ppm, so the full range is 200 ppm, and notably, 0 V is no longer 0 ppm.

\[ CO_2 = V_{output} \left( \frac{CO_{2}\_max-CO_{2}\_min}{V_{range}} \right) + CO_{2}\_min \]  

\(CO_{2}\_max\) is the reading that corresponds with the high voltage output (500 ppm in this case), and \(CO_{2}\_min\) is the reading that corresponds with the low voltage output (300 ppm in this case). With an output voltage of 2.9, the concentration is:

\[ CO_2 = 2.9 \, V \left( \frac{200 \, ppm}{5 \, V} \right) + 300 \, ppm = 416 \, ppm\]  

Example 3: Computing CO\(_2\) from a current output; 0 to 500 ppm range

Unlike the voltage outputs, which are 0 to 2.5 or 0 to 5 V, the current outputs are from 4 to 20 mA (the 4 to 20 mA output is only available when the 0 to 2.5 V range is selected). Therefore, the range is always 16 (20 - 4), and the offset is always 4. Electrical current is indicated with \(I\). In this example, let's configure the output range for 4 mA = 0 ppm and 20 mA = 500 ppm. The concentration is computed from:

\[ CO_2 = (I_{output} - 4) \times \left( \frac{CO_{2}\_range}{I_{range}} \right) \]  

If the current is 16.25 mA, then

\[ CO_2 = (16.25 \, mA - 4) \times \left( \frac{500 \, ppm}{16 \, mA} \right) = 382.8 \, ppm \]
Example 4: Computing CO₂ from a current output; 300 to 500 ppm range

In this example, let's configure the output range so that 4 mA = 300 ppm and 20 mA = 500 ppm (the 4 to 20 mA output is only available when the 0 to 2.5 V range is selected). Now, we need an offset that corresponds with the minimum CO₂ reading (300 ppm).

\[
CO_2 = (I_{\text{output}} - 4) \times \left( \frac{CO_{2_{\text{max}}}-CO_{2_{\text{min}}}}{16 \text{ mA}} \right) + CO_{2_{\text{min}}}
\]

If the current is 16.25 mA, then

\[
CO_2 = (16.25 \text{ mA} - 4) \times \left( \frac{200 \text{ ppm}}{16 \text{ mA}} \right) + 300 \text{ ppm} = 453 \text{ ppm}
\]

Example 5: Computing cell temperature from a voltage output

Cell temperature (T in °C) is calculated from the DAC output with the following:

\[
T = (X_F - X_Z) \frac{V}{V_{\text{max}}} + X_Z
\]

where \(X_F\) is the full scale temperature value that corresponds to the high voltage, \(X_Z\) is the temperature value that corresponds to the low voltage, \(V\) is the voltage reading, and \(V_{\text{max}}\) is the output range (either 2.5 V or 5 V).

Example 6: Computing cell pressure from a voltage output

Cell pressure (kPa) can be computed from a voltage output with the following:

\[
\text{Pressure} = (X_F - X_Z) \frac{V}{V_{\text{max}}} + X_Z
\]

where \(X_F\) is the full scale pressure value that corresponds to the high voltage, \(X_Z\) is the pressure value that corresponds to the low voltage, \(V\) is the voltage reading, and \(V_{\text{max}}\) is the output range (either 2.5 V or 5 V).
Other options

**Enable heater:** The heater should be enabled to maintain the optical bench temperature. Normally the optical bench will be about 51.4 °C. If you disable the heater, the instrument will not perform as specified.

**Pressure compensation:** Pressure compensation should be applied to ensure that the instrument compensates for changes in pressure. Disabling pressure compensation will lead to less accurate measurements.

**Enable pump:** If your instrument is equipped with a pump, this box will be interactive. Check it to turn the pump on. Clear it to turn the pump off.

**Filter:** The digital filter instructs the instrument to average the readings. A value of 0 means no averaging, whereas a value of 20 instructs the instrument to average the previous 20 seconds of measurements. Increasing the averaging time will decrease the variation in each reported value, but will also reduce the instrument response time.

Using the Pump

For instruments that are equipped with the LI-COR-installed pump, the pump is activated through the interface.

**Note:** The pump will not run until the optical cell has reached the operating temperature of about 50 °C, regardless of the setting. If the cell temperature drops below 50 °C, the instrument will disable the pump temporarily until the cell warms back up.

To turn on the pump from the main window, click the pump button so it reads **Pump: On.** Or, click **Settings > Instrument** and check **Enable pump.**
Using alarms

The alarms can be set for CO₂ or H₂O (in the LI-850). Each alarm features an activation level and a deadband. The high alarm is activated if the measured concentration exceeds a high threshold. The high deadband is the range below the threshold that the alarm remains activated. The low alarm is activated if the measured concentration drops below a low threshold. The low deadband is the range above the low threshold that the alarm remains activated.

![Diagram showing high and low alarm thresholds and deadbands for CO₂ concentration.](image)

**Figure 3-2.** An alarm is triggered when a concentration is outside of a threshold. For example, the high alarm is activated if the CO₂ concentration exceeds 700 ppm. The alarm is deactivated when the concentration drops below 600 ppm (high deadband = 100). The low alarm is activated if the concentration drops below 300 ppm and is deactivated when the concentration rises above 350 ppm (low deadband = 50).

![Alarm Settings window](image)

**Figure 3-3.** The Alarm Settings window is where you configure alarms. The settings shown here will give the behavior shown in Figure 3-2 above.
In **TTL mode** (transistor-to-transistor logic), the instrument sends a +5 V signal when the alarm is off and a 0 V signal when it is on. The alarm can be used to activate a relay switch or send a digital signal, for example. In **Open collector mode**, the alarm works like a switch. There is infinite resistance when the alarm is off and low resistance when the alarm is on, allowing current to flow between the pins.

Alarms are configured under **Settings > Alarm Thresholds**. To configure the alarms:

1. **Check Enable Alarms.**
   Select whether the alarm should be for CO₂ or H₂O (with the LI-850). Alarms can be set for one or the other, not both.

2. **Configure the High Alarms.**
   - The **High threshold** is the high level at which the alarm is activated.
   - The **High deadband** is the range below the high threshold that the alarm remains activated.

3. **Configure the Low Alarms.**
   - The **Low threshold** is the low level at which the alarm is activated.
   - The **Low deadband** the range above the low threshold that the alarm remains activated.

4. **Click Apply** when you are done.

---

**Figure 3-4. In this example, the high alarm is used to activate a relay, which turns on a fan.**
Section 4.
Troubleshooting

In this section, we describe how to identify some potential problems. If you can't find a solution here, contact your local distributor or LI-COR technical support for more help.

Instrument will not power on

Power supply adequate? The power supply should source at least 1.2 amps at 12 VDC (minimum of 14 watts during warmup).

Blown fuse? The instrument can be powered from the power jack or pins one and two on the terminal strip. Each connection has its own fuse.

- If you are unable to power on the instrument using the jack, but can power it on using the terminal strip, you probably have a blown power jack fuse.
- If the opposite is true, you probably have a blown terminal strip fuse.
- If you can't power it using either connection, you may have blown both fuses.

Check the resistance across the fuse contacts using an ohm meter. A reading of 0 means that the fuse is good; a reading of 1 indicates that the fuse is blown.

See Replacing a fuse on page 5-6 for complete instructions.

Unable to span or zero the instrument

Dirt in the optics? If the optical cell becomes contaminated, the instrument will drift in either the zero or span.

See Cleaning the optical bench on page 5-4 for complete instructions.
Instrument reports -50 ppm CO₂ or measurements jump around

If the instrument measures -50 ppm or the measurements are going between negative and positive values, or just simply not making any sense, the optical source may have failed or be in the midst of failure. Contact technical support for additional troubleshooting help.
Section 5. Maintenance

The LI-830 and LI-850 will require little maintenance. Typical maintenance procedures are described in this section.

User calibration

If the instrument is not measuring as expected, or if you have disassembled the optical bench for any reason, you should check the zero and span settings and set them if necessary. The zero and span are an offset and slope. The zero value ensures that the instrument shows zero when the gas has a zero concentration. A change in the zero will affect every measurement. The span setting ensures a correct measurement at a known non-zero concentration. A change in the span affects higher concentration measurements more than lower ones.

Setting the CO₂ zero

Always perform the zero first. To set the zero, you'll need either a tank of dry air that is free of CO₂ or a CO₂ scrubbing chemical such as wet soda lime and a desiccant such as Drierite.

1. Plumb the zero-gas tank or scrubber to the air inlet.
   Be sure to use an air filter to prevent contaminants from entering the optical path.
   • If using tank air, the pressure of the tank is sufficient to flow the gas through the analyzer. Allow at least 0.75 liters per minute to flow through the cell (no more than 1.0 lpm).
   • If using a scrubbing chemical, use a pump to draw air through the analyzer.

2. Install a 10 to 20 cm length of tubing to the air outlet.
   This vent prevents ambient air from diffusing upstream into the optical cell.
3 When the CO₂ concentration has stabilized, click the Zero CO₂ button.

Setting the primary CO₂ span

When choosing a span gas, we recommend a gas concentration that is close to the upper limit of what you expect to measure. For example, if you are measuring near-ambient levels, choose a span gas that is near 400 ppm CO₂ (as opposed to 18,000 ppm). Similarly, if you are measuring concentrations near 15,000 ppm CO₂, a span gas with 100 ppm would not be ideal.

4 After zeroing, flow a gas with a known CO₂ concentration through the analyzer at a rate of 0.5 liters per minute.

5 Enter the CO₂ concentration of the span gas into the software.

6 When the CO₂ reading has stabilized, click Span CO₂.
Setting the secondary CO₂ span

You can set a second span (using a gas that has a CO₂ concentration that is higher or lower than the primary span gas) to improve the precision of the analyzer. The process is exactly the same as setting the primary span, only you'll enter a different concentration and click Span2 CO₂.

Setting the H₂O zero and spans (LI-850 only)

The water vapor span can be set with a dew point generator such as the LI-610. The procedure is the same as setting the CO₂ zero and spans, only this uses known concentrations of water vapor rather than CO₂.

Caution: Setting the zero and span incorrectly for either CO₂ or H₂O will adversely impact the performance of your instrument. If you do not have the proper equipment to span the analyzer, it is best to leave it alone.

Recovering from a bad zero or span

If your attempt to zero or span does not go as planned, you can restore the factory default zero and span settings. The information you need is provided on the calibration sheet (included with the instrument or available for download from www.licor.com/env/support/). Under Settings > Calibrations > Advanced, enter the factory zero and span values for your instrument.
Cleaning the optical bench

The optical bench can be removed and cleaned if necessary. If, for example, you are unable to set the span, the optics may be contaminated. Generally speaking, don't undertake this procedure unless you've ruled out other potential problems. You'll have to set the instrument zero and span after reassembling the optical bench.

1. Turn off the instrument.
   Unplug the power cable.

2. Remove the top cover.
   It is attached with 6 captive screws. Loosen each of the screws until the top cover is free of the bottom. If your instrument has a display, carefully rotate the top cover out of the way without straining the cable, and then unplug the display cable. Set the cover aside, being careful not to strain the ribbon cable.

3. Remove the cables from the source and detector.
   Gently grasp the plug and pull it free of the assembly. Leave the tubes in place.

4. Carefully lift the optical bench out of the foam.

5. Remove the screws that secure the source and detector (4 each), then separate the source and detector housings (with circuit boards attached) from the optical path.
6 Clean the parts.

Retrieve an optical path swab from the accessories kit. Dip one end into a 50:50 ethanol-water mixture (mild dish washing soap and water will work too) and carefully swab both ends of the optical path. Dip a Source/Detector swab into the solution and then swab around the source and detector to remove any residue.

**Caution:** Do not use abrasive cleansers. Abrasive cleaners can irreparably damage the gold plating on the optical path, source, or detector.

7 Inspect the hose barbs and tubing.

If the tubes are dirty or damaged, replace them with new tubes (available from LI-COR, part number 6580-041). Carefully remove them from the hose barbs. If the tubes are in good condition and clean, you may be able to reuse them. If the hose barbs are dirty, remove them and clean them with rubbing alcohol or soapy water. Use caution: Do not scratch the hose barbs because scratches may cause leaks.

8 Inspect the O-rings.

If they are smashed flat or damaged in any way, replace them with new O-rings (part number 192-00226) from the accessories kit.

9 Let the optical bench components dry.

10 Reassemble the optical bench.

Attach the source and detector. The orientation of the optical path cylinder is unimportant — either end can be inserted into the source and detector housing. Tighten each of the screws snugly.

11 Place the optical bench in the foam, plug in the source and detector connectors, and re-assemble the case.

Be sure the foam insulation on the top cover is positioned over the optical bench. It is required for thermal stability.

12 Perform a zero and span calibration. See *User calibration* on page 5-1.
Replacing a fuse

The power supply and terminal strip are both protected by fuses. The instrument has one extra fuse (Littelfuse 476 Series Nano® 3 amp 125 V fuse) that can be used in the event that one of the fuses blows. If you are able to power on the instrument using one of these two ways—but not the other—a fuse may be blown. If you can't power the instrument using either way, both fuses may be blown (or you may have a power supply issue).

Before simply replacing a fuse, be sure to identify the problem that caused the fuse to blow in the first place. Otherwise, you'll just blow the spare fuse, in which case, you'll still have the problem and you'll be out of spare fuses. After identifying and solving the problem that caused the fuse to blow, replace the fuse:

1. Open the case.
   Loosen each of the six top cover screws until the top cover is free of the bottom. If your instrument has a display, carefully rotate the top cover out of the way without straining the cable, and then unplug the display cable. Set the cover aside.

2. Locate the fuses.
   With the air-in and -out ports facing away from you, the fuses are mounted to the lower left corner of the lowest circuit board. In the image, ribbon cables and tubes have been removed for clarity.

3. Using a needle-nose pliers, grasp the blown fuse and remove it from the holder.

4. Replace it with a spare fuse.

5. Power on the instrument to verify that the issue is resolved.
Appendix A.
Equation Summary

The LI-830 and LI-850 compute CO₂ concentrations using an equation of the form

\[ c = f(\alpha'')(T + 273.15) \]  

A-1

where \( c \) is concentration, \( f() \) is the calibration function, \( \alpha'' \) is the absorptance, \( g(\alpha,P) \) is the pressure correction, \( S(\alpha) \) is the span, and \( T \) is the temperature (°C) of the gas in the cell, typically 51.5 °C. Absorptance is computed from

\[ \alpha'' = \alpha' g(\alpha',P) \]  

A-2

\( \alpha' \) is a span corrected absorptance, and \( g(\alpha',P) \) is the pressure correction.

\[ \alpha' = \alpha S(\alpha) \]  

A-3

\( S(\alpha) \) is the span function, and raw absorptance \( \alpha \) is computed from

\[ \alpha = \left( 1 - \frac{V}{V_o} Z \right) \]  

A-4

where \( V \) and \( V_o \) are the raw detector sample and reference readings, and \( Z \) is the zeroing parameter.

Span is a linear function of absorptance.

\[ S(\alpha) = S_o + S_1 \alpha \]  

A-5
H₂O Equations (LI-850 only)

Absorptance $\alpha_w$ for water vapor is computed from

$$\alpha_w = \left( 1 - \frac{V_w}{V_{wo}} Z_w \right)$$  \hspace{1cm} (A-6)

$$\alpha'_w = \alpha_w S_w(\alpha_w)$$

$$\alpha''_w = \alpha' g_w(\alpha'_w, P)$$

where $V_w$ and $V_{wo}$ are the sample and reference raw detector readings, and $Z_w$ is the zero parameter. The pressure correction for water vapor is an empirical function $g_w(\cdot)$ of absorptance and pressure $P$:

$$g_w(\alpha'_w, P) = \frac{P_o}{P\left(1+0.8\alpha'_w\left(\frac{P_o}{P}-1\right)\right)}$$  \hspace{1cm} (A-7)

The value of $P_o$ is 99 kPa. When the pressure correction is not enabled, $g_w(\cdot)$ is simply 1.0. Water vapor concentration $W$ (mmol mol⁻¹) is computed from

$$W = f_w(\alpha''_w)(T+273.15)$$  \hspace{1cm} (A-8)

where $f_w(x)$ is a third order polynomial whose coefficients are given on the calibration sheet.

$$f_w(x) = a_{w1}x + a_{w2}x^2 + a_{w3}x^3$$  \hspace{1cm} (A-9)

CO₂ Equations

The measurement of CO₂ is a bit more complicated than for H₂O because of the influence of water vapor. There is a slight direct cross sensitivity in the CO₂ signal to H₂O. This is measured at the factory and accounted for in the computation of absorptance (equation A-10). There is also a band broadening effect that is accounted for in the computation of concentration (equation A-14).

CO₂ absorptance $\alpha_c$ is computed from

$$\alpha_c = \left( 1 - \left( \frac{V_c}{V_{co}} + X_{wc} \left( 1 - \frac{V_c}{V_{wo}} Z_w \right) \right) \right) Z_c$$  \hspace{1cm} (A-10)
\[ \alpha'_{c} = \alpha_{c} S_{c} (\alpha_{c}) \]

\[ \alpha''_{c} = \alpha'_{c} g_{c} (\alpha'_{c}, P) \]

where \( V_{c} \) and \( V_{co} \) are the raw detector signals for sample and reference, \( Z_{c} \) is the CO\(_{2}\) zero parameter, and \( X_{wc} \) is a cross sensitivity parameter for the effect of water vapor on CO\(_{2}\). Its value is reported on the calibration sheet as \( XS = \).

The empirical pressure correction function \( g_{c}(\cdot) \) depends on CO\(_{2}\) absorptance and pressure:

When \( P = P_{o} \), \( g_{c}(\cdot) = 1 \).

When \( P < P_{o} \)

\[ g_{c} (\alpha_{c}, P) = X \]

\[ X = \frac{1}{A+B\left(\frac{1}{z-\alpha_{c}} - \frac{1}{z}\right)} + 1 \]

\[ A = \frac{1}{a(p-1)} \quad \text{A-11} \]

\[ B = \frac{1}{b+cp+d} \]

\[ p = \frac{P_{o}}{P} \]

where \( a = 1.10158 \), \( b = -6.1217E-3 \), \( c = -0.266278 \), \( d = 3.69895 \), and \( z \) is the asymptotic value of absorptance, obtained from the calibration coefficients (equation A-15).

\[ z = a_{c1} + a_{c3} \quad \text{A-12} \]

When \( P > P_{o} \)

\[ g_{c} (\alpha_{c}, P) = \frac{1}{X} \]

\[ P = \frac{P}{P_{o}} \quad \text{A-13} \]

where \( X, A, \) and \( B \) are computed as in equation A-11. CO\(_{2}\) concentration \( C \) (\( \mu \text{mol mol}^{-1} \)) is computed from
\[ C = f_c \left( \frac{\alpha' c}{\psi(W)} \right) \psi (W) (T + 273.15) \quad A-14 \]

where \( f_c(x) \) is a function whose inverse is a double rectangular hyperbola, and whose coefficients \( (a_1...a_4) \) are given on the calibration sheet.

\[ f_c^{-1} (C) = \frac{a_{c1} C}{a_{c2} + C} + \frac{a_{c3} C}{a_{c4} + C} \quad A-15 \]

Solving equation A-15 for \( C \) yields the calibration function

\[ f_c \left( x \right) = \frac{\left(a_2 a_3 + a_1 a_4\right) - \left(a_2 + a_4\right) x - \sqrt{(a_2 - a_4)^2 x^2 + D x + (a_2 a_3 + a_1 a_4)^2}}{2(x-a_1-a_3)} \quad A-16 \]

Where

\[ D = 2(a_2 - a_4)(a_1 a_4 - a_2 a_3) \quad A-17 \]

\[ \psi (W) \] accounts for band broadening by water vapor.

\[ \psi (W) = 1 + (h(\alpha' c) - 1) \frac{W}{1000} \quad A-18 \]

The band broadening coefficient \( h(\alpha' c) \) has been determined to be 1.45 for the instrument for \( \text{CO}_2 \) concentrations near ambient. At higher concentrations, the value decreases. We capture this behavior with an empirical relationship (equation A-19).

\[ h(\alpha' c) = \frac{1}{(0.64b_w - 0.64)e^{-\frac{1}{b_w}} + \frac{1}{b_w}} \quad A-19 \]

Where \( z \) is from equation A-12, and \( b_w \) is the low concentration band broadening coefficient: 1.45. This is the value shown on the calibration sheet as \( BB = 1.45 \). The typical relationship between \( h(\alpha' c) \) and \( \text{CO}_2 \) concentration is shown in Figure A-1 on the facing page. ('Typical' because the exact relationship depends on the relationship between absorptance and \( \text{CO}_2 \), which is the calibration curve.)
Figure A-1. The typical relationship between \( h(\alpha'c) \) and CO\(_2\) concentration.

Note: We formulated equation A-19 with \( 0.64b_w - 0.64 \) instead of the simple equivalent \( 0.29 \) because this allows band broadening corrections to be turned off by setting \( b_w \) to 1. When \( b_w = 1 \), \( h(\alpha_c) = 1 \) everywhere. Also, to avoid computational problems (underflows, overflows, and division by zero) we constrain the argument \( \alpha_c \) when computing \( h(\alpha_c) \) to be \( 0.1 < \alpha_c \leq z \). \( \alpha_c - 0.1 \) is typically equivalent to about 600 ppm.

Calibration Equations

The following equations describe the implementation of zero and span calibrations.

Zeroing H\(_2\)O (LI-850 only)

When the command for zeroing water is received, the LI-850 computes the water zero from equation A-20, where \( \bar{V}_w \) and \( \bar{V}_{wo} \) are averaged for 5 seconds.

\[
Z_w = \frac{\bar{V}_{wo}}{\bar{V}_w} \tag{A-20}
\]

Zeroing CO\(_2\)

When the command for zeroing CO\(_2\) is received, the instrument computes the CO\(_2\) zero term from equation A-21, where \( \bar{V}_c, \bar{V}_{co}, \bar{V}_w, \) and \( \bar{V}_{wo} \) are averaged for 5 seconds.
\[ Z_c = \frac{1}{\left( \frac{V_c}{V_c} + X_{wc}\left(1 - \frac{V_w}{V_w}Z_w\right) \right)} \]  

**Spanning H\textsubscript{2}O (LI-850 only)**

When the command for setting the span for H\textsubscript{2}O is received, along with the target concentration \( W_T \), from the target concentration, the target absorbance \( \alpha_T \) is computed from

\[ \alpha_{wT} = f_w^{-1}\left( \frac{W_T}{T+273.15} \right) \]  

LI-850 computes \( S_{w0} \) from equation A-23, where \( \overline{\alpha}_w \) is averaged over five seconds.

\[ S_{w0} = \frac{\beta_w}{\overline{\alpha}_w} - S_{w1}\overline{\alpha}_w \]  

where

\[ \beta_w = \frac{\alpha_{wT}}{g_w(\alpha_{wT}, P)} \]  

The instrument retains the following values, which are used for subsequent secondary spans:

\[ \alpha_{w1} = \overline{\alpha}_w \]  

\[ \beta_{w1} = \beta_w \]

**Secondary Span H\textsubscript{2}O (LI-850 only)**

When the secondary span command for H\textsubscript{2}O is received, the instrument computes new values for both \( S_{w0} \) and \( S_{w1} \). First, it measures a new \( \overline{\alpha}_w \) and computes a new \( \beta_w \) from equation A-24. Then, it uses these plus the retained values (\( \alpha_{w1} \) and \( \beta_{w1} \) from the previous normal span) to compute

\[ S_{w1} = \frac{\beta_w}{\overline{\alpha}_w} - \frac{\beta_{w1}}{\overline{\alpha}_w} \]  

Given the new span slope \( S_{w1} \), it updates the span offset \( S_{w0} \) by equation A-23.
Spanning CO₂
When the command for setting the CO₂ span is received, along with the target concentration $C_T$, the instrument computes $S_{c0}$ from equation A-28, where $\bar{\alpha}_c$ and $\bar{W}$ are averaged for 5 seconds.

\[
\alpha_{cT} = f_c^{-1} \left( \frac{C_T}{(T+273.15)\psi(\bar{W})} \right) \tag{A-27}
\]

\[
S_{c0} = \frac{\beta_c}{\bar{\alpha}_c} - S_{c1} \bar{\alpha}_c \tag{A-28}
\]

where

\[
\beta_c = \frac{\alpha_{cT}\psi(\bar{W})}{g_c(\alpha_{cT}, P)} \tag{A-29}
\]

Note that

\[
\psi(\bar{W}) = 1 + (h(\alpha_{cT}) - 1) \frac{\bar{W}}{1000}
\]

\[
= \left( 1 + \left( \frac{1}{(0.64b_w - 0.64)e^{-3\left(\frac{\beta_c}{\alpha_{cT}}\right)^{-1}} + \frac{1}{\beta_c}} \right) \frac{\bar{W}}{1000} \right) \tag{A-30}
\]

We need $\alpha_{cT}$ to compute $\psi(\bar{W})$, but $\alpha_{cT}$ depends on $\psi(\bar{W})$. We resolve this by using an approximation (equation A-31) instead when computing equation A-30

\[
\alpha_{cT} \approx f_c^{-1} \left( \frac{C_T}{(T+273.15)} \right) \tag{A-31}
\]

The instrument retains the following values, which are used for subsequent secondary spans, if necessary:

\[
\alpha_{c1} = \bar{\alpha}_c \tag{A-32}
\]

\[
\beta_{c1} = \beta_c \tag{A-33}
\]

Secondary Span CO₂
When the secondary span command for CO₂ is received, the instrument computes new values for both $S_{c0}$ and $S_{c1}$. First, it measures a new $\bar{\alpha}_c$ and computes a new $\beta_c$...
from equation A-29. Then it uses these, plus the retained values ($\alpha_{c1}$ and $\beta_{c1}$ from the previous normal span) to compute

$$S_{c1} = \frac{\beta_{c} - \beta_{c1}}{\bar{\alpha} - \alpha_{c1}}$$  \hspace{1cm} A-34

Given the new span slope $S_{c1}$, it updates the span offset $S_{c0}$ by equation A-28.
Appendix B.
Specifications

CO₂ measurements
Measurement range: 0 to 20,000 ppm
Accuracy
LI-850: Within 1.5% of reading
LI-830: Within 3% of reading
Calibration drift
Zero drift\(^1\): <0.15 ppm / °C
Span drift\(^2\): <0.03% / °C
Total drift at 370 ppm\(^3\): <0.4 ppm / °C
RMS noise at 370 ppm with 1 sec signal filtering: <1 ppm
Sensitivity to water vapor (LI-850 only): <0.1 ppm CO₂ / mmol mol\(^{-1}\) H₂O
Lower limit of detection: 1.5 ppm

H₂O measurements (LI-850 only)
Measurement range: 0 to 60 mmol mol\(^{-1}\)
Accuracy: Better than 1.5% of reading
Calibration drift
Drift at 0 mmol mol\(^{-1}\): <0.005 mmol mol\(^{-1}\) / °C
Span drift at 10 mmol mol\(^{-1}\): <0.006 mmol mol\(^{-1}\) / °C
Total drift at 10 mmol mol\(^{-1}\): <0.016 mmol mol\(^{-1}\) / °C
RMS noise at 10 mmol mol\(^{-1}\) with 1 sec signal filtering: <0.01 mmol mol\(^{-1}\)
Sensitivity to CO₂: <0.0001 mmol mol\(^{-1}\) H₂O / ppm CO₂
Lower limit of detection: 0.015 mmol mol\(^{-1}\)

\(^1\)Zero drift is the change with temperature at 0 concentration.
\(^2\)Span drift is the residual error after re-zeroing following a temperature change.
\(^3\)Total drift is the change with temperature without re-zeroing or re-spanning.
Appendix B. Specifications

General
Output rate: Up to 2 measurements per second
Response time (T90)
  CO₂: <3.5 seconds from 0 to 375 ppm
  H₂O (LI-850 only): <3.5 seconds from 0 to 21 mmol mol⁻¹
Measurement principle: Non-Dispersive Infrared
Traceability
  CO₂: Traceable gases to WMO standards from 0 to 3,000 ppm; traceable gases to EPA protocol gases from 3,000 to 20,000 ppm
  H₂O (LI-850 only): NIST traceable LI-610 Portable Dew Point Generator
Pressure compensation range: 50 to 110 kPa
Maximum gas flow rate: 1 liter min⁻¹
Output signals: Two analog voltage (0 to 2.5 V or 0 to 5 V) and two current (4 to 20 mA)
Digital outputs: TTL (0 to 5 V) or Open Collector
DAC resolution: 16 bits across user-specified range
Power requirements
  Input voltage: 12-30 VDC
  After warmup (without pump): 0.33A @ 12 VDC (4.0 W) average
  After warmup (with pump): 0.42A @ 12 VDC (5.0 W) average
  During warmup: 1.2 A @ 12 VDC (14 W) maximum
Operating temperature range: -20 to 45 °C
Relative humidity range: 0 to 95% RH, Non-condensing
Dimensions: 22.23 cm W × 15.25 cm D × 7.62 cm H
Weight
  No pump, no display: 1.0 kg
  No pump, with display: 1.02 kg
  With pump, no display: 1.3 kg
  With pump, with display: 1.32 kg
Internal optical cell volume: 14.5 mL

Pump specifications (optional)
Operating temperature range: 5 to 45 °C
Storage temperature range: -20 to 60 °C
Operating humidity range: 0 to 80% RH
Nominal flow rate: 0.75 liters minute⁻¹
Power consumption: 1 W (nominally)
Expected life span: 8,000 hrs in standard conditions with a normal load
Display specifications (optional)

Dimensions: 6.7 cm corner-to-corner
Resolution: 400 × 200 px; monochrome
Power consumption: <200 µW
Displayed variables: CO₂ reading, H₂O reading (LI-850 only), optical bench temperature, and pressure.

Specifications subject to change without prior notice.

Dimensional drawings
Serial cable pin assignments

The image below shows the pin assignments for the 9-pin RS-232 serial connector.

![Serial cable pin assignments diagram]

Serial communication parameters

The LI-830 and LI-850 can communicate through a RS-232 serial port on the front of the instrument. You may need to set the communication parameters on your computer or your terminal emulator program. The RS-232 port is configured as Data Terminal Equipment (DTE) with no hardware handshaking. It is bi-directional, meaning information can be transferred both into and out of the instrument. The port is configured as follows:

- Baud Rate: 9600 bps
- Data Bits: 8
- Parity: None
- Stop Bits: 1
- Flow Control: None
Appendix C.

Warranty

Each LI-COR, Inc. instrument is warranted by LI-COR, Inc. to be free from defects in material and workmanship; however, LI-COR, Inc.'s sole obligation under this warranty shall be to repair or replace any part of the instrument which LI-COR, Inc.'s examination discloses to have been defective in material or workmanship without charge and only under the following conditions, which are:

1. The defects are called to the attention of LI-COR, Inc. in Lincoln, Nebraska, in writing within one year after the shipping date of the instrument.
2. The instrument has not been maintained, repaired or altered by anyone who was not approved by LI-COR, Inc.
3. The instrument was used in the normal, proper and ordinary manner and has not been abused, altered, misused, neglected, involved in an accident or damaged by act of God or other casualty.
4. The purchaser, whether it is a DISTRIBUTOR or direct customer of LI-COR or a DISTRIBUTOR'S customer, packs and ships or delivers the instrument to LI-COR, Inc. at LI-COR Inc.'s factory in Lincoln, Nebraska, U.S.A. within 30 days after LI-COR, Inc. has received written notice of the defect. Unless other arrangements have been made in writing, transportation to LI-COR, Inc. (by air unless otherwise authorized by LI-COR, Inc.) is at customer expense.
5. No-charge repair parts may be sent at LI-COR, Inc.'s sole discretion to the purchaser for installation by purchaser.
6. LI-COR, Inc.'s liability is limited to repair or replace any part of the instrument without charge if LI-COR, Inc.'s examination disclosed that part to have been defective in material or workmanship.

There are no warranties, express or implied, including but not limited to any implied warranty of merchantability of fitness for a particular purpose on underwater cables or on expendables such as batteries, lamps, thermocouples, and calibrations.
Other than the obligation of LI-COR, Inc. expressly set forth herein, LI-COR, Inc. disclaims all warranties of merchantability or fitness for a particular purpose. The foregoing constitutes LI-COR, Inc.'s sole obligation and liability with respect to damages resulting from the use or performance of the instrument and in no event shall LI-COR, Inc. or its representatives be liable for damages beyond the price paid for the instrument, or for direct, incidental or consequential damages.

The laws of some locations may not allow the exclusion or limitation on implied warranties or on incidental or consequential damaged, so the limitations herein may not apply directly. This warranty gives you specific legal rights, and you may already have other rights which vary from state to state. All warranties that apply, whether included by this contract or by law, are limited to the time period of this warranty which is a twelve-month period commencing from the date the instrument is shipped to a user who is a customer or eighteen months from the date of shipment to LI-COR, Inc.'s authorized distributor, whichever is earlier.

This warranty supersedes all warranties for products purchased prior to June 1, 1984, unless this warranty is later superseded. To the extent not superseded by the terms of any extended warranty, the terms and conditions of LI-COR’s Warranty still apply.

DISTRIBUTOR or the DISTRIBUTOR's customers may ship the instruments directly to LI-COR if they are unable to repair the instrument themselves even though the DISTRIBUTOR has been approved for making such repairs and has agreed with the customer to make such repairs as covered by this limited warranty.

Further information concerning this warranty may be obtained by writing or telephoning Warranty manager at LI-COR, Inc.