SERVICE MANUAL

Hi Flow STROLLER®

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<table>
<thead>
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</thead>
<tbody>
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</tr>
</tbody>
</table>

Abbreviations

<table>
<thead>
<tr>
<th>FCV</th>
<th>Flow Control Valve</th>
<th>PRV</th>
<th>Primary Relief Valve</th>
</tr>
</thead>
<tbody>
<tr>
<td>LED</td>
<td>Light Emitting Diode</td>
<td>QDV</td>
<td>Quick Disconnect Valve</td>
</tr>
<tr>
<td>LOX</td>
<td>Liquid Oxygen</td>
<td>RMA</td>
<td>Return Materials Authorization</td>
</tr>
<tr>
<td>LPM</td>
<td>Liters Per Minute</td>
<td>RP</td>
<td>Repair Procedure</td>
</tr>
<tr>
<td>NER</td>
<td>Normal Evaporation Rate</td>
<td>R/R</td>
<td>Removal and Replacement</td>
</tr>
<tr>
<td>POI</td>
<td>Patient Operating Instructions</td>
<td>SRV</td>
<td>Secondary Relief Valve</td>
</tr>
<tr>
<td>N2</td>
<td>Nitrogen Gas</td>
<td>O2</td>
<td>Oxygen Gas</td>
</tr>
<tr>
<td>TF</td>
<td>Top Fill</td>
<td>SF</td>
<td>Side Fill</td>
</tr>
<tr>
<td>DF</td>
<td>Dual Fill</td>
<td>PTFE</td>
<td>Polytetrafluoroethylene (Teflon*)</td>
</tr>
</tbody>
</table>

Definition of Terms

WARNING Description of a condition that can result in personal injury or death.

CAUTION Description of a condition that can result in equipment or component damage.

NOTE A statement containing information important enough to emphasize or repeat.

(ITEM) Item numbers used throughout this manual are shown on the illustrations beginning on page 34.

Disclaimer

This manual is intended for use by experienced personnel only. No attempt should be made to fill or maintain this equipment until both this manual and the Patient Operating Instruction booklet have been read and fully understood.
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CAIRE, Inc.’s Hi Flow Stroller is the portable component of the Liberator/Portable supplementary oxygen system. Enclosed in a durable simulated leather case, Hi Flow Strollers incorporate a stainless steel cryogenic container with the valves, plumbing, and associated hardware required to deliver gaseous oxygen to the patient at near ambient temperature.

The Hi Flow Stroller is comprised of five major assemblies, grouped according to function.

1. **Cryogenic Container** – This assembly is a double walled, vacuum insulated dewar for storing liquid oxygen at approximately -180°C (-300°F).

2. **Breathing Circuit** – This circuit consists of the manifold assembly, fixed orifice rotary flow control valve (FCV) and vaporizing coil. It withdraws liquid oxygen from the cryogenic container, warms it to near ambient temperature, and regulates the flow of oxygen gas to the patient. Water that condenses on the cold coils is collected and absorbed by the condensate pad.

3. **Case Assembly** – The case houses and protects the cryogenic container, the breathing circuit, and the liquid level meter.

4. **Liquid Level Meter** – This system uses a capacitance probe and an electronic (LED) readout to measure and display the LOX level by pressing the onboard operate button.

5. **Filling Circuit** – This manual covers both the side fill Hi Flow with the rotary quick disconnect valve and the top fill Hi Flow with the push quick disconnect valve for filling.
<table>
<thead>
<tr>
<th>Specifications (Nominal Values)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>HI FLOW STROLLER</strong></td>
</tr>
<tr>
<td><strong>Capacity</strong></td>
</tr>
<tr>
<td>LOX lbs. (kg)</td>
</tr>
<tr>
<td>Liquid Liters</td>
</tr>
<tr>
<td>Gaseous Liters</td>
</tr>
<tr>
<td><strong>Hi Flow Selectable Flow Rates</strong></td>
</tr>
<tr>
<td>Off</td>
</tr>
<tr>
<td>2,0</td>
</tr>
<tr>
<td>4,0</td>
</tr>
<tr>
<td>10,0</td>
</tr>
<tr>
<td><strong>Flow Rate Accuracy</strong></td>
</tr>
<tr>
<td><strong>Normal Evaporation Rate lbs./day (kg/day)</strong></td>
</tr>
<tr>
<td><strong>Operating Pressure PSIG (Bar)</strong></td>
</tr>
<tr>
<td>Primary Relief Valve Setting PSIG (Bar)</td>
</tr>
<tr>
<td>Secondary Relief Valve Setting PSIG (Bar)</td>
</tr>
<tr>
<td><strong>Filling Time</strong></td>
</tr>
<tr>
<td>Warm minutes:seconds</td>
</tr>
<tr>
<td>Cold minutes:seconds</td>
</tr>
<tr>
<td><strong>Height</strong> in (cm)</td>
</tr>
<tr>
<td><strong>Width</strong> in (cm)</td>
</tr>
<tr>
<td><strong>Weight</strong></td>
</tr>
<tr>
<td>Empty lbs. (kg)</td>
</tr>
<tr>
<td>Full lbs. (kg)</td>
</tr>
<tr>
<td><strong>Fill Connector Types</strong></td>
</tr>
</tbody>
</table>
Oxygen, as it exists at standard atmospheric pressure and temperature, is a colorless, odorless, and tasteless gas. Oxygen constitutes 21% of the atmosphere, by volume. Aside from its well-documented ability to sustain life, oxygen also supports combustion, even though it is nonflammable. Many substances which will burn in air, burn at a faster rate and at a higher temperature in an oxygen enriched atmosphere. Other materials that do not burn in air will burn as oxygen concentration increases. Additionally, many greases and liquid solvents become extremely hazardous materials when placed in an oxygen-enriched environment. In its liquid form, oxygen is still odorless and tasteless, but is pale blue in color. At an operating pressure of 20 psig (1.4 bar), the temperature of liquid oxygen is about -173°C (-280° F). Skin exposed to such a low temperature can become severely frostbitten.

These hazards require certain safety precautions to be taken when working with or around gaseous and/or liquid oxygen:

1. Never permit combustible substances such as greases, oils, solvents, or other compounds not oxygen compatible to contact any component of the unit exposed to higher-than-atmospheric concentrations of gaseous or liquid oxygen. This especially applies to tubing, fittings, and valves.

2. Keep oxygen equipment away from open flames or electrical appliances such as heaters, stoves, toasters, and other devices with heating elements.

3. Never permit smoking in an area where oxygen equipment is repaired, filled, or used.

4. Always wear goggles, a face shield, and insulated gloves when working with or around liquid oxygen.

While CAIRE, Inc. equipment is designed and built to the most rigid standards, no piece of mechanical equipment can ever be made 100% foolproof. Strict compliance with proper safety practices is necessary when using any Hi Flow unit. We recommend that our distributors emphasize safety and safe handling practices to their employees and customers. While safety features have been designed into the unit and safe operations are anticipated, it is necessary that all distributor personnel carefully read and fully understand WARNINGS, CAUTIONS, and NOTES throughout the manual. Periodic review of this information is recommended.

**WARNING:** Excess accumulation of oxygen creates an oxygen-enriched atmosphere (defined by the Compressed Gas Association as an oxygen concentration above 23%). In an oxygen-enriched atmosphere, flammable items may burn vigorously and may explode. Certain items considered non-combustible in air may burn rapidly in such an environment. Keep all organic materials and other flammable substances away from possible contact with oxygen; particularly oil, grease, kerosene, cloth, wood, paint, tar, coal dust, and dirt which may contain oil or grease. DO NOT permit smoking or open flame in any area where oxygen is stored, handled, or used. Failure to comply with this warning may result in serious personal injury.

**WARNING:** In the event a unit is dropped, tipped over, or unreasonably abused; immediately, but cautiously, raise the container to its normal vertical position. If substantial container damage has occurred, remove the liquid oxygen from the vessel in a safe manner (RP14). Purge the unit with an inert gas (nitrogen) and promptly return it to CAIRE, Inc. for inspection. The container should be prominently marked “CONTAINER DROPPED, INSPECT FOR DAMAGE.” Failure to comply with these procedures may result in personal injury and can seriously damage the container.

**WARNING:** Personnel must remove liquid oxygen and depressurize the unit before removing parts or loosening fittings from a unit. Failure to do so may result in personal injury from the extreme cold of liquid oxygen and/or the pressure in the vessel.

**WARNING:** During transfer of liquid oxygen, components will become extremely cold. Care should be used to avoid any contact with these components, as serious frostbite may result.

**WARNING:** Keep filled unit upright at all times. Tip over of filled unit may result in liquid oxygen leakage and/or an oxygen-enriched atmosphere.

**CAUTION:** Only use replacement equipment which is compatible with liquid oxygen and has been cleaned for oxygen use. Do not use regulators, fittings, hoses, etc. which have been previously used in non-oxygen service.
WARNING: Medical electrical Equipment needs special precautions regarding EMC and needs to be installed and put into service according to the EMC information provided in this manual.

WARNING: Portable and mobile RF communications equipment can affect Medical Electrical Equipment.

WARNING: The use of Accessories, transducers, and cables other than those specified by the manufacturer may result in increased Emissions or decreased immunity of the Hi Flow.

WARNING: The Hi Flow should not be used adjacent to or stacked with other equipment, and that if adjacent or stacked use is necessary, the Hi Flow should be observed to verify normal operation in the configuration in which it will be used.

Table 1

Guidance and Manufacturer’s declaration—electromagnetic emissions

The Hi Flow is intended for use in the electromagnetic environment specified below. The customer or the user of the Hi Flow should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Emissions test</th>
<th>Compliance</th>
<th>Electromagnetic environment—guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF emissions</td>
<td>Group 1</td>
<td>The Hi Flow uses RF energy only for internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.</td>
</tr>
<tr>
<td>CISPR 11</td>
<td></td>
<td></td>
</tr>
<tr>
<td>RF emissions CISPR 11</td>
<td>Class B</td>
<td>The Hi Flow is suitable for use in all establishments, including domestic establishments and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.</td>
</tr>
<tr>
<td>Harmonic emissions</td>
<td></td>
<td></td>
</tr>
<tr>
<td>IEC 61000-3-2</td>
<td>Not applicable</td>
<td></td>
</tr>
<tr>
<td>Voltage fluctuations/</td>
<td></td>
<td></td>
</tr>
<tr>
<td>flicker emissions</td>
<td></td>
<td></td>
</tr>
<tr>
<td>IEC 61000-3-3</td>
<td>Not applicable</td>
<td></td>
</tr>
</tbody>
</table>
### Table 2

**Guidance and manufacturers declaration—electromagnetic immunity**

The Hi Flow is intended for use in the electromagnetic environment specified below. The customer or the user of the Hi Flow should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Immunity test</th>
<th>IEC 60601 test level</th>
<th>Compliance level</th>
<th>Electromagnetic environment—guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrostatic discharge (ESD)</td>
<td>±6 kV contact</td>
<td>±6 kV contact</td>
<td>Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30%.*</td>
</tr>
<tr>
<td>IEC 61000-4-2</td>
<td>±8 kV air</td>
<td>±8 kV air</td>
<td></td>
</tr>
<tr>
<td>Electrical fast transient/burst</td>
<td>±2 kV for power supply lines</td>
<td>Not applicable</td>
<td>Not applicable</td>
</tr>
<tr>
<td>IEC 61000-4</td>
<td>±1 kV for input/output lines</td>
<td>Not applicable</td>
<td></td>
</tr>
<tr>
<td>Surge</td>
<td>±1 kV line(s)</td>
<td>Not Applicable</td>
<td>Not Applicable</td>
</tr>
<tr>
<td>IEC 61000-4-5</td>
<td>±2 kV line(s) to line(s)</td>
<td>DC powered device</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Voltage dips, short interruptions and voltage variations on power supply input lines</td>
<td>&lt;5% UT (&gt;95% dip in UT) for 0,5 cycle</td>
<td>Not Applicable</td>
<td>Not Applicable</td>
</tr>
<tr>
<td>IEC 61000-4-11</td>
<td>40% UT (60% dip in UT) for 5 cycles</td>
<td>No data input/output lines</td>
<td></td>
</tr>
<tr>
<td></td>
<td>70% UT (30% dip in UT) for 25 cycles</td>
<td>DC powered device</td>
<td></td>
</tr>
<tr>
<td></td>
<td>&lt;5% UT (&gt;95% dip in UT) for 5 sec</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Power frequency (50/60 Hz) magnetic field</td>
<td>3 A/m</td>
<td>3 A/m</td>
<td>Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.</td>
</tr>
<tr>
<td>IEC 61000-4-8</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Note: UT is the a.c. mains voltage prior to application of the test level.

* This statement indicates that the required testing was performed in a controlled environment and the Hi Flows are found to be compliant with regulations.
Table 4*

**Guidance and manufacturers declaration—electromagnetic immunity**

The Hi Flow is intended for use in the electromagnetic environment specified below. The customer or the user of the Hi Flow should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Immunity test</th>
<th>IEC 60601 test level</th>
<th>Compliance level</th>
<th>Electromagnetic environment—guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conducted RF</td>
<td>3Vrms</td>
<td>Not Applicable</td>
<td>Portable and mobile RF communications equipment should be used no closer to any part of the Hi Flow, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter.</td>
</tr>
<tr>
<td>IEC 61000-4-6</td>
<td>150kHz to 80 MHz</td>
<td>Battery powered device</td>
<td></td>
</tr>
<tr>
<td>Radiated RF</td>
<td>3 V/m</td>
<td>3 V/m</td>
<td>Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey, should be less than the compliance level in each frequency range.</td>
</tr>
<tr>
<td>IEC 61000-4-3</td>
<td>80 MHz to 2,5 GHz</td>
<td></td>
<td>Interference may occur in the vicinity of equipment marked with the following symbol:</td>
</tr>
</tbody>
</table>

\[
d = 1.2 \sqrt{P}
\]

\[
d = 1.2 \sqrt{P} \quad 80 \text{ MHz to } 800 \text{ MHz}
\]

\[
d = 2.3 \sqrt{P} \quad 800 \text{ MHz to } 2,5 \text{ GHz}
\]

where $P$ is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and $d$ is the recommended separation distance in meters (m).

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

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

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

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

* This table is included as a standard requirement for equipment which has been tested to specific test levels and over specific frequency ranges and been found compliant with regulations.
Table 6*

---

**Safety**

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

<table>
<thead>
<tr>
<th>Rated maximum output power of transmitter (W)</th>
<th>Separation distance according to frequency of transmitter (m)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>150 kHz to 80 MHz</td>
</tr>
<tr>
<td></td>
<td>(d = 1.2 \sqrt{P})</td>
</tr>
<tr>
<td>0.01</td>
<td>0.12 m</td>
</tr>
<tr>
<td>0.1</td>
<td>0.38 m</td>
</tr>
<tr>
<td>1</td>
<td>1.2 m</td>
</tr>
<tr>
<td>10</td>
<td>3.8 m</td>
</tr>
<tr>
<td>100</td>
<td>12 m</td>
</tr>
</tbody>
</table>

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

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

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

* This table is included as a standard requirement for equipment which has been tested to specific test levels and over specific frequency ranges and been found compliant with regulations.
**VI Theory of Operation**

**Filling**

1. **Method**

The Hi Flow Stroller is a portable unit designed to be filled by the patient from a Liberator, Low Loss Reservoir, or other compatible unit.

The unit is filled by coupling the quick disconnect valve of the Hi Flow Stroller with the quick disconnect valve on the Reservoir and opening the vent valve on the portable. Liquid oxygen is forced up the Reservoir fill tube, through the coupled quick connectors, and into the inner vessel of the Hi Flow Stroller.

There will be some oxygen vaporized during filling. This gas is discharged through the vent valve. When the unit is full, liquid oxygen is expelled through the vent valve. Closing the vent valve and separating the units terminates the fill process.

2. **Saturation Pressure**

The liquid oxygen saturation pressure can seriously affect the overall efficiency and operation of the Hi Flow Stroller.

a. If the saturation pressure of the liquid in the fill source is greater than 30 psig (2,1 bar), high filling losses and/or frozen open relief devices may result.

b. If the saturation pressure of the liquid in the fill source is below 18 psig (1,2 bar), below tolerance flow rates may result.

**Oxygen Withdrawl**

With oxygen in the unit, and the vent valve closed, the pressure in the inner vessel will remain at or near the primary relief valve pressure of 19.7 - 20.6 (1,35 - 1,42).

At operating pressure and with the flow control valve at any setting other than Off, pressure forces liquid oxygen up the liquid withdrawl tube and into the breathing coil. In the breathing coil, liquid oxygen absorbs heat and vaporizes, warming to almost ambient temperature by the time gas is dispensed by the flow control valve to the patient.
Liquid Level Measurement

1. LED

Hi Flow Strollers are equipped with a unique liquid level measurement system. This system measures the level of liquid oxygen inside the unit with a capacitance probe and displays that liquid level on the level meter’s LEDs.

The liquid level probe consists of two concentric stainless steel cylinders extending inside the inner vessel. As the liquid oxygen level rises, the capacitance of this assembly goes up. The level meter then displays the liquid level in the cylinder based on a calibration relating capacitance to level. The higher the liquid level in the dewar, the more LEDs are activated, beginning at the leftmost LED.

Electrical connection between the G4 meter and the probe is made via a single conductor JST connector. The male plug is attached to its female counterpart extending from the probe, creating a watertight connection. A single ground wire is connected from the meter to a male spade terminal on the mounting bracket.

The meter is powered by an internal battery, offering an estimated battery life of 5 years or more at 30 actuations per day. The meter has a low battery (LOW BATT) indicator which signals the need for battery replacement. The meter battery is covered under a 2-year limited warranty. If battery failure occurs within 2 years of the Hi Flow Stroller shipment date, contact CAIRE, Inc. customer service for a replacement meter. If the meter battery is no longer under warranty, the CR2032 coin cell battery can be replaced (RP2). The battery can be found at most hardware stores, or they can be ordered through CAIRE, Inc. customer service.

The new level meter improves upon the previous meter by integrating all components within its casing, simplifying removal and replacement (RP8). Even more importantly from a technical service point of view, there is a much improved calibration procedure requiring no additional tools and a range of error reporting codes which can be read directly from the LEDs to report calibration errors. These calibration codes can be found in the calibration procedure (RP5).

FIGURE 4: Liquid Level Meter
Unpacking

1. Inspect the carton for shipping damage. Report any damage to the freight company before signing the bill of lading.
2. Check the description on the carton against your order.
3. Unpack the unit, including the Patient Operating Instructions (POI).
4. Set aside the packing material in case the unit must be returned to the factory.

Setup

If desired, the flow control knob (Item 20) can be adjusted so it will not exceed the maximum prescribed flow rate.

1. To remove the flow control knob, firmly grip and pull straight up away from the portable.
2. Remove the flow rate decal number disc (Item 21).
3. Remove the 2 Phillips head screws (Item 22) from the flow lock plate (Item 26) and remove the plate.
4. Remove the locking pin (Item 23) from its storage position on the flow lock plate and place in the underside of the hole corresponding to the maximum allowable flow rate.
5. Replace the flow lock plate (Item 26) and tighten screws (Item 22). Replace number disk (Item 21) and push-on knob (Item 20). Verify the flow lock is at the correct position.

FIGURE 5: Flow Control Knob

Refer to Patient Operating Instructions (POI) Manual.
There are two schedules for routine maintenance which the home health care distributor may follow. These schedules allow the distributor maximum flexibility while assuring that the equipment is operating properly.

**Schedule A - Biennial**

A. Introduction

Routine maintenance is a series of steps used to assure that equipment is functioning properly.

1. If a unit fails a given test, one of two things may be done:
   a. Refer to Troubleshooting section (Pages 17-21) of this manual.
   b. Return the unit to CAIRE, Inc. for repair.

2. Schedule A – Maximum of two years between routine maintenance testing. Unit should be tested whenever a problem is suspected.

B. Procedure

Follow the steps in order listed. If the unit fails any step, refer to the Troubleshooting section (Pages 17-21) of this manual.

1. Visual Inspection:
   a. Look for damaged or missing parts.
   b. Dry condensation pad. Replace if soiled.

2. Full Unit:
   a. Fill, vent full, of LOX.
   b. Check for audible or visual leaks in QDV and vent valve.
   c. Verify that the meter reads full (8 LED).

3. Check Efficiency of Unit:
   a. Allow the unit to stabilize after filling (10–15 minutes).
   b. Inspect the bottle for cold sweaty condition, and for excessive venting from relief valve (some venting is normal).
   c. If either condition is observed, conduct NER test (see RP28 in this manual).

4. Flow Test (Table 1):
   a. Set FCV to the highest flow setting, run for at least 20 minutes.

### TABLE 1: Flow Test Acceptable Ranges

<table>
<thead>
<tr>
<th>Unit Type</th>
<th>FCV Setting</th>
<th>LPM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Standard</td>
<td>OFF</td>
<td>0</td>
</tr>
<tr>
<td>Standard</td>
<td>0,50</td>
<td>0,40 to 0,60</td>
</tr>
<tr>
<td>Standard</td>
<td>1,00</td>
<td>0,90 to 1,10</td>
</tr>
<tr>
<td>Standard</td>
<td>2,00</td>
<td>1,80 to 2,20</td>
</tr>
<tr>
<td>Standard</td>
<td>2,50</td>
<td>2,20 to 2,25</td>
</tr>
<tr>
<td>Standard</td>
<td>3,00</td>
<td>2,70 to 3,30</td>
</tr>
<tr>
<td>Standard</td>
<td>4,00</td>
<td>3,60 to 4,40</td>
</tr>
<tr>
<td>Standard</td>
<td>6,00</td>
<td>5,40 to 6,60</td>
</tr>
<tr>
<td>Standard</td>
<td>8,00</td>
<td>7,20 to 8,80</td>
</tr>
<tr>
<td>Standard</td>
<td>10,00</td>
<td>9,00 to 11,00</td>
</tr>
<tr>
<td>Standard</td>
<td>12,00</td>
<td>10,8 to 13,20</td>
</tr>
<tr>
<td>Standard</td>
<td>15,00</td>
<td>13,50 to 16,50</td>
</tr>
</tbody>
</table>

b. Check all flow settings to chart above and check pressure to be at least 18 psig (1,2 bar).

5. Prepare for Use:
   a. Empty contents by setting FCV at the highest flow setting and running unit until dry and warm to room temperature (approximately one hour).
   b. Verify that meter reads empty (1 LED) and that the low battery LED is not lit.
   c. Clean case with household glass cleaner and lint free cloth (do not get in any valves).
**Maintenance (Schedule B, Continuous)**

**A. Introduction**

Continuous maintenance is a set of tests and inspections performed periodically to ensure the equipment is functioning properly. It can be done with equipment in service by drivers or other personnel.

1. If a unit fails a given test, it should be taken out of service. Refer to the Troubleshooting section (Pages 17-21) or contact Technical Service.

2. Schedule B – Checks are made when the driver visits patients and when equipment is transferred between patients.

**B. Procedure**

These inspections are to be performed at least once per year by the driver or other personnel when the Hi Flow is in use by the patient (LOX in the unit).

1. **Visual Inspection:**
   - a. Case not damaged or dirty. Flow control knob turns easily with firm detent feel.
   - b. Condensation pad is not soaking wet or excessively dirty.
   - c. QDV pin not bent (side fill only).

2. Verify that the meter reads the correct liquid level.

3. Check flow rate, Erie liter meter (± 0.25 LPM) can be used.

**These inspections/tests are to be done between patients.**

1. **Visual Inspection:**
   - a. Damaged case.
   - b. Condensation pad (replace if soiled).
   - c. QDV pin not bent (side fill only).
   - d. Inspect interior of unit for dirt or contaminants.

2. Verify that liquid level meter reads empty.

3. Fill Unit:
   - a. Verify that liquid level gauge reads full.
   - b. Check for audible/visual leaks in QDV and vent valves.

4. **Flow Test (Table 1):**
   - a. Set flow control knob to highest flow setting, run for at least 20 minutes.
   - b. Check all flow settings with chart below and check pressure to at least 18 psig (1.2 bar).

5. **Prepare for Use:**
   - a. Empty contents by turning flow selection knob to highest flow setting and running unit until dry and warm to room temperature (approximately one hour).
   - b. Verify that the meter reads empty.
   - c. Clean case per RP27.
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**Service Tools/Equipment/Supplies** ...................................................................................................... 33
Introduction

1. These procedures are designed to be performed only by qualified personnel with proper equipment.

2. Any failure during routine maintenance checks will refer you to this section. See troubleshooting chart for appropriate procedure.
## Troubleshooting Chart

<table>
<thead>
<tr>
<th>Symptom</th>
<th>Probable Cause</th>
<th>Corrective Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>1) Unable to start fill or excessively long fill times</td>
<td>a) QDV not properly engaged on the reservoir</td>
<td>Make sure the QDV on the portable and reservoir are properly aligned. Ensure that a downward force is being applied to the portable for top fill. Ensure the portable is completely engaged for side fill.</td>
</tr>
<tr>
<td></td>
<td>b) Reservoir is empty</td>
<td>Swap or re-fill the reservoir</td>
</tr>
<tr>
<td></td>
<td>c) Vent valve not open</td>
<td>Ensure that the vent valve lever is fully in the open position. The lever must be open to begin a fill.</td>
</tr>
<tr>
<td></td>
<td>d) FCV is open</td>
<td>Be sure that the FCV knob is in the off (&quot;0&quot;) position. If the valve is open, fill times can increase.</td>
</tr>
<tr>
<td></td>
<td>e) Reservoir saturation pressure is too low</td>
<td>Swap reservoirs or allow the reservoir time to stabilize and build pressure</td>
</tr>
<tr>
<td></td>
<td>f) Vent valve is obstructed</td>
<td>Inspect the vent tubes for blockages. Clean by blowing out with compressed gas or replace parts if necessary.</td>
</tr>
<tr>
<td></td>
<td>g) Leak in the system</td>
<td>Check the portable for leaks (RP18) and repair if needed.</td>
</tr>
<tr>
<td></td>
<td>h) QDV damaged or faulty</td>
<td>Inspect the QDV and be sure the poppet opens properly and smoothly. If necessary, replace the QDV (RP15 or RP16)</td>
</tr>
<tr>
<td></td>
<td>i) Faulty vent valve</td>
<td>Replace the vent valve (RP19)</td>
</tr>
<tr>
<td>2) Liquid leaks from the coupled QDVs during the fill</td>
<td>a) Worn or damaged lip seal</td>
<td>Replace the QDV lip seal if top fill (RP16). See reservoir technical manual for replacement of side fill lip seal on reservoir.</td>
</tr>
<tr>
<td>3) Unable to disconnect the portable from the reservoir</td>
<td>a) Pop-off assembly not being utilized (Top fill only)</td>
<td>Ensure that the pop-off assembly on the reservoir is being used (top fill only). Do not use force to separate the QDVs on top fill or side fill.</td>
</tr>
<tr>
<td></td>
<td>b) QDVs are frozen together</td>
<td>Leave the units coupled with the vent valve closed and let them sit until they warm up enough to disconnect. Always ensure that male and female QDV’s are cleaned and dried prior to each fill.</td>
</tr>
<tr>
<td>4) Liquid leaks from the QDV poppet after filling</td>
<td>a) Ice crystal preventing the QDV from closing properly.</td>
<td>Engage and disengage the portable onto the reservoir several times to dislodge the ice crystal. Always be sure that the male and female QDV’s are wiped clean and dry before filling.</td>
</tr>
<tr>
<td></td>
<td>b) Dirty or damaged QDV poppet</td>
<td>Replace the QDV (RP15 or RP16)</td>
</tr>
<tr>
<td>5) Liquid leaks from the vent valve tube/outlet</td>
<td>a) Vent valve is not fully closed</td>
<td>Ensure that the vent valve lever is fully in the closed position.</td>
</tr>
<tr>
<td></td>
<td>b) The portable has been transported or laid in an improper operating position</td>
<td>Return the portable to an upright or acceptable operating position and allow several minutes for stabilization.</td>
</tr>
<tr>
<td></td>
<td>c) Vent valve is frozen open</td>
<td>Allow the portable to warm until the vent valve can close. After the warm up, allow up to 60 minutes for the portable to stabilize and build pressure before operating.</td>
</tr>
<tr>
<td></td>
<td>d) Faulty vent valve</td>
<td>Replace the vent valve (RP19)</td>
</tr>
<tr>
<td>Symptom</td>
<td>Probable Cause</td>
<td>Corrective Action</td>
</tr>
<tr>
<td>---------</td>
<td>----------------</td>
<td>-------------------</td>
</tr>
<tr>
<td>6) <strong>Excessive venting from relief valves (hissing sound)</strong></td>
<td>a) The portable has been transported or laid in an improper operating position</td>
<td>Return the portable to an upright or acceptable operating position and allow several minutes for stabilization.</td>
</tr>
<tr>
<td></td>
<td>b) Saturation pressure too high.</td>
<td>Inspect the saturation pressure of the reservoir used for filling. Allow at least 30 minutes at no flow for the portable to saturate properly.</td>
</tr>
<tr>
<td></td>
<td>c) Relief valve frozen open</td>
<td>Allow the portable to warm and thaw. Attempt to re-fill the portable.</td>
</tr>
<tr>
<td></td>
<td>d) Faulty relief valve</td>
<td>Test the operating pressure (RP25) and replace the relief valve if necessary (RP22).</td>
</tr>
<tr>
<td></td>
<td>e) Partial or complete loss of vacuum</td>
<td>Conduct the NER test (RP28) and return the unit to CAIRE, Inc. if necessary.</td>
</tr>
<tr>
<td>7) <strong>No Flow</strong></td>
<td>a) Portable is empty</td>
<td>Check the contents indicator/level gauge and fill the portable if needed.</td>
</tr>
<tr>
<td></td>
<td>b) Flow control valve turned off</td>
<td>Ensure the flow control knob is not in the off (&quot;0&quot;) position.</td>
</tr>
<tr>
<td></td>
<td>c) Nasal cannula kinked or disconnected</td>
<td>Ensure proper nasal cannula functionality and positioning.</td>
</tr>
<tr>
<td></td>
<td>d) Saturation pressure is too low</td>
<td>Inspect the saturation pressure of the reservoir used for filling. Allow at least 30 minutes at no flow for the portable to saturate properly.</td>
</tr>
<tr>
<td></td>
<td>e) Leak in the system</td>
<td>Perform a leak check on the plumbing (RP18). Repair leaks as necessary.</td>
</tr>
<tr>
<td></td>
<td>f) Relief valve is open</td>
<td>Ensure that there is no venting from the relief valves. If there is refer to the corrective actions for “Excessive venting from relief valves (hissing sound)”</td>
</tr>
<tr>
<td></td>
<td>g) Vent valve is open</td>
<td>Ensure that there is no venting from the vent valve outlet/tube. If there is refer to the corrective actions for “Liquid leaks from the vent valve tube/outlet”</td>
</tr>
<tr>
<td></td>
<td>h) Blockage in the liquid withdrawal circuit</td>
<td>Check the warming coils and withdrawal tubes for blockages. Replace if necessary.</td>
</tr>
<tr>
<td></td>
<td>i) FCV Faulty</td>
<td>Replace the FCV (RP20)</td>
</tr>
<tr>
<td>8) <strong>Low flow at all LPM settings</strong></td>
<td>a) Nasal cannula kinked or leaking</td>
<td>Inspect the functionality of the nasal cannula.</td>
</tr>
<tr>
<td></td>
<td>b) Saturation pressure is too low</td>
<td>Inspect the saturation pressure of the reservoir used for filling. Allow at least 30 minutes at no flow for the portable to saturate properly.</td>
</tr>
<tr>
<td></td>
<td>c) Leak in the system</td>
<td>Perform a leak check on the plumbing (RP18). Repair leaks as necessary.</td>
</tr>
<tr>
<td></td>
<td>d) PRV faulty</td>
<td>Test the operating pressure (RP25) and replace the relief valve if necessary (RP22).</td>
</tr>
<tr>
<td></td>
<td>e) Blockage in the liquid withdrawal circuit</td>
<td>Check the warming coils and withdrawal tubes for blockages. Replace if necessary.</td>
</tr>
<tr>
<td></td>
<td>f) FCV faulty</td>
<td>Replace the FCV (RP20)</td>
</tr>
<tr>
<td>Symptom</td>
<td>Probable Cause</td>
<td>Corrective Action</td>
</tr>
<tr>
<td>---------</td>
<td>----------------</td>
<td>------------------</td>
</tr>
<tr>
<td>9) Increased NER</td>
<td>a) Saturation Pressure is too high</td>
<td>Inspect the saturation pressure of the reservoir used for filling. Allow at least 30 minutes at no flow for the portable to saturate properly.</td>
</tr>
<tr>
<td></td>
<td>b) Leak in the system</td>
<td>Preform a leak check on the plumbing (RP18). Repair leaks as necessary.</td>
</tr>
<tr>
<td></td>
<td>c) Relief valve open</td>
<td>Ensure that there is no venting from the relief valves. If there is refer to the corrective actions for “Excessive venting from relief valves (hissing sound)”</td>
</tr>
<tr>
<td></td>
<td>d) Partial or complete loss of vacuum</td>
<td>Conduct the NER test (RP28) and return the unit to CAIRE, Inc. if necessary.</td>
</tr>
<tr>
<td>10) Excessive Frost</td>
<td>a) Frost is acceptable</td>
<td>Some frost on the outer case and on the plumbing is acceptable, especially at high flow rates. This is due to the evaporation of LOX to gas and the temperature difference between the LOX and room temperature.</td>
</tr>
<tr>
<td>NOTE: Minimal frost on the case and on the plumbing is normal, especially at higher flow rates. This symptom applies to frost that is much greater than what is normally observed.</td>
<td>b) High humidity level</td>
<td>High humidity levels can increase frost accumulation.</td>
</tr>
<tr>
<td></td>
<td>c) Saturation pressure is too high</td>
<td>Inspect the saturation pressure of the reservoir used for filling. Allow at least 30 minutes at no flow for the portable to saturate properly.</td>
</tr>
<tr>
<td></td>
<td>d) Leak in the system</td>
<td>Preform a leak check on the plumbing (RP18). Repair leaks as necessary.</td>
</tr>
<tr>
<td></td>
<td>e) Relief valve open</td>
<td>Ensure that there is no venting from the relief valves. If there is refer to the corrective actions for “Excessive venting from relief valves (hissing sound)”</td>
</tr>
<tr>
<td></td>
<td>f) Partial or complete loss of vacuum</td>
<td>Conduct the NER test (RP28) and return the unit to CAIRE, Inc. if necessary.</td>
</tr>
<tr>
<td>11) Unit will not maintain acceptable pressure when in use</td>
<td>a) Saturation pressure is unaccept- able</td>
<td>Inspect the saturation pressure of the reservoir used for filling. Allow at least 30 minutes at no flow for the portable to saturate properly.</td>
</tr>
<tr>
<td></td>
<td>b) Leak in the system</td>
<td>Preform a leak check on the plumbing (RP18). Repair leaks as necessary.</td>
</tr>
<tr>
<td></td>
<td>c) PRV faulty</td>
<td>Test the operating pressure (RP25) and replace the relief valve if necessary (RP22).</td>
</tr>
</tbody>
</table>
To use the Troubleshooting Chart:

- Start at the upper left corner.
- Unless otherwise noted by the arrows, the flow through the chart is down or to the right.
**RP1 – General**

The following procedures have been carefully prepared to allow proper removal and replacement of defective components and should be used in conjunction with the Troubleshooting Chart and the tests listed in this section.

**WARNING:** Make sure the unit is empty and vent valve is open before replacing any component, except case assembly components or lip seals.

**WARNING:** The technician’s hands, tools, and clothing should be free of all oils and greases.

**WARNING:** Parts that are welded in place must not be replaced in the field. Should these parts fail, return complete assembly or sub-assembly to factory for repair. DO NOT use solder or silver solder to repair broken welds.

**WARNING:** The manufacturer of fluorolubricant warns users not to allow fluorolubricant to contaminate tobacco products. Wash fluorolubricant from hands before smoking.

**WARNING:** Do not use glue type thread locking compounds or unapproved sealants on any repairs.

**CAUTION:** When replacing components, make sure the new part is oriented exactly the same as the original part prior to installation.

**CAUTION:** Some components require a specific amount of torque when assembling. Follow torque requirements where specified.

**NOTE:** All replacement parts must be factory approved, cleaned for oxygen service, and stored in sealed plastic bags. The repair area must be clean and separate from other areas. Room air should be filtered, and free from dust, soot, and other contaminants.

**NOTE:** When replacing components with pipe threads, use PTFE tape thread sealant. Apply two rounds of PTFE tape to threads near end of component, avoiding first thread.

**NOTE:** When assembling new compression fittings, tighten 1/8”, 1/4” and 1/2” nuts eight flats past finger tight and 3/16” nuts five flats past finger tight. When reassembling previously used compression fittings, tighten nuts one to two flats past finger tight.

**WARNING:** Parts that are welded in place must not be replaced in the field. Should these parts fail, return complete assembly or sub-assembly to factory for repair. DO NOT use solder or silver solder to repair broken welds.

**CAUTION:** Some components require a specific amount of torque when assembling. Follow torque requirements where specified.

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**WARNING:** Make sure the unit is empty and vent valve is open before replacing any component, except case assembly components or lip seals.

**WARNING:** The technician’s hands, tools, and clothing should be free of all oils and greases.

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**NOTE:** When assembling new compression fittings, tighten 1/8”, 1/4” and 1/2” nuts eight flats past finger tight and 3/16” nuts five flats past finger tight. When reassembling previously used compression fittings, tighten nuts one to two flats past finger tight.

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**WARNING:** The technician’s hands, tools, and clothing should be free of all oils and greases.

**WARNING:** Make sure the unit is empty and vent valve is open before replacing any component, except case assembly components or lip seals.

**WARNING:** The technician’s hands, tools, and clothing should be free of all oils and greases.

**WARNING:** Parts that are welded in place must not be replaced in the field. Should these parts fail, return complete assembly or sub-assembly to factory for repair. DO NOT use solder or silver solder to repair broken welds.

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**NOTE:** All replacement parts must be factory approved, cleaned for oxygen service, and stored in sealed plastic bags. The repair area must be clean and separate from other areas. Room air should be filtered, and free from dust, soot, and other contaminants.

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**WARNING:** The technician’s hands, tools, and clothing should be free of all oils and greases.
RP3 – Condensate Pad R/R

a. Unzip bottom case zipper.

b. Remove condensate pad and wring out any absorbed moisture. Allow pad to dry completely or dispose (if pad will not be reused).

c. Replace condensate pad and rezip condensate cup.

RP4 – Soft Case R/R

a. Remove strap from the top of the unit.

b. Unzip the top and side zippers.

c. Carefully remove the unit from case.

d. To replace, reverse the above steps.

CAUTION: Be careful not to pinch wires!
Repair Procedures

RP5 – G4 Meter Empty Capacitance Calibration (Figure 9)

a. Completely empty dewar.

NOTE: The meter must first be calibrated empty and then the span must be set for your particular model. The unit must be empty, warm, and dry for calibration.

b. Allow dewar to warm to room temperature.

c. Enter Calibration Mode

1. Press and hold the hidden calibrate button located on the center of the meter face.

2. While holding the calibrate button, press and hold the green operate button.

3. Continue holding both buttons until LED 1 and LED 8 alternately flash. The meter is now in calibration mode.

4. Release both buttons.

d. Press and hold the hidden calibrate button for 3 seconds. LED 1 will flash for 3 seconds followed by a short pause.

NOTE: Step d must be performed within 45 seconds or calibration mode will exit.

e. If successful, LED 1 will flash again for 3 seconds and exit calibration mode storing the new empty value.

f. If calibration is unsuccessful, multiple LEDs will flash. The LEDs indicate what error has occurred (see Table 2).

RP6 – G4 Meter Span Setting Adjustment (Figure 9)

a. Enter Calibration Mode

NOTE: The meter must first be calibrated empty and then the span must be set for your particular model. The unit must be empty, warm, and dry for calibration.

TABLE 2: Calibration Error Codes

<table>
<thead>
<tr>
<th>Error Code</th>
<th>Reason for Fault</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>LED 1 &amp; LED 4</td>
<td>Empty to full span is too low.</td>
<td>Ensure tank is empty. Recalibrate.</td>
</tr>
<tr>
<td>LED 2 &amp; LED 3</td>
<td>Open Circuit</td>
<td>Check meter connections and wire integrity.</td>
</tr>
<tr>
<td>LED 6 &amp; LED 7</td>
<td>Empty capacitance is too high.</td>
<td>Remove moisture from inner vessel. Decontaminate probe.</td>
</tr>
<tr>
<td>LED 5 &amp; LED 8</td>
<td>Empty to full span is too high.</td>
<td>Ensure tank is empty. Recalibrate.</td>
</tr>
<tr>
<td>LED 2, LED 4 &amp; LED 6</td>
<td>High Capacitance</td>
<td>Remove moisture from inner vessel. Decontaminate probe.</td>
</tr>
<tr>
<td>LED 1, LED 3, LED 5 &amp; LED 7</td>
<td>Calibrated full value less than empty value.</td>
<td>Dry harness assembly. Decontaminate.</td>
</tr>
</tbody>
</table>
1. Press and hold the hidden calibrate button located on the center of the meter face.

2. While holding the calibrate button, press and hold the green operate button.

3. Continue holding both buttons until LED 1 and LED 8 alternately flash. The meter is now in calibration mode.

4. Release both buttons.

b. Press the green operate button 3 times within a 5 second period.

c. One of the meter LED’s will light continuously.

d. Press the hidden calibrate button until the LED which matches the Hi Flow Stroller being calibrated is continuously lit. See Table 3 below for the correct LED setting for your unit.

e. Once the correct LED for your unit is lit, press the green operate button to store the setting and exit calibration mode.

NOTE: Step d must be performed within 45 seconds or calibration mode will exit and the meter will NOT be calibrated properly. The calibration procedure will then need to be repeated from the beginning.

c. Press the green operate button 3 times within a 5 second period.

d. Press the hidden calibrate button until the LED which matches the Hi Flow Stroller being calibrated is continuously lit. See Table 3 below for the correct LED setting for your unit.

e. Once the correct LED for your unit is lit, press the green operate button to store the setting and exit calibration mode.

TABLE 3: Preset Capacitance Span LED Setting

<table>
<thead>
<tr>
<th>Model</th>
<th>LED Setting</th>
</tr>
</thead>
<tbody>
<tr>
<td>High Flow Stroller</td>
<td>LED 8</td>
</tr>
</tbody>
</table>

FIGURE 10: Manifold Capacitance Test Setup

RP7 – Manifold Capacitance Test (Figure 10)

a. Capacitance Meter Setup:

1. Connect wire adapter with jumper wire (Item D) to capacitance meter, following color code.

2. Turn on capacitance meter and select 200 pF range.

3. Move zero adjustment on front of meter until display reads zero.

b. Capacitance Test:

WARNING: Make sure the unit is empty, dry, and at room temperature before testing.

1. Remove shoulder strap, unzip the top case zipper, and fold the top flap back in order to access manifold harness connection.

2. Connect wire adapter (Item D) to manifold harness connection (Item A).

3. Connect alligator clip to bracket or any other properly grounded surface.

4. Read manifold capacitance and verify that it is 145–180 pF. If it does not meet specifications, call Technical Support.
Repair Procedures

RP 8 – G4 Meter RR

a. Remove shoulder strap, unzip top case zipper, and fold back top flap.
b. Remove 2 meter mounting screws.
c. Remove meter assembly.
d. Unplug the wiring connection.
e. Disconnect ground wire spade terminals.
f. To replace G4 meter, reverse above procedures.

CAUTION: Be careful not to pinch wires!

RP 9 – Manifold Assembly RR (Figure 11)

WARNING: The unit must be empty, warm and vented before starting procedure.

a. Remove soft case and Gen 4 meter. (RP 4,8).
b. Remove (3) bracket screws (Item 29) and lift bracket (Item 28) from manifold.
c. Loosen and disconnect breathing coil nut (Item 38) from withdrawl fitting and FCV (Item 24). Be careful not to bend or kink Teflon tubing.
d. Loosen and disconnect primary relief valve tube (Item 46) and vent outlet tube from manifold fitting.
e. For top fill units, loosen and remove QDV tube from manifold fitting.
f. Use a hex wrench to loosen and remove the (4) manifold retaining screws (Item 31).
g. Remove manifold assembly (Item 41) by lifting straight up.
i. To replace manifold, reverse above procedure.

NOTE: If substantial force is required, closely examine wire and probe for damage.


RP 10 – Resolder Feed-thru Wire (Figure 11)

a. Remove manifold assembly (See RP 9).
b. Strip approximately 1/8” (0,3 cm) of insulation from feed-thru wire if necessary.
c. Apply small amount of Stay-Clean flux to tinned area of probe using a cotton swab.
d. Resolder feed-thru wire to tinned area of probe. Add small amount of lead-free solid wire solder if necessary.
e. Clean flux residue with distilled water and cotton swab. Dry thoroughly.
f. Replace manifold following listed procedure (RP9).

FIGURE 11: Feed-thru Wire
RP11 – Clean/Dry Probe and Dewar

Procedure 1:

a. Empty and warm dewar per RP 14).
b. Open vent valve.
c. Loosen feed-thru nut and remove harness.
d. After vent valve (Item 50) is thawed, if all moisture is not removed, continue with procedure #2.

Procedure 2:

a. Remove manifold assembly (See RP 9).
b. Blow off probe assembly with clean, dry nitrogen gas.
c. Blow out inside of dewar with clean, dry nitrogen gas until inside is clean and dry.
d. Replace manifold assembly (See RP 9).

RP12 – LOX Fill

Refer to POI for proper fill procedure.

NOTE: Filling source must contain a minimum of 5 liters of properly saturated LOX.

RP13 – Liberator Lip Seal RR (Side Fill)

Refer to Liberator Service Manual.

RP14 – Empty and Warm

a. Turn FCV knob to highest flow setting.
b. Allow unit to sit for 24 hours before proceeding.

RP15 – QDV RR (Side Fill) (Figure 12)

WARNING: The unit must be empty and vented before starting procedure.

a. Remove soft case (RP 4).
b. Loosen QDV retaining nut (Item A).
c. To replace QDV, reverse above procedure, aligning pin in the upright position. Apply thin film of fluorolubricant to O-ring (Item 36). Torque retaining nut to 140–150 inch-lbs (1550–1700 N-cm).
d. Replace soft case (RP 4).

RP16 – QDV and Lip Seal RR (Top Fill) (Figure 13)

WARNING: The unit must be empty and vented before starting procedure.

a. Remove soft case (RP 4).
b. Remove two retaining nuts and lift bottom fill tube out of coil.
c. Replace QDV by substituting old QDV (Item 62) with the new one, tightening Item 57 to 140–150 inch-lbs (1550–1700 N-cm) and Item 58 to 300 inch-lbs (3000 N-cm).
d. Replace lip seal by removing Item 65 and Item 64. Apply a thin film of fluorolubricant to new Item 64 and insert into QDV. Tighten Item 65 to 140–150 inch-lbs (1550–1700 N-cm).
**RP17 – Pressure Retention Test (Figure 14)**

a. Assemble oxygen regulator, pneumatic adapter and hose. Connect assembly to oxygen gas source.

b. Assemble pressure gauge (Item J) and adapter assembly. Thread hose barb onto pressure gauge (use Teflon® tape). Push tubing onto hose bard and attach with clamp.

c. Connect gauge assembly (Item J) to hose barb on FCV outlet. Open FCV (Item 24) to highest flow setting.

d. Connect the proper pneumatic adapter to QDV (Item 37) for side fill units or (Item 62) for top fill.

e. Increase pressure to 19 psig (1.3 bar).

f. Turn FCV valve (Item 24) to Off setting.

g. Allow unit to sit undisturbed for 60 minutes.

h. Turn FCV valve (Item 24) to highest flow setting.

i. If pressure gauge (Item J) is at or above 10 psig (0.7 bar), unit passes test.

**RP18 – Plumbing Leak Test (Figure 14)**

**WARNING:** The unit must be empty and vented before starting procedure.

a. Remove soft case (RP 4)

b. Assemble oxygen regulator, pneumatic adapter and hose. Connect assembly to oxygen gas source.

c. Assemble pressure gauge (Item J) and adapter assembly. Thread hose barb onto pressure gauge (use Teflon® tape). Push tubing onto hose bard and attach with clamp.

d. Connect gauge assembly (Item J) to hose barb on FCV outlet. Open FCV (Item 24) to highest flow setting.

e. Connect the proper pneumatic adapter to QDV (Item 37) for side fill units or (Item 62) for top fill.

f. Increase pressure to 19 psig (1.3 bar).

g. Leak test all connections, joints, and valves with leak test solution.

h. Close FCV (Item 24) by turning to Off position. Remove pressure gauge assembly

i. Disconnect pneumatic adapter from QDV.

j. Leak test QDV poppet and FCV outlet.

k. Repair all leaks by following appropriate repair procedures.

NOTE: PRV (Item 49) and SRV (Item 43) may leak slowly. Repair all other leaks first and retest for pressure retention before changing relief valves.
**RP19 – Vent Valve RR (Figure 15)**

**WARNING:** The unit must be empty and vented before starting procedure.

- b. Open the vent valve.
- b. Remove vent valve (Item 16) by turning counterclockwise.
- c. To replace vent valve, reverse above procedure. Torque nut to 90-100 inch-lbs (1000–1100 N-cm).

**RP20 – FCV RR**

**WARNING:** The unit must be empty and vented before starting procedure.

- b. Disconnect coils from FCV inlet by loosening compression fitting (Item 44), and removing coils.
- c. Remove FCV knob and flow decal.
- d. Disconnect and remove the FCV from bracket by removing two screws.
- e. Remove SRV from FCV.
- f. To replace FCV, reverse above procedure. Tighten screws to 4–6 inch-lbs. (45–70 N-cm). Record FCV serial number.

**RP21 – SRV RR**

- b. Disconnect SRV from FCV (Item 24).
- c. Replace SRV in reverse order. When reassembling SRV, tighten approximately 10 to 20 degrees after the SRV contacts the FCV body (20-30 inch-lbs, 225-350 N-cm minimum).

**RP22 – PRV RR (Figure 18)**

- b. Remove PRV (Item 49) while holding adapter (Item 48).
- c. Replace the PRV using two rounds of Teflon® tape on the threads and tighten 6-7 flats past snug.
- d. Reassemble in the reverse order.
Repair Procedures

RP23A – Breathing Coil RR (Side Fill) (Figure 18)

**WARNING:** The unit must be empty and vented before starting procedure.

a. Remove soft case (RP 4).

b. Loosen the two compression fitting nuts (Items 38 and 44) from liquid withdrawal fitting (Item 41) and FCV (Item 24). Disconnect breathing coil (Item 67), being careful not to kink breathing coil or internal Teflon® tube.

c. Carefully remove breathing coil outlet from around top of dewar.

d. To replace breathing coil, reverse above procedure.

**CAUTION:** The vaporizer coil is fragile. Do not drop coil. Do not hit or mar coil with wrenches.

RP23B – Breathing Coil RR (Top Fill) (Figure 19)

**WARNING:** The unit must be empty and vented before starting procedure.

a. Remove soft case (RP 4).

b. Loosen the two compression fitting nuts (Items 38 and 44) from liquid withdrawal fitting (Item 41) and FCV (Item 24). Disconnect breathing coil (Item 67), being careful not to kink breathing coil or internal Teflon® tube.

c. Loosen QDV tube (Item 56) nut.

d. Loosen QDV jam nut (Item 58). Remove QDV (Item 62) from fill tube.

e. Cut two cable ties holding coil in place. Remove breathing coil by sliding straight down from fill line. Be careful to properly remove breathing coil outlet from around top of dewar.

f. To replace coil, reverse above procedure. Tighten QDV jam nuts to 140–150 inch-lbs. (1550–1700 N-cm).
RP24 – Flow Rate Testing (Figure 20)

a. Unit must be at least 1/2 full with correctly saturated LOX.

b. Set FCV to 2 LPM setting and allow unit to operate for 20–60 minutes.

c. Connect FCV outlet (Item N) to flowmeter inlet (Item M) with respiratory tubing. Make sure flowmeter outlet is open, unobstructed, and properly positioned.

d. Test flow rate at each FCV position. Record all flow rates.

e. Flow rates must be nominal values within tolerances listed in specification section of this manual or unit fails flow rate test.

NOTE: Be careful to allow for accuracy tolerances of flowmeter.

RP25 – Operating Pressure Test (Figure 20)


b. Connect gauge assembly (Item J) to hose barb on FCV outlet. Open FCV (Item 24) to highest flow setting.

c. Read operating pressure on pressure gauge (Item J).

NOTE: If testing operating pressure because of improper flow rates, test immediately after flow rate test.

d. Operating pressure must be 18–22 psig (1.2–1.5 bar) or unit fails test.

RP26 – Flow Meter Verification

a. Flow meter accuracy is best verified by a calibration laboratory. Equipment should indicate liter per minute oxygen gas at atmospheric pressure and 21°C (70°F).

b. Flow meter accuracy may also be tested by comparison to one or more new, unused, calibrated flow meters. This method will increase confidence in accuracy of readings, but not necessarily verify accuracy.
Repair Procedures

RP27 – Case Cleaning

NOTE: Clean case only after unit is empty and warm. Do not clean in oxygen enriched atmosphere.

a. Clean using household glass cleaner or Simple Green D and lint-free cloth. Do not get glass cleaner inside case or onto any plumbing components. Simple Green D is available at www.simplegreend.com

b. If necessary, a soft bristle brush can be used to help remove dirt and debris. If desired, Caire, Inc. Technical Service can supply a specific list of additional recommended cleaning tools.

b. Allow unit to dry thoroughly before filling.

RP28 – Normal Evaporation Rate Test

a. Fill unit following procedure found in the Patient Operating Instructions (POI).

b. Push button and check that all 8 LEDs are on, or verify that the spring scale reads full.

c. Check for leaks in QDV and vent valve.

d. Let unit sit 16–18 hours.

e. Make sure FCV and vent valves are closed.

f. Do not move unit during this time.

g. If the Hi Flow is at least half full (4 LEDs), the unit passes. Maximum loss rate prior to dewar servicing is 1.75 lbs./day (0.8 kg/day).
Required Tools
1. Hex Wrenches (various sizes)
2. Flat Blade Screwdriver
3. 5/16” Nut Driver
4. Open End Wrenches (1/2” to 1-1/8”)
5. Side Cutters
6. Phillips Screwdriver
7. Pliers
8. Torque Driver/Wrenches:
   - 12-17 N-cm (10-15 in-lbs)
   - 23-29 N-cm (20-25 in-lbs)
   - 69-92 N-cm (60-80 in-lbs)
   - 104-115 N-cm (90-100 in-lbs)
   - 6.2-6.9 N-m (45-50 ft-lbs)
9. Jeweler’s Screwdriver

Required Fixtures/Equipment
1. Capacitance Meter
2. Soldering Iron
3. Oxygen Regulator/Hose Kit
4. Pressure Gauge
5. Pressure Gauge Adapter
6. Flowmeter
7. 02 Gas Source (High Pressure bottle)
8. 02 Liquid Source
9. N2 Gas or Clean, Dry Compressed Air Source
10. Tubing (02 compatible)

Required Supplies
1. Stay-Clean Flux
2. Cotton Swabs
3. Lead-free Solder
4. Distilled Water
5. Household Glass Cleaner
6. Lint-Free Cloth
7. Teflon® Tape
8. Fluorolubricant
9. Leak Detection Fluid

Tools and Accessories available from CAIRE, Inc.

<table>
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<tr>
<th>Part No</th>
<th>Description</th>
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<tbody>
<tr>
<td>CA200071</td>
<td>240 AC Fluorolubricant</td>
</tr>
<tr>
<td>CA200072</td>
<td>&quot;Snoop&quot; Leak Detection Fluid</td>
</tr>
<tr>
<td>CA400004</td>
<td>Replacement Filter/Male Transfer Line Adapter</td>
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<tr>
<td>CA406308</td>
<td>10.3 bar (150 psi) Relief Valve Assembly</td>
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<tr>
<td>CA406310</td>
<td>Teflon® Tape</td>
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<tr>
<td>CA406398</td>
<td>10.3 bar (150 psi) Relief Valve only</td>
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<td>Service Manual</td>
</tr>
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<tr>
<td>10018138</td>
<td>C1000 Backpack</td>
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<tr>
<td>97112026</td>
<td>Female Side Fill Pneumatic Test Adapter</td>
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<tr>
<td>97200076</td>
<td>Erie “Liter Meter”</td>
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<tr>
<td>97217007</td>
<td>Pressure Gauge Adapter</td>
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<tr>
<td>97212021</td>
<td>Male Side Fill Pneumatic Test Adapter</td>
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<tr>
<td>97403015</td>
<td>Capacitance Meter</td>
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<tr>
<td>97403016</td>
<td>Jeweler’s Screwdriver</td>
</tr>
<tr>
<td>97405279</td>
<td>Pneumatic Hose with DISS Fittings</td>
</tr>
<tr>
<td>97403574</td>
<td>Dewar Cap</td>
</tr>
<tr>
<td>97403577</td>
<td>0–4.1 bar (0-60 psig) Pressure Gauge</td>
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<tr>
<td>12917730</td>
<td>Stroller Cart</td>
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<tr>
<td>97405147</td>
<td>0–45 psig Oxygen Regulator</td>
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<tr>
<td>97405431</td>
<td>Liquid Oxygen Transfer Line – 2 m (6 ft)</td>
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<td>97404564</td>
<td>Transfer Line Swivel Connector</td>
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<tr>
<td>97405590</td>
<td>Lip Seal Service Tool</td>
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<tr>
<td>97406471</td>
<td>Tandem Tee Kit</td>
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<td>97406555</td>
<td>Super Flex Liquid Oxygen Transfer Line – 6’</td>
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<td>97406630</td>
<td>Dual Fill Head Tee</td>
</tr>
<tr>
<td>13329091</td>
<td>G4 Capacitance Meter Adaptor Kit</td>
</tr>
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</table>
Contact Customer Service or visit www.cairemedical.com to obtain your parts list.
Ordering Information

The following steps should be used when ordering a new Hi Flow, or replacement parts for an existing unit:

1. **Compile a list of all equipment and replacement parts to be ordered.** An exploded view of assemblies for easy parts identification, along with parts/price lists can be found on www.cairemedical.com.

   For US/Canada Part Numbers, please call 770 721 7759.
   For European Part Numbers, please call +44(0) 1344 403100.
   For Asian, Australia, New Zealand Part Numbers, please call +61 297 494333.

2. **Fill out a purchase order containing the following information:**
   a. Purchase order number.
   b. Name and address of billing location.
   c. Name and address of shipping location.
   d. Quantity, part number, description, and unit cost for each item ordered.

3. **Telephone or fax CAIRE, Inc. at one of the numbers listed below to begin immediate processing of the order:**
   - **USA**
     - Toll Free Phone: 800 48 CAIRE (800 482 2473)
     - Toll Free Fax: 888 WE CAIRE (888 932 2473)
     - Phone: 770 721 7758
     - Fax: 770 721 7759
   - **Asia, Australia, Pacific Rim**
     - Phone: +61 297 494333
     - Fax: 888-932-2473
   - **Europe**
     - Phone: +44(0) 1344 403100
     - Fax: +44(0) 1344 429224

4. **Mail or fax the completed purchase order for confirmation to:**
   - CAIRE, Inc.
     - 2200 Airport Industrial Drive
     - Suite 500
     - Ball Ground, GA 30107
   - Chart BioMedical Ltd.
     - Unit 6, Ashville Way
     - Wokingham, Berkshire, RG41 2PL
     - United Kingdom

All new equipment will be shipped either “prepaid”, F.O.B. Canton, Georgia, F.O.B. Bracknell, UK, or collect via your specified carrier. All replacement parts will be sent by UPS “prepaid”, and the shipping charges for equipment and parts will be added to the final invoice. Payment for replacement parts are located on CAIRE, Inc.’s invoice with payment date indicated. All shipments will originate from Canton, Georgia or Bracknell, UK. If a particular carrier or method of shipment is desired, specify when placing order.

For additional ordering and contact information, visit www.cairemedical.com
When a Hi Flow Stroller is received, it should be inspected immediately, as outlined in Section VII, Unpacking and Setup Instructions.

If a problem with the unit should be encountered, reference should be made to the Troubleshooting Charts on pages 17-21. If these procedures do not provide a solution for the problem, the following steps should be taken:

1. Call CAIRE, Inc. Customer Service. State the problem with the unit. If it is determined that the problem cannot be solved by the distributor, a Return Material Authorization (RMA) number will be assigned to the unit or part(s). If a Purchase Order Number is to be referenced, please give this number to the Customer Service Representative at that time.

2. Carefully package the parts, or repack the unit in its original shipping container, precisely as shipped.

3. Write the Return Material Authorization Number on the top of the shipping container.

4. Return the unit or parts by professional carrier to:
   
   CAIRE, Inc.
   2205 Airport Drive
   Ball Ground, GA 30107

   Chart BioMedical Ltd.
   Unit 6, Ashville Way
   Wokingham, Berkshire, RG41 2PL
   United Kingdom

   All equipment returned to CAIRE, Inc. must be shipped "prepaid".

When the defective item(s) is received at CAIRE, it will be serviced and returned to the distributor as soon as possible. A copy of the "Repair Cost Sheet" will be enclosed giving a detailed listing of any maintenance performed.

Restocking Policy

If it becomes necessary to cancel an order with CAIRE, Inc. after the shipment has been received, use the following "Restock Policy" procedure:

1. Notify the Customer Service Department at CAIRE, Inc. using the toll-free number. When contacting Customer Service personnel, it will be necessary to relay the following information:
   
   a. State the quantity and description of equipment to be returned.
   b. Give the Serial Number of each unit to be returned.
   c. State the equipment purchase date.

2. An RMA number will be issued in the name of the distributor by CAIRE, Inc. for the equipment to be returned. When the equipment is shipped to the factory, the RMA number must appear on the packing slip.

3. All equipment must be returned "prepaid" to the original shipping location:
   
   CAIRE, Inc.
   2205 Airport Drive
   Ball Ground, GA 30107

   Chart BioMedical Ltd.
   Unit 6, Ashville Way
   Wokingham, Berkshire, RG41 2PL
   United Kingdom

   4. Finally, a "Credit Memo", minus a 15% restocking fee, will be issued to the distributor when all equipment has been received, inspected, and restocked by CAIRE, Inc.

Return of Unused Non-Defective Merchandise

CAIRE, Inc., at its discretion, charges a 15% restocking fee for unused non-defective merchandise that is returned. A RMA number must be obtained from CAIRE, Inc. Customer Service prior to return of any goods. Merchandise cannot be returned for credit after sixty (60) days. Customer to pay all freight charges. Tracking capability and insurance on all returned goods is advised. CAIRE, Inc. will not be responsible for misdirected shipments.