INOmax DS\textsuperscript{IR} Plus

Operation Manual
(English)
Series 3 software
User Responsibility

This Product will perform in conformity with the description contained in this operating manual and accompanying labels and/or inserts, when assembled, operated, maintained and repaired in accordance with the instructions provided. This Product must be checked prior to use following the Pre-Use Checkout procedure described in section two. A defective Product should not be used. Parts that are broken, missing, visibly worn, distorted or contaminated should be replaced immediately.

Should such repair or replacement become necessary, the manufacturer recommends that a telephone request for service advice be made to the local distributor. This Product or any of its parts should not be repaired other than in accordance with written instructions provided by the manufacturer or local distributor. The Product must not be altered.

The user of this Product shall have the sole responsibility for any malfunction which results from improper use, faulty maintenance, improper repair, damage, or alteration by anyone other than INO Therapeutics LLC.

Caution: U.S. Federal and Canadian law restrict this device to sale by or on the order of a licensed medical practitioner. Outside the U.S.A. and Canada, check local laws for any restrictions that may apply.

Inhaled Nitric Oxide mixtures must be handled and stored in compliance with federal, state and local regulations.

INO Therapeutics LLC products have unit serial numbers with coded logic which indicate the year of manufacture and a sequential unit number for identification.

Important:
Before using the INOmax DSIR, Plus read through this manual.
Read through the manuals for the ventilator, humidifier and any other accessory items used. Follow the manual instructions and obey the Warnings and Cautions.
Keep this manual readily available to answer questions.

<table>
<thead>
<tr>
<th>SN 20151234</th>
<th>The first four numeric digits indicate the year of product manufacture, and the next 4 digits are the sequential unit number produced.</th>
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</thead>
<tbody>
<tr>
<td>Ref 10023</td>
<td>INOmax DSIR, 800 ppm, English - Australia</td>
</tr>
<tr>
<td>Ref 10085</td>
<td>INOmax DSIR, 400 ppm, English - Europe</td>
</tr>
<tr>
<td>Ref 10086</td>
<td>INOmax DSIR, 800 ppm, English - Europe</td>
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Open Source Software
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No license is conveyed, either expressed or implied, with the purchase or usage hereof under any patent or patent application covering this product, including but not limited to U.S. Patent 5,485,827, 5,873,359, 5,558,083 and any respective foreign equivalents thereof.
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Changing Cylinders

- Only use manufacturer supplied drug cylinders, regulators and adapters (see Changing INOMAX Cylinders and Purging the Regulator Assembly, Section 3/ Patient Application).

High Frequency Oscillatory and Jet Ventilator Circuits

- Some high frequency ventilator circuits require a one-way valve to prevent high NO delivery.
- Place the Bunnell Life Pulse in Standby prior to suctioning the patient to avoid NO delivery transiently exceeding the set dose by up to 30 ppm for 800 ppm cylinders. Press ENTER to reestablish ventilation as soon as the catheter is removed from the airway. This will limit the extent of over delivery above the NO set dose.
- Do not use dose settings above 40 ppm when using the HFO option with the Acutronic Fabian HFO ventilator. Bidirectional flow through the Injector Module may cause over-delivery which can lead to measured NO values greater than 100 ppm.

Integrated Pneumatic Backup

- The integrated pneumatic backup is intended for short term use when the electronic delivery system fails until a replacement NO delivery device can be brought to the bedside.
- The integrated pneumatic backup delivers a variable concentration of NO to the patient depending on the ventilator flow being used.
- When using the integrated pneumatic backup with breathing circuit gas flows of 5 L/min, the delivered NO dose will be approximately 40 ppm (800 ppm cylinder) or 20 ppm (400 ppm cylinder). Breathing circuit gas flows less than 5 L/min will deliver an NO dose greater than 40 ppm (800 ppm cylinder) or 20 ppm (400 ppm cylinder).
- The integrated pneumatic backup (250 mL/min.) should not be used with the Bunnell Life Pulse as ventilator flow rates can be very low at times, creating a potential delivered NO dose greater than 80 ppm.
**Maintenance**

- Handle and dispose of sensors according to facility biohazard policies. Do not incinerate.
- Use only RS 232 cables that are shielded (see Section 8/ Specifications for more detail).
- If the injector module has been used in the wet/humidified part of the breathing circuit, it should be sterilized between each patient use.

**Manually Bagging a Patient with an Injector Module**

- The hyperinflation bag will, under some conditions, contain NO₂ in excess of one ppm. Use of large tidal volume breaths may expose the patients to the NO₂ present in the bag for part of the breath. In general, if the inspiratory flow rate induced by the manual ventilation does not exceed the fresh gas flow rate, the patient should not be exposed to the concentrations of NO₂ present in the hyperinflation bag.
- Adult and infant hyperinflation bags generate more NO₂ when used at lower minute ventilation. If use of the bag is interrupted (for example to adjust the tracheal tube), before resuming ventilation of the patient, the user should squeeze the bag several times to empty residual gas from the bag.
- Because of the potential for inhalation of excessive concentrations of NO₂, and the difficulty in monitoring the peak inhaled NO₂ concentrations, ventilation with a hyperinflation bag or self-inflating bag is intended only for short term use.
- The monitoring system within the INOmax DSIR Plus will not detect generation of NO₂ within the hyperinflation bag or self-inflating bag devices and the alarms for excessive NO₂ cannot warn of NO₂ produced within the manual bag system.
- To minimize the delivered concentration of NO₂, the following steps should be taken for use with the manual resuscitator bags:
  - Concentrations greater than 20 ppm NO should not be used because of excessive NO₂ generation.
  - Use the smallest bag adequate to deliver the desired tidal volume.
  - Oxygen tubing lengths greater than 72 inches should not be used (between the injector module and the bag).
  - Use the highest fresh gas flow rate (up to 15 L/min) that is practical.
  - Use the lowest practical inspired oxygen concentration.
  - After starting fresh gas flow, squeeze the bag several times to empty residual gas in the bag prior to using the system to ventilate a patient.

**WARNING:**
**WARNING:**

*Manually Bagging a Patient with the INOblender*

- The purge procedure must be followed to help ensure NO₂ is purged from the system before the manual resuscitator bag is connected to the patient.
- The manual bag should be squeezed repeatedly during use to avoid NO₂ building up in the bag.
- If the bag is not squeezed repeatedly while delivering INOMAX, the bag should be removed from the patient and the bag purge procedure performed before continuing.
- The INOblender should be upright when setting the oxygen flowrate for accurate setting.
- Do not use pneumatically powered nebulizers with the INOblender. This will result in significant over delivery of INOMAX in excess of 80 parts per million (ppm) with 800 ppm cylinders and 40 parts per million with 400 ppm cylinders.
  - The INOblender outlet pressure has been validated for use up to 400 millibar (5.8 psig) pressure. The amount of back-pressure generated by pneumatic nebulizers is significantly greater 1.4 to 2.0 bar (20-30 psig) and will result in over delivery of INOMAX in excess of 80 ppm. The user adjusted dose setting on the INOblender will not correlate with, or have an effect on the actual delivered dose.
  - In addition, the INOblender flowmeter is not back-pressure compensated and will display a lower flow rate than actual when pressure is applied to the outlet.

*Purging the INOmax DSIR Plus*

- All INOmax DSIR Plus devices must be purged before use to ensure the patient does not receive an excess level of NO₂.
- If the INOmax DSIR Plus is not going to be used on a patient within 10 minutes, depressurize the regulator supply line.
- If the INOmax DSIR Plus is not used and is pressurized for more than 10 minutes, repeat automated or manual purge procedure.
- If the INOmax DSIR Plus is depressurized and not used within 12 hours, repeat pre-use procedure.

*Transport*

- If the INOmax DSIR Plus or INOblender is to be used in a transport vehicle, they should be affixed to the transport mounting post, which is part of the transport mounting bracket assembly (part number 50041).
- The transport mounting post and/or the transport mounting bracket assembly should be secured to the transport isolette/transport gurney in a manner which will secure the INOmax DSIR Plus/INOblender.
- Only use one length of extension hose (part number 10014) between devices to minimize the risk of NO₂ formation within the hose.
**WARNING:**

**Troubleshooting or Calibrating**

- If an alarm occurs, safeguard the patient first before troubleshooting or repair procedures.
- Use caution when troubleshooting the INOmax DS_{IR} Plus delivery system while in use for a patient.
- Abrupt discontinuation of INOMAX may lead to worsening oxygenation and increasing pulmonary artery pressure, i.e., Rebound Pulmonary Hypertension Syndrome. To avoid abrupt discontinuation, utilize the NOblender or integrated pneumatic backup if necessary. If Rebound Pulmonary Hypertension occurs, reinstate INOMAX therapy immediately (See the INOMAX prescribing Information for further details.)
- If the high NO_2 alarm activates, the delivery system should be assessed for proper set up while maintaining INOMAX delivery. Adjust the dose as described in the INOMAX Prescribing Information-The Effects of Nitrogen Dioxide. If unable to determine the cause of the increased NO_2 levels, call Customer Support, do not discontinue therapy.
- Do not change any sensor while delivering NO to a patient.
- Loss of communication between the INOmax DS_{IR} Plus and the INOMAX cylinder for more than one hour will result in interruption of INOMAX delivery.

**Use Outside of Product Labeling**

- The INOmax DS_{IR} Plus must only be used in accordance with the indications, usage, contraindications, warnings and precautions described in the INOMAX (nitric oxide) drug package inserts and labeling. Refer to this material prior to use.
- Helium/oxygen mixtures should not be used with the INOmax DS_{IR} Plus.
- The use of devices which radiate high-intensity electrical fields may affect the operation of the INOmax DS_{IR} Plus. Constant surveillance of all monitoring and life support equipment is mandatory whenever interfering devices are in operation on or near a patient.
- The approved patient population for the INOmax DS_{IR} Plus, as specified in the drug labeling for INOMAX (nitric oxide) for inhalation, is limited to neonates. The INOmax DS_{IR} Plus is not intended to be used in other patient populations.
- Outside of the United States, use of the INOmax DS_{IR} Plus is limited to the use in accordance with INOMAX or INOflo, nitric oxide for inhalation prescribing information as established with the national health authority.
WARNING:

Ventilators and Breathing Devices

- The INOmax DSIR Plus subtracts gas from the breathing circuit via the gas sampling system at 230 mL per minute which can cause the ventilator to auto-trigger. Adjusting the flow sensitivity may be necessary. The trigger sensitivity of the ventilator should be checked after connecting the INOmax DSIR Plus to the breathing circuit.
- Set the INOmax DSIR Plus alarm thresholds for the current patient conditions to monitor any inadvertent changes in treatment.
- Be certain all cables and hoses are positioned to help prevent damaging or occluding them.
- The use of pediatric and neonatal ventilator settings with adult size breathing circuits can result in higher levels of NO₂. Always use the size of breathing circuit that is appropriate for the patient.
- The humidifier chamber volume should not be more than 480 mL to prevent elevated NO₂ values.
- The INOmax DSIR Plus should not be used with the BiPap Vision system or other single-lumen breathing systems with bidirectional flow, as over-dose of INOMAX (nitric oxide) and interruption of drug delivery to the patient may occur.
- The patient gas sample tee must have the INOmax DSIR Plus sample line attached or be capped off to avoid loss of ventilator circuit pressure.
- Avoid recirculation of gases. Undesired recirculation of gases will occur if fresh gas flows are less than the patient minute volume.
- Only use parts/accessories designated for use with this system.
INOmax DS_{IR} Plus

1/ General Information
INOmax DS<sub>IR</sub>® Plus

1/ General Information
1/ General Information

Indications for Use

• The INOmax DS_{IR} Plus (delivery system) delivers INOMAX® (nitric oxide for inhalation) therapy gas into the inspiratory limb of the patient breathing circuit in a way that provides a constant concentration of nitric oxide (NO), as set by the user, to the patient throughout the inspired breath. It uses a specially designed injector module, which enables tracking of the ventilator waveforms and the delivery of a synchronized and proportional dose of NO. It may be used with most ventilators.

• The INOmax DS_{IR} Plus provides continuous integrated monitoring of inspired O_2, NO_2, and NO and a comprehensive alarm system.

• The INOmax DS_{IR} Plus incorporates a battery that provides up to six hours of uninterrupted INOMAX delivery in the absence of an external power source.

• The INOmax DS_{IR} Plus includes a backup NO delivery capability that provides a fixed flow of 250 mL/min of 800 ppm NO which along with user supplied 10 L/min of oxygen provides 20 ppm (10 ppm with 400 ppm NO gas) in the gas flow to a patient’s breathing circuit. It may also use the INOblender for backup.

• Use of the INOmax DS_{IR} Plus is limited to the use in accordance with INOMAX nitric oxide for inhalation prescribing information as approved with the national health authority.
# Introduction to this Manual

## Definitions and abbreviations

<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>% v/v</td>
<td>% volume/volume</td>
</tr>
<tr>
<td>Breathing circuit</td>
<td>Part of ventilator or breathing system that connects to the INOmax DSIR® Plus.</td>
</tr>
<tr>
<td>Breathing system</td>
<td>Non-invasive breathing devices.</td>
</tr>
<tr>
<td>Control wheel</td>
<td>Rotary control used to change and confirm settings.</td>
</tr>
<tr>
<td>Cylinder</td>
<td>Aluminum cylinder containing INOMAX® therapy gas.</td>
</tr>
<tr>
<td>HFOV</td>
<td>High frequency oscillatory ventilation.</td>
</tr>
<tr>
<td>INOblender®</td>
<td>Backup to the INOmax DSIR® Plus. Allows manual ventilation of the patient, providing uninterrupted delivery of INOMAX.</td>
</tr>
<tr>
<td>INOMAX</td>
<td>Nitric oxide for inhalation.</td>
</tr>
<tr>
<td>INOmeter®</td>
<td>Counter mounted on a cylinder that records the amount of time the INOMAX cylinder valve is open.</td>
</tr>
<tr>
<td>Infrared (IR)</td>
<td>Infra-red technology by which the INOmax DSIR® Plus communicates with the INOmeter mounted on each cylinder.</td>
</tr>
<tr>
<td>N₂</td>
<td>Nitrogen.</td>
</tr>
<tr>
<td>NO</td>
<td>Nitric oxide.</td>
</tr>
<tr>
<td>NO₂</td>
<td>Nitrogen dioxide.</td>
</tr>
<tr>
<td>O₂</td>
<td>Oxygen.</td>
</tr>
<tr>
<td>ppm</td>
<td>Parts per million.</td>
</tr>
<tr>
<td>Pre-use circuit</td>
<td>Connectors and tubing assembly required for INOmax DSIR® Plus for pre-use checkout.</td>
</tr>
<tr>
<td>psig</td>
<td>Pounds per square inch gauge.</td>
</tr>
<tr>
<td>Set NO</td>
<td>Dose of INOMAX set by the user.</td>
</tr>
</tbody>
</table>
This manual shows the Set NO displays associated with the 0-80 ppm range.

Figure 1-1 INOmax DS$_{IR}$ Plus Front View

Figure 1-2 INOmax DS$_{IR}$ Plus Rear View

1. Sample Line Inlet
2. Main Power Indicator
3. Display Screen
4. Alarm Speaker (under front label)
5. Integrated Pneumatic Backup Switch
6. Control Wheel
7. Injector Module Tubing Outlet
8. Injector Module Cable Inlet
9. Water Bottle
10. Purge Port
11. INOMAX Gas Inlets
12. INOblender Gas Outlet
13. Ethernet Port
14. Infrared Connector
15. USB Port
16. Water Separator Cartridge
17. Water Bottle
18. Sample Gas Outlet Port
19. Clamp Assembly
20. Electrical Cord Inlet
21. Equipotential Terminal
22. ON/Standby Switch
23. RS 232 Port
Figure 1-3 INOmax DSIR Plus and Cart (shown with 88-size, 800 ppm cylinders)
Navigating the Display Screens

**Note:** The specific level is identified by the highlighted card on the Menu Button. The red arrows indicate going back to a previous screen.
Menu Screen (second level)

Pre-Use Wizard

Automated Purge Screen

Alarm History Screen

Low Calibration Screen

High Calibration Screen

Settings Screen
Main Display Screen

- On the main screen the user can view alarm messages, monitored values and graphical information.
- By pressing the “Menu Button” on the touch screen (top right hand corner), the user can access the menu screen (see Figure 1-5).

Menu Screen (second level)

- On the menu screen the user can access the Pre-Use Checkout (#1) and the Auto Purge (#2) wizards.

  Note: The Pre-Use Checkout and Auto Purge buttons are inactive (greyed out) if a dose is set.

- To review the complete alarm history, press the Alarm History button (#5), (refer to Section 5/ Alarms and Troubleshooting).
- To initiate a low (room air) or high calibration, press either the Low Cal (#9) or High Cal (#7) buttons. (refer to Section 6/ Calibration).
- Press the Settings button (#6) to view circuit flow and calculated delivery graphs, change display brightness, change alarm volume, change time zone and view software revision (see Figure 1-6).
**Settings Screen (third level)**

- The circuit flow graph, combined with calculated delivery graph, is a user level tool to ascertain NO delivery system limitations in the context of mechanical ventilation.

- The circuit flow rate graph displays the real time peak and average flow rate in the breathing circuit over a 10 second time period, as measured by the injector module. The area in green represents the circuit flow range where the INOMax DSIR Plus system is rated to deliver NO from 1-80 ppm for 800 ppm cylinders and 1-40 ppm for 400 ppm cylinders. (see maximum NO delivery graph page 1-25). Display graphic areas in yellow represents where some inaccuracy of NO delivery is to be expected.

- The calculated delivery graph displays the delivered dose as calculated by the delivery system. The system calculates the dose using the known variables of flow through the injector module, INOMAX cylinder concentration and set dose. The green zone represents that the delivered dose is within +/- 20% of the set dose, the yellow indicates a delivered dose greater (+) or less than (-) 20% of the set dose (see formula below).

**Note:** If the NO dose is not set, the Calculated Delivery graph will remain inactive.

**Calculated Delivery Formula:**

\[
\text{Calculated Delivery} = \left( \frac{\text{NO flow}}{\text{NO flow} + \text{ventilator flow}} \right) \times \text{cylinder concentration}
\]
Display and user controls

The INOmax DSIR Plus has a color touch screen display and a control wheel for adjusting and entering user settings. The buttons on the touch screen and the control wheel perform a variety of functions using a three-step procedure (see “Setting and making changes on the INOmax DSIR Plus” see page 1-12).

The touch screen buttons and control wheel are used to:

- Set the concentration of delivered NO
- Adjust alarm limits
- Silence alarms
- Calibrate the sensors
- Review alarm history
- Define setup options
- Enter patient information

Note:

- If a button has been selected and no activity has been sensed within 20 seconds, the display will return to its previous condition. If a button is greyed out, it is not active.
- Position delivery system so user screen is unobstructed and the speaker is not covered.

When a value is being changed, pressing the "Cancel Active Status" button during editing will stop the change and return the parameter to its original value (similar to the escape key on a computer).
Main Screen

Cylinder icons are not visible and the NO delivery setpoint button will remain inactive until the INOmax DSIR Plus recognizes an INOMAX cylinder.

**Caution:** High frequency and/or high intensity light emission, in the area of the INOmeter, may interfere with communication between the INOmax DSIR Plus and the INOmeter on the INOMAX cylinder (see Section 5/ Alarms and Troubleshooting).

The cylinder icons will appear on the main screen in relation to their position on the cart when the user is facing the INOmax DSIR Plus.

**Note:** When using the transport regulator/cap assembly (PN 10022, CGA or 10041, ISO) only one cylinder will be displayed.

When an INOMAX cylinder valve is opened, the cylinder handle graphic will turn green representing an open INOMAX cylinder valve.
Setting and making changes on the INOmax DSIR Plus

Dose settings

The range of NO dose settings for the INOmax DSIR Plus is configured by the distributor for a specific INOMAX (nitric oxide) concentration, e.g., 400 ppm or 800 ppm. If configured for 400 ppm the range of NO dose setting is 0 to 40 ppm and if configured for 800 ppm the range is 0 to 80 ppm (see below).

Adjusting Parameters (example: dose setting)

1. SELECT (press) a button on the touch screen associated with the desired function. (An audible beep will sound when a button is selected, and the button will be displayed in inverse video.)

2. ROTATE the control wheel clockwise or counterclockwise to adjust the value.
3. CONFIRM the selection by pressing the control wheel or the button associated with the desired function again.

**Caution:** A two minute monitoring alarm delay will prevent the low NO monitoring alarm from occurring while the measured values stabilize.

**Note:**
- After confirming a desired dose, the NO dose setting indicator will fill to the set dose, and the alarm setting (high and low) will automatically be set for the first setting only.
- Any other changes will require the high and low alarm settings to be adjusted.
Settings Screen Adjustments

Access the settings screen (third menu level).

Display Brightness setting
1. Select the display brightness button on the touch screen.
2. Rotate the control wheel to indicate the display brightness level desired. Choices range from one (darkest) to 10 (brightest).
3. Confirm the selection by pressing the control wheel or the display brightness button again.
4. When finished with the menu screen, push the return to previous level button on the touch screen.

Alarm Volume setting
1. Select the alarm volume button on the touch screen.
2. Rotate the control wheel to indicate the volume level desired. Choices range from one (softest) to five (loudest).
3. Confirm the selection by pressing the control wheel or the alarm volume button again.
4. When finished with the menu screen, push the return to previous level button on the touch screen.

Time Adjust setting
If the "Time" button is pressed the Time Adjust screen will appear.
1. Select the Hour or Minute button on the touch screen.
2. Rotate the control wheel to adjust the displayed hour or minute.
3. Confirm the selection by pressing the control wheel or the Hour or Minute buttons again.
4. When finished with the menu screen, push the return to previous level button on the touch screen.

Note: Changing the displayed time does not impact the time written to the INOmeter since the time written to the INOmeter is GMT time, not the displayed time.
Infrared Communication between the INOMAX Cylinders and the INOmax DS\textsubscript{IR} Plus

**WARNING:** Loss of communication between the INOmax DS\textsubscript{IR} Plus and the INOmeter for more than one hour will result in interruption of INOMAX delivery.

The INOmax DS\textsubscript{IR} Plus has an interface using infrared (IR) technology which allows the INOmax DS\textsubscript{IR} Plus to communicate with the INOmeter (which is mounted to each INOMAX cylinder). The INOmax DS\textsubscript{IR} Plus checks the INOMAX cylinder for the correct expiration date and cylinder concentration. The INOmax DS\textsubscript{IR} Plus also transmits a confirmed patient identifier to the INOmeter on any open INOMAX cylinder.

The INOmax DS\textsubscript{IR} Plus cart has a cover (see Figure 1-7, 1) with an infrared transceiver mounted directly above each INOMAX cylinder. When INOMAX cylinders are loaded, communication will take place between the INOmax DS\textsubscript{IR} Plus and the INOmeter (see Figure 1-7, 2) after the boot up phase of the INOmax DS\textsubscript{IR} Plus is complete. A cylinder icon will be displayed on the main screen when an INOMAX cylinder is recognized by the INOmax DS\textsubscript{IR} Plus (see "Loading INOMAX Cylinders onto the INOmax DS\textsubscript{IR} Plus Cart", page 1-17).

**Caution:** Nothing should be placed between the INOmeter and the cart to which it is attached.

**IR Communication Interference**

The INOmax DS\textsubscript{IR} Plus transceiver is located under the cart cover and should be protected from outside IR sources. The INOmax DS\textsubscript{IR} Plus cart was designed to protect the INOmeter from external light/IR energy sources. The INOmax DS\textsubscript{IR} Plus transceiver transmits via a 30 degree transmission cone projecting towards the floor (see dotted lines in Figure 1-7). The specifications of the IR beam call for it to have a range of 20 cm (7.9 in). Based on these specifications it should not affect other devices in the vicinity of the INOmax DS\textsubscript{IR} Plus.

The INOmeter uses a lower energy source which results in a lower IR beam range than that of the INOmax DS\textsubscript{IR} Plus cart. The INOmeter does not transmit IR signals unless it is mounted on the INOmax DS\textsubscript{IR} Plus cart.

**Caution:** A strong magnetic field could affect the ability of the INOmeter to detect if the cylinder valve is opened or closed. This may affect the ability of the INOmax DS\textsubscript{IR} Plus to detect the position (open or closed) of the cylinder valve.

If there is interference with the INOmax DS\textsubscript{IR} Plus/INOmeter communication, the cylinder icon on the user screen will not be displayed and a “Cylinder Not Detected” alarm will activate if there is a set INOMAX dose.

If IR communication interference occurs, we recommend taking the following actions:

- Move the external IR source.
- Move the INOmax DS\textsubscript{IR} Plus cart to reduce the external IR source in the area of the INOmeter.
- Shield the INOmeter from the suspect IR source.

If the actions listed above do not remedy this issue, the transport regulator/cap assembly (PN 10022, CGA or 10041, ISO) may be utilized.
External Light Interference

**Caution:** High frequency and/or high intensity light emission, in the area of the INOmeter, may interfere with communication between the INOmax DS<sub>IR</sub> Plus and the INOmeter on the INOMAX cylinder.

If there is interference with the INOmax DS<sub>IR</sub> Plus/INOmeter communication, the cylinder icon on the user screen will not be displayed and a “Cylinder Not Detected” alarm will activate if there is a set INOMAX dose.

Test results have demonstrated susceptibility to unintended infrared energy from artificial light sources. Most notably, various compact fluorescent lighting fixtures that focus or reflect light, increasing the light intensity in the vicinity of the INOmax DS<sub>IR</sub> Plus cart, could affect INOmeter communications.

If external light interference occurs, we recommend taking the following actions:

- Move the interfering light source.
- Move the INOmax DS<sub>IR</sub> Plus cart to reduce the high intensity light in the area of the INOmeter.
- Shield the INOmeter from the suspect light source.

If the actions listed above do not remedy this issue, the transport regulator/cap assembly may be utilized.
Loading INOMAX Cylinders Onto the INOMax DSIR Plus Cart

Ensure all INOMAX gas cylinders contain more than 35 bar (500 psig).

Note:
- The INOMax DSIR Plus checks INOMAX cylinders for the correct product identity labels, cylinder concentration and expiration date.
- The INOMax DSIR Plus recognizes the drug as expired on the first day of labeled expiration month on the INOMAX cylinder.

Loading the first INOMAX cylinder on the cart 1 will result in a cylinder icon displayed on the screen 2.

Loading a second INOMAX cylinder onto the cart 3 will result in a second cylinder icon displayed on the screen 4.
INOmeter Operation

• The INOmeter is a time-metric device which records the amount of time the INOMAX cylinder valve is opened.

• When used with INOmax DS\textsubscript{IR} Plus, two-way infrared (IR) communication occurs between the INOmax DS\textsubscript{IR} Plus and the INOmeter. The INOmeter communicates the INOMAX cylinder concentration and the expiration date to the INOmax DS\textsubscript{IR} Plus. Patient ID (when confirmed) and dose information are communicated from the INOmax DS\textsubscript{IR} Plus to the INOmeter.

Note:
• Cylinders are shipped with the INOmeter covered in a tamper-proof seal.
• A valve lock is secured to the cylinder by a lanyard.
• The lock must be removed to open the cylinder valve for use.

1. Remove and properly dispose of tamper-proof seal or covering (see Figure 1-8).

2. The lock is secured to the cylinder by a lanyard (see Figure 1-9).
5. When the INOmeter is turned ON (cylinder open) the display will show a "+" (positive) sign (see Figure 1-12) and alternate between:
   a. The event time XX.X in hours since turned ON (eight seconds).
   b. The total cumulative time XXX in hours of all the ON events (four seconds).
   c. The display alternates between a and b for the indicated times in parenthesis above.

   **Note:**
   - If display is blank, replace cylinder.

6. When the INOmeter is turned OFF (cylinder closed) the display will show a "-" (negative) sign (see Figure 1-13) and alternate between:
   a. Showing - - - on the display (eight seconds).
   b. The total cumulative time XXX in hours of all the ON events (four seconds).
   c. The display alternates between a and b for the indicated times in parenthesis above.

   **Note:**
   - If display is blank, replace cylinder.

---

3. Press lock downward to remove from the INOmeter (see Figure 1-10).

4. The cylinder must be closed to reinsert the lock. Align directly across from the iButton and press upward into socket to attach lock (see Figure 1-11).

   **Note:**
   - The INOmeter is used to open and close the cylinder valve.
   - Counter-clockwise rotation of the INOmeter serves to open the cylinder valve (see Figure 1-12).
   - Clockwise rotation acts to close the cylinder valve (see Figure 1-13).

---

Note: If display is blank, replace cylinder.
When the cylinder valve is open and delivery is normal, the main screen shows the handle as green (see Figure 1-14).

**Note:** When two INOMAX cylinders are loaded onto the cart and if both cylinder images do not appear on user screen, check to see if IR or light interference is suspected (see Section 5/ Alarms and Troubleshooting). If there is no light interference, replace suspected right or left INOMAX cylinder.
### Symbols used in this manual or on the system

Symbols replace words on the equipment and/or in this manual. These symbols include:

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="Alarm Silence" /></td>
<td>Alarm Silence</td>
</tr>
<tr>
<td><img src="image" alt="134°C" /></td>
<td>Autoclavable</td>
</tr>
<tr>
<td><img src="image" alt="Average Flow Rate" /></td>
<td>Average Flow Rate</td>
</tr>
<tr>
<td><img src="image" alt="Calculated Dose Greater than 20% of the Set Dose" /></td>
<td>Calculated Dose Greater than 20% of the Set Dose</td>
</tr>
<tr>
<td><img src="image" alt="Calculated Dose Less than 20% of the Set Dose" /></td>
<td>Calculated Dose Less than 20% of the Set Dose</td>
</tr>
<tr>
<td><img src="image" alt="CE European Representative" /></td>
<td>CE European Representative</td>
</tr>
<tr>
<td><img src="image" alt="CE Mark" /></td>
<td>CE Mark</td>
</tr>
<tr>
<td><img src="image" alt="Do Not Push" /></td>
<td>Do Not Push</td>
</tr>
<tr>
<td><img src="image" alt="EHR" /></td>
<td>Electronic Health Record</td>
</tr>
<tr>
<td><img src="image" alt="Equipotential Stud" /></td>
<td>Equipotential Stud</td>
</tr>
<tr>
<td><img src="image" alt="Ethernet Port" /></td>
<td>Ethernet Port</td>
</tr>
<tr>
<td><img src="image" alt="Fuse Rating" /></td>
<td>Fuse Rating</td>
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<tr>
<td><img src="image" alt="Infrared Input/Output" /></td>
<td>Infrared Input/Output</td>
</tr>
<tr>
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<td>Injector Module</td>
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<tr>
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<td>Keep Dry</td>
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<td>Lot Number</td>
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<td>Main Power Connected</td>
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<td><img src="image" alt="Maximum" /></td>
<td>Maximum</td>
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<tr>
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<td>NO Backup OFF</td>
</tr>
<tr>
<td><img src="image" alt="NO Backup ON" /></td>
<td>NO Backup ON</td>
</tr>
<tr>
<td><img src="image" alt="NO Gas Inlet" /></td>
<td>NO Gas Inlet</td>
</tr>
<tr>
<td><img src="image" alt="NO Gas Outlet" /></td>
<td>NO Gas Outlet</td>
</tr>
<tr>
<td><img src="image" alt="On" /></td>
<td>On</td>
</tr>
<tr>
<td><img src="image" alt="Peak Flow Rate" /></td>
<td>Peak Flow Rate</td>
</tr>
<tr>
<td><img src="image" alt="Pneumatic Inlet" /></td>
<td>Pneumatic Inlet</td>
</tr>
<tr>
<td><img src="image" alt="Pneumatic Outlet" /></td>
<td>Pneumatic Outlet</td>
</tr>
<tr>
<td><img src="image" alt="Prescription use only" /></td>
<td>Prescription use only</td>
</tr>
<tr>
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<td>Purge Location</td>
</tr>
<tr>
<td><img src="image" alt="Refer to Instructions" /></td>
<td>Refer to Instructions</td>
</tr>
<tr>
<td><img src="image" alt="Running on Battery" /></td>
<td>Running on Battery</td>
</tr>
<tr>
<td><img src="image" alt="Sample Gas Inlet Port" /></td>
<td>Sample Gas Inlet Port</td>
</tr>
<tr>
<td><img src="image" alt="Sample Gas Outlet Port" /></td>
<td>Sample Gas Outlet Port</td>
</tr>
<tr>
<td><img src="image" alt="Separate Collection" /></td>
<td>Separate Collection</td>
</tr>
<tr>
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<td>Serial Number</td>
</tr>
<tr>
<td><img src="image" alt="Standby" /></td>
<td>Standby</td>
</tr>
<tr>
<td><img src="image" alt="Stock Number" /></td>
<td>Stock Number</td>
</tr>
<tr>
<td><img src="image" alt="Type B Electrical Equipment" /></td>
<td>Type B Electrical Equipment</td>
</tr>
<tr>
<td><img src="image" alt="Use by yyyy-mm" /></td>
<td>Use by yyyy-mm</td>
</tr>
<tr>
<td><img src="image" alt="USB Port" /></td>
<td>USB Port</td>
</tr>
<tr>
<td><img src="image" alt="Water Separator Cartridge" /></td>
<td>Water Separator Cartridge</td>
</tr>
</tbody>
</table>
Theory of Operation

The INOmax DSIR Plus provides a constant dose of INOMAX into the inspiratory limb of the ventilator circuit. The INOmax DSIR Plus uses a “dual-channel” design to ensure the safe delivery of INOMAX. The first channel has the delivery CPU, the flow controller and the injector module to ensure the accurate delivery of NO. The second channel is the monitoring system, which includes a separate monitor CPU, the gas sensors (NO, NO₂, and O₂ sensors) and the user interface, including the display and alarms. The dual-channel approach to delivery and monitoring permits INOMAX delivery independent of monitoring. This allows the monitoring system to shutdown INOMAX delivery, if it detects a fault in the delivery system. For example, INOMAX delivery will be interrupted should the monitored NO concentration become greater than 100 ppm for greater than 12 consecutive seconds. (See Figure 1-15 for a schematic diagram).

1. INOMAX drug is stored as a gas mixture of NO/N₂ in an aluminum cylinder at a concentration of either 800 or 400 ppm.

2. The cylinder is attached to a high pressure regulator, which incorporates a pressure gauge that indicates cylinder pressure when the cylinder valve is open. The cylinder regulator is attached via tubing to the INOmax DSIR Plus using one of the two NO/N₂ quick connect inlets on the back of the device.

3. The INOmax DSIR Plus checks the INOMAX cylinder for the correct expiration date and cylinder concentration.

4. The INOMAX enters the back of the INOmax DSIR Plus, passes through a filter, then a safety shutoff valve, which is open under normal operation.

5. An injector module is placed in the ventilator gas flow between the ventilator inspiratory outlet and the humidifier. Based on the ventilator flow, the INOMAX cylinder concentration and set INOMAX dose, the proportional solenoid valve delivers 800 or 400 ppm INOMAX into the ventilator circuit via the injector module where it mixes with the breathing circuit gas flow to achieve the set dose. This allows the INOmax DSIR Plus to deliver a constant dose of INOMAX regardless of the ventilator flow pattern or breath rate (see Figure 1-16).

6. An internal flow sensor verifies the INOMAX flow from the proportioning valve, providing feedback to adjust the flow real time. This assures the calculated INOMAX flow necessary to achieve a given dose based on reported injector module flow. A one-way valve separates the flow sensor from potential reverse flow that may come from the ventilator circuit.

7. Gas Monitoring - The INOmax DSIR Plus gas monitoring system provides monitored values for inspired NO, NO₂, and O₂. The sample gas is withdrawn from the breathing circuit and goes through a water bottle, a zero valve, a sample pump and finally a sample flow sensor to the gas monitoring sensors.

7a. The zero valve allows the gas sensors to be zeroed (during low calibration) without having to disconnect the sample line from the breathing circuit.

7b. The pump and sample flow sensor ensure a sample gas flow rate is maintained to the monitoring sensors.

7c. The gas monitoring sensors are electrochemical; they are specific to each gas and provide an electronic signal which is proportional to the concentration of the gas present.

8. Integrated Pneumatic Backup - The INOmax DSIR Plus has an integrated pneumatic backup system that will supply a fixed flow of INOMAX at 250 mL per minute into the injector module. This system is completely pneumatic and does not rely on electronic control or power. The system will not allow a dose to be set on the INOmax DSIR Plus if the integrated pneumatic backup is in use.
Figure 1-15 Schematic Diagram of INOmax DSIR Plus

Figure 1-16 INOMAX injection method provides a constant NO concentration
Effect of the INOmax DS$_{IR}$ Plus in a ventilator circuit

There are two main effects of connecting and using the INOmax DS$_{IR}$ Plus in a ventilator breathing circuit.

1. The INOmax DS$_{IR}$ Plus adds NO/N$_2$ gas to the breathing circuit in proportion to the NO setting and the ventilator flowrate. For example, at an NO setting of 20 ppm with an 800 ppm NO cylinder, the INOmaxDS$_{IR}$ Plus adds 2.5% more gas to that delivered by the ventilator and proportionally less for lower NO settings.

2. The INOmax DS$_{IR}$ Plus subtracts gas from the breathing circuit via the gas sampling system at a nominal flow rate of 0.23 L/min.

These two effects of adding and subtracting gas from the ventilator breathing circuit have the following effects:

**Oxygen Dilution**

The INOmax DS$_{IR}$ Plus adds gas to the breathing circuit in proportion to the NO setting as described above. The NO/N$_2$ mixture added to the ventilator gas dilutes the oxygen in proportion to the set INOMAX dose. At the INOMAX dose setting of 20 ppm (800 ppm cylinder), the added gas is 2.5%. Thus, the O$_2$ concentration is reduced by 2.5% of its original value. For example, if the original O$_2$ concentration was 60% v/v, then the O$_2$ value after injection, at the maximum setting, is 58.5% v/v.

<table>
<thead>
<tr>
<th>Set Dose (ppm)</th>
<th>Oxygen Dilution % v/v</th>
</tr>
</thead>
<tbody>
<tr>
<td>800 ppm cylinder</td>
<td></td>
</tr>
<tr>
<td>80</td>
<td>10</td>
</tr>
<tr>
<td>40</td>
<td>5</td>
</tr>
<tr>
<td>20</td>
<td>2.5</td>
</tr>
</tbody>
</table>

**Minute Volume**

When using volume ventilation with the INOmax DS$_{IR}$ Plus, the measured tidal volume delivered to the patient shows small changes depending on the NO setting being used due to the addition and subtraction of gases by the delivery system. Some minor ventilator adjustments to the minute volume may be required. The net result of the INOmax DS$_{IR}$ Plus on the delivered minute ventilation can be calculated as follows:

If the patient’s minute ventilation is 10 L/min (500 cc X 20 breaths/min)

The additional minute volume due to the INOMAX can be calculated as follows:

\[
\text{Additional INOMAX volume added per minute} = \frac{\text{INOmax dose} \times \text{Minute Volume}}{\text{Cylinder Concentration} - \text{INOmax Dose}}
\]

For a dose of 20 ppm (800 ppm cylinder) the additional volume would be

\[
(20 \times 10 / 800 - 20) = 0.26 \text{ L/min}
\]

To calculate the net change in minute volume:

\[
0.26 \text{ L/min INOMAX added} - 0.23 \text{ L/min removed (sample system)} = 0.03 \text{ L/min (net change)}
\]

This formula may be used when calculating the changes to continuous flow on continuous flow ventilators as well (using the continuous flow in place of minute ventilation).

**Trigger Sensitivity**

The addition and subtraction of gases by the INOmax DS$_{IR}$ Plus may affect the trigger sensitivity of the ventilator when using synchronized modes of ventilation. This may cause the ventilator to auto-trigger in ventilators which have flow trigger modes, especially where the trigger flow is set to less than one L/min. The trigger sensitivity of the ventilator should be checked after connecting the INOmaxDS$_{IR}$ Plus delivery system.
Circle Anesthesia Ventilator Systems

The use of the INOmax DSIR Plus with circle anesthesia ventilator systems (which use volume ventilation) causes small changes in the delivered minute volume as noted previously (see Minute Volume, page 1-24). Recirculation of INOMAX in circle breathing systems should be avoided. The gas in the ventilator bellows may also contain undesirable levels of NO₂ which may not be removed by the CO₂ absorbent.

Recirculation of gases may lead to a rapid increase in INOMAX dose levels creating a shutdown of the INOmaxDSIR Plus. This can be avoided by using a fresh gas flow rate equal to or above that of the patient's minute volume. This will ensure that there is sufficient fresh gas in the absorber such that no accumulated gas from the ventilator bellows reaches the patient through the inspiratory limb of the breathing circuit.

Maximum NO Delivery

The INOmax DSIR Plus is limited to a maximum NO flow of 6.35 L/min. Maximum deliverable dose is 80 ppm (800 ppm cylinders) or 40 ppm (400 ppm cylinders) when the breathing gas flow is 60 lpm or less. Breathing gas flows greater than 60 lpm will reduce the delivered dose (resulting in a lower monitored NO value). See the graph below for estimated dosing based on breathing gas circuit flow rate.

When intermittent inspiratory flow rates are used, peak ventilator flows which exceed 120 L/min may be achieved. Peak inspiratory flow rates are transient and extremely short in duration. As a result, the portion of the breath which is not matched by the INOmax DSIR Plus is extremely small and the effect on the delivered concentration of NO within the entire range of the breath is small.

Does acid form in the humidifier or breathing circuit when delivering INOMAX?

A long term test was performed at Datex-Ohmeda to determine if acid would build up in a breathing circuit over time when delivering inhaled Nitric Oxide.

The test equipment was a Sechrist IV-100B neonatal ventilator and a Fisher Paykel MR500 humidifier. The ventilator settings were Rate 0 breaths per minute, Flow 6 L/min and Oxygen 100% v/v and the humidifier was set to 36 degree’s C.

The pH level was measured at the humidifier (the water in the humidifier chamber), at the patient Y (the condensate in the breathing circuit) and at the exhalation valve back at the ventilator (the condensate in the breathing circuit).

For the test distilled water was used which had an initial pH of 5.75 and the pH was measured with Hydrion Paper (4.5 to 7.5).

A control test without NO being delivered was run initially to see if the pH would change over time due to the slightly acidic nature of distilled water. The control test was run for six days with no change in the pH at any of the test points. The test was then repeated with 80 ppm of NO being delivered continuously for nine days with the pH being tested daily at each of the test points. There was no change of pH at any of the test points for any of the daily tests.
Environmental Effects

The National Institute for Occupational Safety and Health (NIOSH) have recommended exposure limits as follows (Ref. 1).

<table>
<thead>
<tr>
<th>NO</th>
<th>time-weighted (8 hours) average concentration limit of 25 ppm</th>
</tr>
</thead>
<tbody>
<tr>
<td>NO₂</td>
<td>ceiling limit of 1 ppm</td>
</tr>
</tbody>
</table>

The environmental build up of NO in a well ventilated ICU room can be evaluated using the following calculation.

<table>
<thead>
<tr>
<th>Room size</th>
<th>1000 ft³</th>
</tr>
</thead>
<tbody>
<tr>
<td>Room volume</td>
<td>28,300 L</td>
</tr>
<tr>
<td>Room ventilation (6 complete exchanges/hour)</td>
<td>2,830 L/min</td>
</tr>
<tr>
<td>NO flow into the room</td>
<td>80 ppm at 14 L/min</td>
</tr>
<tr>
<td>Average NO room concentration</td>
<td>(80 x 14) ÷ 2,830 = 0.396 ppm (0.4 ppm)</td>
</tr>
</tbody>
</table>

This theoretic calculation can be supplemented by measurements as performed by Hess et al (Ref. 2). The NO and NO₂ concentrations were measured using a chemiluminescence analyzer when 100 ppm of NO at 8 L/min was delivered into a room with no scavenging being used. The maximum NO and NO₂ concentrations measured over a one hour period were 0.12 ppm of NO and 0.03 ppm of NO₂.

Both these methods show that the exposure levels are significantly less than the levels recommended by NIOSH.

If the location for using NO has uncertain ventilation then the location should be evaluated for NO and NO₂ build up prior to use.

References:

(Ref. 1) Centers for Disease Control, Atlanta, GA 30333 USA. NIOSH Recommendations for Occupational Safety and Health Standards 1988. August 26, 1988 / vol. 37 / No. 9.

(Ref. 2) Hess et al, Use of Inhaled Nitric Oxide in patients with Acute Respiratory Distress Syndrome. Respiratory Care, 1996, vol. 41, No. 5, pg. 424-446.
INOmax DS$^\text{IR}$ Plus

2/ Automated Pre-Use Checkout
INOmax DS$_{IR}$ Plus

2/ Automated Pre-Use Checkout
2/ Automated Pre-Use Checkout

Connect the INOmax DSIR Plus power cord to a hospital-grade AC outlet. The power cord must always be connected to an electrical outlet to maintain a full battery charge.

1. Turn ON INOmax DSIR Plus.

An INOmax DSIR Plus splash screen will appear once the device is turned ON followed by the speaker sounding.

Note:

- Low calibration automatically starts following the INOmax DSIR Plus self test.
- A Pre-Use wizard will be displayed on the main screen, which will provide step-by-step instructions to complete the pre-use procedure.
- Pressing the NEXT button initiates the Pre-Use wizard.
- Pressing the CANCEL button exits the Pre-Use wizard. If you cancel out of the pre-use wizard, the manual pre-use checkout procedure can be found in Section 9/Appendix.
Initial connections

1. Confirm the water bottle and water separator cartridge are in place \(1a\).
   Connect the patient gas sample line with Nafion to the sample line inlet port on the front of the INOmax DSIR Plus \(1b\).

2. Insert the injector module cable into the INOmax DSIR Plus and the injector module, lining up the red dots on both ends \(2a\).

3. Connect the injector tubing to the INOmax DSIR Plus and the injector module \(2b\).

Note:
- It is recommended to disinfect or sterilize the injector module prior to initial setup.
- To remove this type of connector, the knurled sleeve \(2c\) on the connector must be pulled outward before removing the connector from the injector module or the front panel.

Check cables and hoses for signs of wear and damage.
3. Verify the power supply indicator is illuminated (3a).

Caution: Keep the power cord off of the ground and away from moving parts.

4. Load two INOMAX drug cylinders onto cart and check for correct product identity labels, cylinder concentration (800 or 400 ppm) and expiration date.

Note:
- For the CGA-type INOMAX regulator connector, ensure the white plastic tip is not chipped or cracked. Remove and replace as necessary. (see Replacing the tip on the INOMAX regulator, Section 7/ Maintenance).
- For the ISO-type regulator connector, ensure that the O-ring is present and is not damaged (see Replacing the tip on the INOMAX regulator, Section 7/ Maintenance).

5. Connect an INOMAX regulator to one of the INOMAX cylinders, and hand tighten the fitting to the INOMAX cylinder.

Caution: If using the transport regulator/cap assembly (PN 10022, CGA or 10041, ISO) see Figure 4-9, Section 4/ Transport.
6. Connect the INOMAX regulator hose to one of the INOMAX inlets.
7. Connect the INOblender inlet hose to the INOmax DS_{IR} Plus INOblender outlet.
8. Slide the quick-connect cover into place.
9. Connect oxygen supply (3.4 bar or 50 psig) hose to O_2 inlet fitting on back of INOblender.
10. Connect the Infrared cable from the INOmax DS_{IR} Plus cart or transport regulator/cap assembly (PN 10022, CGA or 10041, ISO) to the back of the INOmax DS_{IR} Plus.

**Note:** Do not attempt to connect the transport regulator cap assembly electrical plug to the INOblender outlet port. This will damage the electrical pins on the connector plug.
High Pressure Leak Test and Automated Purge

**WARNING:** All INOmax DSIR Plus devices must complete an automated or manual purge procedure prior to the start of therapy, to ensure the patient does not receive an excess level of NO₂.

1. Verify one of the high pressure regulators is connected to an INOMAX cylinder.
2. Open and then close the cylinder valve. Verify cylinder has at least 34 bar (500 psig).
3. Monitor pressure gauge for 30 seconds for any signs of pressure decrease. If no pressure decrease is observed, the high pressure leak test is successful. If there is an observed pressure decrease, see Section 7/ Maintenance; Cylinder Leak Check.

1. Confirm injector module is out of the pre-use circuit. Press NEXT button to start purge process.

**Note:** Perform auto-purge with device plugged in. Failure to do so may result in the procedure not completing.
5. Low Cylinder Pressure alarm may activate following purge sequence.

6. Open cylinder valve when purge is completed.

Note: If low calibration is still running after the automated purge completes, wait for low calibration to complete.
Integrated Pneumatic Backup INOMAX Delivery Test

1. Assemble pre-use set-up connectors and tubing (press SHOW DIAGRAM button if needed).
   Set the oxygen flowmeter to 10 L/min. (#1 in Figure 2-1).

1. O₂ Flowmeter (Connected to wall/tank)
2. Injector Module Electrical Cable
3. NO/N₂ Injector Tube
4. Patient Gas Sample Line with Nafion
5. O₂ Tubing
6. 15M x 4.5 mm Adapter
7. 22M / 15F x 22M / 15F Adapter
8. Injector Module
9. 300 mm of 22 mm Hose
10. Gas Sample Tee

Figure 2-1
1. Using the pre-use set-up connectors, verify that the O₂ flowmeter is set to 10 L/min.

2. Press NEXT button to automatically set the INOMAX dose to 40 ppm for 800 ppm cylinders or 20 ppm for 400 ppm cylinders.

3. Allow monitored values to stabilize (may take up to 3 minutes). Verify the NO and NO₂ readings are within the following ranges:

<table>
<thead>
<tr>
<th>Cylinder Concentration (ppm)</th>
<th>800 ppm</th>
<th>400 ppm</th>
</tr>
</thead>
<tbody>
<tr>
<td>NO (ppm)</td>
<td>14 - 26</td>
<td>7 - 13</td>
</tr>
<tr>
<td>NO₂ (ppm)</td>
<td>≤ 1.0</td>
<td>≤ 1.0</td>
</tr>
</tbody>
</table>

4. Performance test is complete. Press NEXT button to set the INOMAX dose to zero.

**Performance Test**

<table>
<thead>
<tr>
<th>Cylinder Concentration (ppm)</th>
<th>800 ppm</th>
<th>400 ppm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Set Dose (ppm)</td>
<td>40</td>
<td>20</td>
</tr>
<tr>
<td>Acceptable NO Value (ppm)</td>
<td>35 - 45</td>
<td>17 - 23</td>
</tr>
<tr>
<td>Acceptable NO₂ Value (ppm)</td>
<td>&lt; 1.5</td>
<td>&lt; 1</td>
</tr>
<tr>
<td>Acceptable FiO₂ (%)</td>
<td>95 ± 3</td>
<td>95 ± 3</td>
</tr>
</tbody>
</table>

*Note:* If a monitored value is outside the range indicated, see Section 5/ Alarms and Troubleshooting.
**INOblender Test**

1. Remove oxygen tubing from O₂ flowmeter and connect to front of INOblender.
2. Remove the injector module from the pre-use set-up and reconnect the adapters.
3. On the INOblender, set the INOMAX dose and flow to:

<table>
<thead>
<tr>
<th>Cylinder Concentration (ppm)</th>
<th>800</th>
<th>400</th>
</tr>
</thead>
<tbody>
<tr>
<td>INOblender Set Dose (ppm)</td>
<td>40</td>
<td>20</td>
</tr>
<tr>
<td>INOblender Flow</td>
<td>10 L/min</td>
<td></td>
</tr>
</tbody>
</table>

4. Allow monitored values to stabilize (may take up to 3 minutes) and verify the NO value on the INOmax DSIR Plus using the table below:

5. Turn the dose and oxygen flow to zero.
6. Remove the pre-use set-up from the INOblender.

---

**Pre-use checkout complete.**

**WARNING:**

- If the INOmax DS<sub>IR</sub> Plus is not going to be used on a patient within 10 minutes, depressurize the regulator supply line (see next page "Depressurizing the Regulator Supply Line").
- If the INOmax DS<sub>IR</sub> Plus is not used and is pressurized for more than 10 minutes, repeat automated or manual purge procedure.
- If the INOmax DS<sub>IR</sub> Plus is depressurized and not used within 12 hours, repeat pre-use procedure.

The INOmax DS<sub>IR</sub> Plus is now ready to connect to the patient. Proceed to Section 3/ Patient Application.
2. At the back of the INOmax DSIR Plus, remove the regulator hose from the purge port and connect it to the INOMAX gas inlet. This depressurizes the regulator.

3. When the regulator pressure gauge reads zero, remove the regulator hose from the purge port and connect it to the INOMAX gas inlet.

**Note:** If difficulties are encountered in connecting the regulator hose, refer to page 3-9.
3/ Patient Application
INOmax DSIR® Plus

3/ Patient Application
3/ Patient Application

Before Operation

Complete the initial connections and Pre-Use Checkout procedure as described in the previous sections before connecting the INOmax DSIR Plus into the patient’s breathing circuit. (See the ventilator/breathing device manual for its setup and operation)

**WARNING:**

- The use of pediatric and neonatal ventilator settings with adult size breathing circuits can result in higher levels of NO₂. Always use the size of breathing circuit that is appropriate for the patient.
- Abrupt discontinuation of INOMAX may lead to worsening oxygenation and increasing pulmonary artery pressure, i.e., Rebound Pulmonary Hypertension Syndrome. To avoid abrupt discontinuation, utilize the INOblender or integrated pneumatic backup if necessary. If Rebound Pulmonary Hypertension occurs, reinstate INOMAX therapy immediately (See the INOMAX prescribing Information for further details.)
- If the high NO₂ alarm activates, the delivery system should be assessed for proper set up while maintaining INOMAX delivery. Adjust the dose as described in the INOMAX Prescribing Information-The Effects of Nitrogen Dioxide. If unable to determine the cause of the increased NO₂ levels, call Customer Support, do not discontinue therapy.

*Use Outside of Product Labeling*

- The INOmax DSIR Plus must only be used in accordance with the indications, usage, contraindications, warnings and precautions described in the INOMAX (nitric oxide) drug package inserts and labeling. Refer to this material prior to use.
- Helium/oxygen mixtures should not be used with the INOmax DSIR Plus.
- The use of devices which radiate high-intensity electrical fields may affect the operation of the INOmax DSIR Plus. Constant surveillance of all monitoring and life support equipment is mandatory whenever interfering devices are in operation on or near a patient.
- The approved patient population for the INOmax DSIR Plus, as specified in the drug labeling for INOMAX (nitric oxide) for inhalation, is limited to neonates. The INOmax DSIR Plus is not intended to be used in other patient populations.
Connection to the ventilator breathing circuit

**WARNING:** The INOmax DS$_{IR}$ Plus subtracts gas from the breathing circuit via the gas sampling system at 230 mL per minute which can cause the ventilator to auto-trigger. Adjusting the flow sensitivity may be necessary. The trigger sensitivity of the ventilator should be checked after connecting the INOmax DS$_{IR}$ Plus to the breathing circuit.

**Caution:**
- For proper gas flow measurement, the injector module should not be connected directly to the ventilator's inspiratory outlet.
- Connect the injector module directly to the humidifier inlet.
- If it is not possible to connect the injector module to the inlet of the humidifier or if a humidifier is not used, a length of tubing (at least 6 inches) must be placed between the ventilator patient outlet and the injector module.

**Note:** Connections to various ventilators are unique to each manufacturer as well as their corresponding disposable circuits. Adapter diagrams can be found on page 7-13.
Connect the INOmax DS$_{IR}$ Plus into the breathing circuit as shown in the appropriate connection diagrams later in this section.

1. To ensure correct flow measurement, use breathing circuit tubing between the ventilator inspiratory port and the injector module (Fig. 3-1).

2. Connect the injector module to the humidifier inlet, note the airflow direction indicator on the injector module (see Figure 3-2).

3. The distance between the injector module and the sample tee must be greater than 24 inches. This ensures proper gas mixing, minimizes the sampling of mixed inspired/expired concentrations, and ensures correct patient NO/NO$_2$ measurement.

4. Insert the sample tee on the inspiratory side of the ventilator circuit, 150-300 mm (6-12 inches) from the patient wye. Make sure that the sample tee port points upward. This helps to avoid fluid accumulation in the sample line.

5. Select the dose button on the screen. On the INOmax DS$_{IR}$ Plus, rotate the control wheel to set the NO dose.

6. Confirm the change by pressing the Control wheel or dose button on the screen.

7. Set the user-adjustable alarm settings on the INOmax DS$_{IR}$ Plus and on the ventilator or breathing system.

Note: The first time a dose is set from zero, the upper and lower NO alarm limits are set 50% above and 50% below the set dose.
INOblender Operation

Important: Read the INOblender Operation Manual PN 20181 before using the INOblender. Follow instructions and obey all Warnings and Cautions.

**WARNING:**

• The purge procedure must be followed to help ensure NO₂ is purged from the system before the manual resuscitator bag is connected to the patient.

• The manual resuscitator bag should be squeezed repeatedly during use to avoid NO₂ building up in the bag.

• If the bag is not squeezed repeatedly while delivering INOMAX, the bag should be removed from the patient and the bag purge procedure performed before continuing.

• The INOblender should be upright when setting the oxygen flowrate for accurate setting.

• Do not use pneumatically powered nebulizers with the INOblender. This will result in significant over delivery of INOMAX in excess of 80 parts per million with 800 ppm cylinders and 40 parts per million with 400 ppm cylinders.
  - The INOblender outlet pressure has been validated for use up to 400 millibar (5.8 psig) pressure. The amount of back-pressure generated by pneumatic nebulizers is significantly greater 1.4 to 2.0 bar (20-30 psig) and will result in over delivery of INOMAX in excess of 80 ppm. The user adjusted dose setting on the INOblender will not correlate with, or have an effect on the actual delivered dose.
  - In addition, the INOblender flowmeter is not back-pressure compensated and will display a lower flow rate than actual when pressure is applied to the outlet.

**Caution:**

• When not in use, the oxygen flowmeter should be turned off.

• A user may determine that some clinical conditions may necessitate the use of an oxygen/air blender with the INOblender to achieve FiO₂ levels less than 100%.

• Delivered INOMAX dose from the INOblender is affected by varying oxygen concentrations (see table below):

<table>
<thead>
<tr>
<th>FiO₂</th>
<th>INOblender Accuracy Specification (at 50 psig)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.0</td>
<td>+/- 20% of set value or 2 ppm whichever is greater</td>
</tr>
<tr>
<td>0.21 to 0.95</td>
<td>+/- 30% of set value or 3 ppm whichever is greater</td>
</tr>
</tbody>
</table>
Integrated Pneumatic Backup NO Delivery

**WARNING:**
- When using the integrated pneumatic backup with breathing circuit gas flows of 5 L/min, the delivered NO dose will be approximately 40 ppm (800 ppm cylinder) or 20 ppm (400 ppm cylinder). Breathing circuit gas flows less than 5 L/min will deliver an NO dose greater than 40 ppm (800 ppm cylinder) or 20 ppm (400 ppm cylinder).
- The integrated pneumatic backup is intended for short term use when the electronic delivery system fails until a replacement NO delivery device can be brought to the bedside.
- The integrated pneumatic backup delivers a variable concentration of NO to the patient depending on the ventilator flow being used.

**Integrated Pneumatic Backup NO Delivery Description**

The integrated pneumatic backup delivery provides a fixed flow of 250 ml/min of INOMAX directly into the ventilator circuit through the injector module. The integrated pneumatic backup is not reliant on the operation of the main system (see Figure 3-5).

- The integrated pneumatic backup delivery is activated through the backup switch on the front panel. When activated, the set INOMAX dose will be automatically turned OFF. The high and low NO alarms are automatically set to 90 and 5 ppm for 800 ppm cylinders (50 and 5 ppm for a 400 ppm cylinder) respectively (see Figure 3-3).
- The estimated backup dose graphic (if displayed) represents the estimated dose the patient is receiving, by displaying a dose indicator. The estimated backup dose is calculated by using the circuit flow measured by the injector module (see Figure 3-3).
- The estimated NO dose table is also displayed on the main screen.

**Note:**
- If the injector module is not functioning, the estimated backup dose graphic will be inactive.
- The estimated backup dose graphic and the estimated backup dose based on ventilator flow table, will not be present during a Cylinder Concentration Mismatch alarm.
When the backup switch is turned OFF, the dose and alarm settings will return to the previous values (see Figure 3-4).

**Figure 3-4**

**Figure 3-5**
Cylinder Information

**WARNING:**
- Only use manufacturer supplied drug cylinders, regulators and adapters.
- Cylinders should be stored between 15-30 degrees C.
- Always secure a cylinder when not using it.
- Never lift a cylinder by its valve.
- Never drop a cylinder.
- Never use a hammer, pry or wedge to loosen a valve or protection cap. The valve and protection cap should be operated by hand.
- Never let oil, grease or other combustibles come in contact with a cylinder or valve.
- Never remove or deface cylinder labeling or markings.
- Never attempt to repair a leak on a cylinder valve or its safety relief device.
- Never operate equipment that is leaking.
- Never ship a leaking cylinder.
- Never store cylinders:
  - Where damage can result from the elements, such as standing water or temperatures over 51 degrees C (125 degrees F).
  - Where they can contact corrosive substances.
  - Where they can be cut or abraded by an object.
  - Next to a walkway, elevator or platform edge.

**Purging the INOmax DS\textsubscript{IR} Plus**
- All INOmax DS\textsubscript{IR} Plus devices must be purged before use to ensure the patient does not receive an excess level of NO\textsubscript{2}.
- If the INOmax DS\textsubscript{IR} Plus is not going to be used on a patient within 10 minutes, depressurize the regulator supply line.
- If the INOmax DS\textsubscript{IR} Plus is not used and is pressurized for more than 10 minutes, repeat automated or manual purge procedure.
- If the INOmax DS\textsubscript{IR} Plus is depressurized and not used within 12 hours, repeat pre-use procedure.

**Note:**
- Use a properly designed cart to move a cylinder and properly secure the cylinder when moving it.
- Apply a proper pressure regulating device to the cylinder before using it.
- Periodically check the cylinder pressure.
- Apply the valve outlet cap and valve protective cap to a cylinder when it is not connected.
Changing INOMAX Cylinders and Purging the Regulator Assembly

**Caution:** Replace an INOMAX cylinder when its pressure is less than 14 bar (200 psig).

1. Check the INOMAX drug cylinders for the correct product identity, cylinder concentration, and expiration date. Verify cylinder has at least 34 bar (500 psig) and tighten the fitting to the I120A; cylinder, attach a second INOMax DSIR Plus regulator (hand-tighten only) which is currently not in use.

   - Do not attach the regulator hose to the INOMax DSIR Plus at this time.
   - For the CGA-type regulator connector, ensure the white plastic tip is in place on the regulator connector and not chipped or cracked. Remove and replace as necessary (see page 7-8).
   - For the ISO-type regulator connector, check that the O-ring is present and is not damaged (see Replacing the tip on the INOMAX regulator, page 7-9).

2. Open and then close the valve on the new INOMAX cylinder. Check for adequate cylinder pressure. Monitor pressure gauge for 30 seconds for any signs of leakage. If there is a decrease, check for leaks around the hose connections and cylinder valve connector using soapy water. (see Section 7/ Maintenance; Cylinder Leak Check).
3. Insert the NO/N₂ quick-connect fitting into the purge port on the back of the INOmax DSIR Plus and firmly push until the regulator pressure gauge reads zero (this purges any NO₂ that has accumulated in the hose and regulator).

**WARNING:** All INOmax DSIR Plus devices must be purged before use to ensure the patient does not receive an excess level of NO₂.

4. Prior to connecting a regulator hose, ensure the inlet connectors, on the INOmax DSIR Plus unit, have the knurled sleeve set in the back position (toward the INOmax DSIR Plus unit, see Figure 3-6).
5. Open the cylinder valve on the new cylinder (this may activate the “Two Cylinders Open” alarm until the empty cylinder valve is closed).

6. Close the cylinder valve on the empty cylinder and remove the supply line from the back of the INOmax DSIR Plus.

7. Depressurize by using the purge port on the back of the INOmax DSIR Plus prior to removing the regulator from the empty cylinder.

8. Replace the empty cylinder with a full cylinder on the cart.
Oxygen Dilution Chart

For delivery using *800 ppm or **400 ppm cylinder (Illustrative Only)

---

### Cylinder Conc. | 800  | 400
--- | --- | ---
5 2.5 | 0.21 | 0.40
10 5 | 0.21 | 0.40
20 10 | ![Warning] 0.20 | 0.39
40 20 | ![Warning] 0.20 | 0.38
80 40 | ![Warning] 0.19 | 0.036

### Set FiO2

<table>
<thead>
<tr>
<th>0.21</th>
<th>0.40</th>
<th>0.60</th>
<th>0.80</th>
<th>1.00</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.21</td>
<td>0.40</td>
<td>0.60</td>
<td>0.80</td>
<td>0.99</td>
</tr>
<tr>
<td>0.21</td>
<td>0.40</td>
<td>0.59</td>
<td>0.79</td>
<td>0.99</td>
</tr>
<tr>
<td>![Warning] 0.20</td>
<td>0.39</td>
<td>0.59</td>
<td>0.78</td>
<td>0.98</td>
</tr>
<tr>
<td>![Warning] 0.20</td>
<td>0.38</td>
<td>0.57</td>
<td>0.76</td>
<td>0.95</td>
</tr>
<tr>
<td>![Warning] 0.19</td>
<td>0.036</td>
<td>0.54</td>
<td>0.72</td>
<td>0.90</td>
</tr>
</tbody>
</table>

### Actual FiO2

---

⚠️ Caution FiO₂ less than 21%

Please Note:
The calculations on this chart have been determined based on 800 and 400 ppm cylinders of INOMAX (nitric oxide) for inhalation.
This chart is representative of a range of doses available on the INOmax DSᵢᵣ Plus. Doses higher than 20 ppm as a therapeutic dose are not recommended.
Calculations are considered estimates and may vary under clinical conditions.
All numbers have been rounded to the nearest hundredth.
## Duration Chart

### INOMAX Cylinder Luxfer 10L-Size

**For a Luxfer-Size *800 ppm and **400 ppm Cylinder Concentrations*** *(Illustrative Only)*

<table>
<thead>
<tr>
<th>INOMAX Dose (ppm)</th>
<th>Cylinder Conc.</th>
<th>*800</th>
<th>**400</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>5 L/min</td>
<td>10 L/min</td>
</tr>
<tr>
<td>5</td>
<td>2.5</td>
<td>30.7 Days</td>
<td>15.4 Days</td>
</tr>
<tr>
<td>10</td>
<td>5</td>
<td>15.3 Days</td>
<td>7.6 Days</td>
</tr>
<tr>
<td>20</td>
<td>10</td>
<td>7.5 Days</td>
<td>3.8 Days</td>
</tr>
<tr>
<td>40</td>
<td>20</td>
<td>3.7 Days</td>
<td>1.8 Days</td>
</tr>
<tr>
<td>80</td>
<td>40</td>
<td>1.7 Days</td>
<td>20.9 Hours</td>
</tr>
</tbody>
</table>

This chart is representative of a range of doses available on the INOMax DSIR Plus. Doses higher than 20 ppm as a therapeutic dose are not recommended.

***All calculations for the table above are based on a full cylinder of 155 bar (2248 psig) Luxfer 10 Liter cylinder, with a cylinder change at 14 bar (200 psig). The figures are calculated based on a total continuous breathing circuit gas flow and a cylinder conversion factor of 10 liters per bar (0.69 liters per psig).

- INOMAX flow = [Desired dose × total ventilator flow] + [Cylinder concentration - desired dose]
- Cylinder volume = Cylinder conversion factor × cylinder pressure (bar/psig)
- Cylinder duration (hours) = (Cylinder volume ÷ INOMAX flow rate) ÷ 60

Calculations are considered estimates and may vary under clinical circumstances.
**Duration Chart**

**INO MAX Cylinder Luxfer 2L-Size**

**For a Luxfer-Size *800 ppm and **400 ppm Cylinder Concentrations*** *(Illustrative Only)*

<table>
<thead>
<tr>
<th>INOMAX Dose (ppm)</th>
<th>Cylinder Conc.</th>
<th>FLOW 5 L/min</th>
<th>FLOW 10 L/min</th>
<th>FLOW 20 L/min</th>
<th>FLOW 40 L/min</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>*800<strong>400</strong></td>
<td>5 L/min</td>
<td>10 L/min</td>
<td>20 L/min</td>
<td>40 L/min</td>
</tr>
<tr>
<td>5</td>
<td>2.5</td>
<td>6.1 Days</td>
<td>3.1 Days</td>
<td>1.5 Days</td>
<td>18.4 Hours</td>
</tr>
<tr>
<td>10</td>
<td>5</td>
<td>3.1 Days</td>
<td>1.5 Days</td>
<td>18.3 Hours</td>
<td>9.2 Hours</td>
</tr>
<tr>
<td>20</td>
<td>10</td>
<td>1.5 Days</td>
<td>18.1 Hours</td>
<td>9.0 Hours</td>
<td>4.5 Hours</td>
</tr>
<tr>
<td>40</td>
<td>20</td>
<td>17.6 Hours</td>
<td>8.8 Hours</td>
<td>4.4 Hours</td>
<td>2.2 Hours</td>
</tr>
<tr>
<td>80</td>
<td>40</td>
<td>8.3 Hours</td>
<td>4.2 Hours</td>
<td>2.1 Hours</td>
<td>1.0 Hours</td>
</tr>
</tbody>
</table>

This chart is representative of a range of doses available on the INOmax DSIR Plus. Doses higher than 20 ppm as a therapeutic dose are not recommended.

*** All calculations for the table above are based on a full cylinder of 155 bar (2248 psig), Luxfer 2 Liter cylinder, with a cylinder change at 14 bar (200 psig). The figures are calculated based on a total continuous breathing circuit gas flow and a cylinder conversion factor of 2.0 liters per bar (0.14 liters per psig).

- INOMAX flow = [Desired dose × total ventilator flow] ÷ [Cylinder concentration - desired dose]
- Cylinder volume = Cylinder conversion factor × cylinder pressure (bar/psig)
- Cylinder duration (hours) = (Cylinder volume ÷ INOMAX flow rate) × 60

Calculations are considered estimates and may vary under clinical circumstances.
Duration Chart
INOMAX Cylinder 88-Size

For a 88-Size *800 ppm and **400 ppm Cylinder Concentrations
***(Illustrative Only)

<table>
<thead>
<tr>
<th>Cylinder Conc. (ppm)</th>
<th>*800</th>
<th>**400</th>
<th>FLOW</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>5</td>
<td>2.5</td>
<td>5 L/min</td>
</tr>
<tr>
<td></td>
<td>10</td>
<td>5</td>
<td>10 L/min</td>
</tr>
<tr>
<td></td>
<td>20</td>
<td>10</td>
<td>20 L/min</td>
</tr>
<tr>
<td></td>
<td>40</td>
<td>20</td>
<td>40 L/min</td>
</tr>
<tr>
<td></td>
<td>80</td>
<td>40</td>
<td></td>
</tr>
</tbody>
</table>

This chart is representative of a range of doses available on the INOmax DSiR Plus. Doses higher than 20 ppm as a therapeutic dose are not recommended.

*** All calculations in the table above are based on a full cylinder, 128 bar (1850 psig), 1963 liter “88” cylinder, with a cylinder change at 14 bar (200 psig). The figures are calculated based on a total continuous breathing circuit gas flow and a cylinder conversion factor of 15.7 liters per bar (1.08 liters per psig).

- INOMAX flow = [Desired dose × total ventilator flow] ÷ [Cylinder concentration - desired dose]
- Cylinder volume = Cylinder conversion factor × cylinder pressure (bar/psig)
- Cylinder duration (hours) = (Cylinder volume ÷ INOMAX flow rate) ÷ 60

Calculations are considered estimates and may vary under clinical circumstances.
Duration Chart
INOMAX Cylinder D-Size

For a D-Size *800 ppm and **400 ppm Cylinder Concentrations
*** (Illustrative Only)

<table>
<thead>
<tr>
<th>Cylinder Conc. (ppm)</th>
<th>*800</th>
<th>**400</th>
<th>5 L/min</th>
<th>10 L/min</th>
<th>20 L/min</th>
<th>40 L/min</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>2.5</td>
<td>7.0 Days</td>
<td>3.5 Days</td>
<td>1.8 Days</td>
<td>21 Hours</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>5</td>
<td>3.5 Days</td>
<td>1.7 Days</td>
<td>21 Hours</td>
<td>10.5 Hours</td>
<td></td>
</tr>
<tr>
<td>20</td>
<td>10</td>
<td>1.7 Days</td>
<td>20.7 Hours</td>
<td>10.3 Hours</td>
<td>5.2 Hours</td>
<td></td>
</tr>
<tr>
<td>40</td>
<td>20</td>
<td>20 Hours</td>
<td>10 Hours</td>
<td>5 Hours</td>
<td>2.5 Hours</td>
<td></td>
</tr>
<tr>
<td>80</td>
<td>40</td>
<td>9.5 Hours</td>
<td>4.8 Hours</td>
<td>2.4 Hours</td>
<td>1.2 Hours</td>
<td></td>
</tr>
</tbody>
</table>

Typically used in transport

This chart is representative of a range of doses available on the INOmax DSIR Plus. Doses higher than 20 ppm as a therapeutic dose are not recommended.

*** All calculations in the table above are based on a full cylinder, 128 bar (1850 psig), “D” cylinder, with a cylinder change at 14 bar (200 psig). The figures are calculated based on a total continuous breathing circuit gas flow using a cylinder conversion factor of 2.8 liters per bar (0.19 liters per psig).

- INOMAX flow = [Desired dose × total ventilator flow] ÷ [Cylinder concentration - desired dose]
- Cylinder volume = Cylinder conversion factor × cylinder pressure (bar/psig)
- Cylinder duration (hours) = (Cylinder volume + INOMAX flow rate) ÷ 60

Calculations are considered estimates and may vary under clinical circumstances.
Emptying the Water Bottle

The water bottle (see Figure 3-7) collects fluids separated from the patient gas sample.

- Empty and clean the water bottle before each patient use and empty whenever the bottle is more than half full.
- Empty the water bottle routinely. Allowing it to fill and overflow may cause system errors.
- A “Water Bottle Full” message will remind the user to empty and clean the bottle should it become full.

Note: Monitoring will be temporarily interrupted when the “Water Bottle Full” message is indicated.

To empty the Water Bottle:

1. Remove the bottle by pulling it straight down (see Figure 3-7).
2. Discard the contents according to an approved fluid waste disposal policy.
3. Clean the bottle.
4. Replace the bottle by pushing it up into position.
5. Check for leaks by running the system and occluding the sample line until the “Sample Line/Filter Block” alarm message appears.

Figure 3-7

During delivery of INOMAX to a patient

1. The disposable water separator cartridge on the rear of the water bottle housing protects the monitoring system from moisture and other contaminants and may need to be replaced occasionally while in use (refer to Section 7/ Maintenance).
2. To avoid medications interfering with the gas monitoring system, administer any aerosolized medications distal to the sampling tee in the breathing circuit (refer to page 3-18).
When operating on the battery, a battery icon \(1\) is displayed on the screen along with the message “Running on Battery” \(2\) in the text message area.

- The low battery alarm will alert the user when there are approximately 30 minutes remaining.
- A fully charged battery will run the INOmax DS\(_{IR}\) Plus for up to six hours in optimal conditions.
- Battery life can be extended by keeping the display brightness and the audio alarm volume to the minimum. (Display brightness and alarm volume can be changed by accessing the settings screen. See section 1/ General Information for instructions).
- Connect the system to an electrical outlet for at least ten hours to fully charge the battery when depleted.
Inspired Gas Sampling During Aerosol Delivery

Caution: Pneumatic nebulizers will dilute the delivered INOMAX dose.

To sample inspired gas during aerosol delivery:
- Place the medication nebulizer downstream of the sample tee on the inspiratory limb (see Figure 3-8).

This avoids over saturation of the water separator cartridge, or contamination of the sample system. In addition, it prevents the Sample Line/Filter Block alarm from occurring.

1. Place the 1.0 micron disk filter on the INOmax DS IR Plus sample line inlet (Fig. 3-9).
2. Connect the patient gas sample line to the filter. If monitored values change, check that all sample line and filter connections are secure.

This disk filter has been validated for this purpose.

Note:
- Change the filter every 12 hours or more often if necessary (due to Sample Line/Filter Block alarm). Replacing the disk filter reduces the replacement frequency of the water separator cartridge.
- Do not operate the INOmax DS IR Plus without the water separator cartridge. Always use the disk filter in conjunction with the INOmax DS IR Plus water separator cartridge.
The INOmax DSIR Plus monitoring system can measure the environmental levels of NO and NO₂.

1. Disconnect the sample line connector from the sample tee.
2. Cap the Luer fitting on the sample tee.

**WARNING:** The patient gas sample tee must have the INOmax DSIR Plus sample line attached or be capped off to avoid loss of ventilator circuit pressure.

3. Sample the room air with the sample line and read the NO and NO₂ readings.
4. After environmental monitoring, remove the Luer fitting cap on the sample tee and reconnect the sample line.

**Note:** Monitoring alarms may occur during the performance of this test.
Entering Patient Information

The following are instructions of how to use the patient identifier screen.

**Note:**
- Any identifier entered will be linked with each INOMAX cylinder used during treatment.

A patient identifier and patient details can be entered at any time during the treatment of a patient by pressing the patient information button in the right-lower corner of the main screen.

**Note:** If patient identifier has not been entered a “Patient Info Incomplete” indicator will stay illuminated in the text message area of the screen, unless an alarm condition is present. (see Figure 3-10).

After pressing the patient information button, the patient information screen will appear (see Figure 3-11).

Press the "Enter Patient Identifier" button to access keyboard.

The patient identifier screen (see Figure 3-12) allows a unique alphanumeric patient identifier to be entered that contains six to eight characters (note: spaces will be accepted).

**Note:** For compliance to European Directive 95/46/EC, do not use identifiers traceable to a specific patient. Consult and comply with your internal hospital guidelines when entering a patient identifier.
Pressing the keys on the keyboard allows the user to enter a sequential alphanumerical identifier.

Prior to confirming the identifier, digits can be changed either by pressing the backspace button 1 or pressing the digit that has been entered and typing over it.

The CONFIRM button 2 will illuminate when six characters have been entered.

**Note:** Once the CONFIRM button has been pressed, the identifier remains unchangeable until therapy is ended by turning the device to Standby (OFF).

Press the Select Patient Type button and rotate the control wheel to select either neonate, pediatric or adult. Press the control wheel or button to confirm.

Once the patient type is confirmed the Select Diagnosis button will appear (see Figure 3-14).
Press the Select Diagnosis button and rotate the control wheel to select the patient diagnosis. Press the control wheel or button to confirm.

Figure 3-14

Press the CONFIRM button to enter the patient details selected (see Figure 3-15).

Figure 3-15

If the user has entered Neonate as the patient type, a screen will appear to enter Gestational Age at Birth and Weight at Birth (see Figure 3-16).

Prior to confirming the gestational age and birth weight, digits can be changed either by pressing the backspace button 1 or pressing the number that has been entered and selecting a new number.

The CONFIRM button 2 will illuminate when age and weight have been entered.
Once the CONFIRM button has been pressed, the patient details are stored (see Figure 3-17) and the identifier remains unchangeable until therapy is ended by turning the device to Standby (OFF).

To access patient identifier information, press the patient information button on the main screen.

Press the EXIT button to return to the main screen.
Connection to Various Breathing Systems

Acutronic Medical Systems AG Fabian +nCPAP Evolution

Note: Validated for use outside of the United States.

1. Fabian+ nCPAP Evolution
2. Patient Gas Sample Line with Nafion
3. Injector Module Electrical Cable
4. INOmax DSIR Plus
5. NO/N₂ Injector Tube
6. Connecting Tube (15 inches)
7. Injector Module
8. 22F X 15M Adapter
9. Humidifier
10. Inspiratory Breathing Circuit Hose
11. Gas Sample Tee
12. Patient Wye
13. Proximal Pressure Tube
14. Expiratory Breathing Circuit Hose

Figure 3-18 Example: Acutronic Medical Systems AG Fabian +nCPAP Circuit Diagram
(for conventional modes ONLY)
WARNING: Do not use dose settings above 40 ppm while using the HFO option. Bidirectional flow through the injector module may cause over-delivery which can lead to measured NO values greater than 100 ppm.

Note:
- Only use the circuit configuration below.
- Connect the injector module directly to the inspiratory outflow port on the front of the ventilator to prevent over-delivery from bidirectional flow during HFOV (gas flow from connection #7).
- Use of a one-way valve distal to the injector module IS NOT necessary.
- Validated for use outside of the United States.

Figure 3-19 Example: Acutronic Medical Systems AG Fabian HFO Circuit Diagram (for high frequency mode ONLY)
A-Plus Medical Babi-Plus Bubble CPAP Circuit

1. Oxygen Source
2. Oxygen Tubing
3. Pressure Relief Manifold
4. Injector Module
5. Temperature Probe
6. 90 Degree Sample Port Adapter
7. Nasal Prongs
8. Babi-Plus Bubble PAP Valve
9. Tee Adapter
10. Breathing Circuit
11. Humidifier
12. NO/N₂ Injector Tube
13. Injector Module Electrical Cable
14. INOmax DSIR Plus
15. Patient Gas Sample Line with Nafion

Figure 3-20 Example: A-Plus Medical Babi-Plus Bubble CPAP Circuit Diagram
Bagging Systems While Using the Injector Module

**WARNING:**

- The hyperinflation bag will, under some conditions, contain NO₂ in excess of one ppm. Use of large tidal volume breaths may expose the patients to the NO₂ present in the bag for part of the breath. In general, if the inspiratory flow rate induced by the manual ventilation does not exceed the fresh gas flow rate, the patient should not be exposed to the concentrations of NO₂ present in the hyperinflation bag.

- Adult and infant hyperinflation bags generate more NO₂ when used at lower minute ventilation. If use of the bag is interrupted (for example to adjust the tracheal tube), before resuming ventilation of the patient, the user should squeeze the bag several times to empty residual gas from the bag.

- Because of the potential for inhalation of excessive concentrations of NO₂, and the difficulty in monitoring the peak inhaled NO₂ concentrations, ventilation with a hyperinflation bag or self inflating bag is intended only for short term use.

- The monitoring system within the INOmax DSIR Plus will not detect generation of NO₂ within the hyperinflation bag or self-inflating bag devices and the alarms for excessive NO₂ cannot warn of NO₂ produced within the manual bag system.

To minimize the delivered concentration of NO₂, the following steps should be taken for use with the manual resuscitator bags:

- Use the smallest bag adequate to deliver the desired tidal volume.
- Oxygen tubing lengths greater than 182 cm (72 inches) should not be used (between the injector module and the bag).
- Use the highest fresh gas flow rated (up to 15 L/min) that is practical.
- Use the lowest practical inspired oxygen concentration.
- After starting fresh gas flow, squeeze the bag several times to empty residual gas in the bag prior to using the system to ventilate a patient.
Bagging Systems While Using the Injector Module

Caution: New O₂ tubing must be used each time for optimal fit on the 4.5 mm adapter.

![Diagram of bagging system connection]

- **1. O₂ Flowmeter (wall outlet or cylinder)**
- **2. O₂ Tubing**
- **3. 15M X 4.5 mm Adapter**
- **4. 22M/15F X 22M/15F Adapter**
- **5. Injector Module**
- **6. 15M X 4.5 mm Adapter**
- **7. O₂ Tubing**
- **8. O₂ Tubing Sample Tee**
- **9. Patient Gas Sample Line with Nafion**
- **10. NO/N₂ Injector Tube**
- **11. Resuscitator Bag with O₂ Reservoir**
- **12. Injector Module Electrical Cable**

Figure 3-21 Example: Self-inflating Manual Bagging System Connection Diagram

Testing has been conducted using the following hyperinflation and self-inflating bag systems:
- Hudson RCI Hyperinflation 1L Adult # 5404
- Hudson RCI Hyperinflation 0.5L Neonatal # 5403
- Nellcor-Puritan Bennett Self-inflating 1.76 L Adult # 655005
- Nellcor-Puritan Bennett Self-inflating 0.52 L Infant # 616416
To minimize the delivered concentration of NO₂, the following steps should be taken for use with the manual resuscitator bags:

• Use the smallest bag adequate to deliver the desired tidal volume.
• Oxygen tubing lengths greater than 182 cm (72 inches) should not be used (between the injector module and the bag).
• Use the highest fresh gas flow rated (up to 15 L/min) that is practical.
• Use the lowest practical inspired oxygen concentration.
• After starting fresh gas flow, squeeze the bag several times to empty residual gas in the bag prior to using the system to ventilate a patient.

Figure 3-22 Example: Hyperinflation Manual Bagging System Diagram

Testing has been conducted using the following hyperinflation and self-inflating bag systems.

• Hudson RCI Hyperinflation 1 L Adult # 5404
• Hudson RCI Hyperinflation 0.5 L Neonatal # 5403
• Nellcor-Puritan Bennett Self-inflating 1.76 L Adult # 655005
• Nellcor-Puritan Bennett Self-inflating 0.52 L Infant # 616416
Bunnell Life Pulse High Frequency Ventilator Circuit

WARNING:

• The integrated pneumatic backup (250 mL/min.) should not be used with the Bunnell Life Pulse as ventilator flow rates can be very low at times, creating a potential delivered NO dose greater than 80 ppm.

• Place the Bunnell Life Pulse in standby prior to suctioning the patient to avoid NO delivery transiently exceeding the set dose by up to 30 ppm for 800 ppm cylinders. Press ENTER to reestablish ventilation as soon as the catheter is removed from the airway. This will limit the extent of over delivery above the NO set dose.

Caution:

• If set dose is below 5 ppm for 800 ppm cylinders and the Servo pressure is 140 mbar (2.0 psig) or less, this will result in flow rates outside of the specification of the injector module and fluctuating NO values may result.

• A one-way valve should be placed between the injector module and the humidifier chamber to prevent water from backing up into the injector module if the Life Pulse is either put into Standby or cycled OFF.

• There are higher pressures in the breathing circuit than normal; use only parts provided in disposable package #50046 and tightly secure all connections.

Connection Instructions:

1. Connect the sample Tee as shown in Figure 3-24.
2. Connect the injector module as shown in Figure 3-25. The one-way valve prevents water from backing up into the injector module if the Life Pulse is either put into standby or cycled OFF.
Connecting INOmax DS_{IR} Plus Sample Tee to the Bunnell Life Pulse Circuit

1. From Patient Box
2. Cut Green tube at midpoint (approximately six inches from the Life Port Adapter)
3. Patient Gas Sample Line with Nafion
4. Insert Sample Tee
5. Life Port Adapter
6. Endotracheal Tube

Figure 3-24

Connecting INOmax DS_{IR} Plus Injector Module to the Bunnell Life Pulse Circuit

1. Injector Module Electrical Cable
2. NO/N_{2} Injector Tube
3. Gas Out Tube from Vent
4. 15M X 4.5 mm I.D. Adapter
5. 22M/15F X 22M/15F Adapter
6. Injector Module
7. 15M X 4.5 mm I.D. Adapter
8. Three cm Piece of Green Gas Out Tube
9. One-Way Valve
10. Green Gas Out Tube to Humidifier

Figure 3-25
Figure 3-26 Example: CareFusion Infant Flow CPAP System Circuit Diagram

1. INOmax DSIR Plus
2. Heated Delivery Circuit
3. Infant Flow System
4. Infant Flow Generator
5. Sample Tee
6. Temperature Probe
7. Patient Gas Sample Line with Nafion
8. Humidifier
9. 22F X 15M Adapter
10. Injector Module
11. NO/N₂ Injector Tube
12. Injector Module Electrical Cable
CareFusion Infant Flow SiPAP

- The INOmax DSIR Plus adds NO/N₂ gas flow to the breathing circuit flow in proportion to the NO setting (up to 10% at 80 ppm for a 800 ppm cylinder and 40 ppm for a 400 ppm cylinder) and subtracts gas from the breathing circuit via gas sampling at a nominal flow rate of 0.23 L/min.

- These effects change the flow going to the nasal adapter and can therefore impact the CPAP level established by specific flow settings (See table below). The maximum flow error is approximately 11% at two L/min which is within the accuracy of the flow meter specification (+/-15%).

- It is recommended that after an NO setting change the user checks the CPAP level on the Infant Flow SiPAP front panel display and adjusts as necessary.

<table>
<thead>
<tr>
<th>SiPAP Flow Setting</th>
<th>Flow (L/min)</th>
<th>Flow (L/min)</th>
<th>Flow (L/min)</th>
<th>Flow (L/min)</th>
<th>Flow (L/min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>After INOmax DSIR Plus Set at 0 ppm</td>
<td>1.77</td>
<td>3.77</td>
<td>5.77</td>
<td>7.77</td>
<td>9.77</td>
</tr>
<tr>
<td>Actual Flow vs Set</td>
<td>-11.5%</td>
<td>-5.8%</td>
<td>-3.8%</td>
<td>-2.9%</td>
<td>-2.3%</td>
</tr>
<tr>
<td>After INOmax DSIR Plus Set at 80/800 ppm cylinder or 40/400 ppm cylinder</td>
<td>1.97</td>
<td>4.17</td>
<td>6.37</td>
<td>8.57</td>
<td>10.77</td>
</tr>
<tr>
<td>Actual Flow vs Set</td>
<td>-1.5%</td>
<td>4.3%</td>
<td>6.2%</td>
<td>7.1%</td>
<td>7.7%</td>
</tr>
</tbody>
</table>
CareFusion Infant Flow SiPAP

Figure 3-27 Example: CareFusion Infant Flow SiPAP Circuit Diagram
Circle Anesthesia System

**WARNING:**

- Avoid recirculation of gases. Undesired recirculation of gases will occur if fresh gas flows are less than the patient minute volume and may result in:
  - Higher NO₂ levels due to the limited ability of the carbon dioxide absorbent to remove NO₂.
  - Higher NO concentrations than those set due to NO recirculated through the absorber.
  - Reduction in O₂ concentration because nitrogen is the balance gas for nitric oxide and will be present in the re-circulated gases.
- If the injector module has been used in the wet/humidified part of the breathing circuit, it should be sterilized between each patient use.

**Caution:**

- Note the airflow direction arrow on the injector module: the flow out of the absorber must pass through the injector module in the direction of the arrow on the module.
- Nitrous Oxide (N₂O) will also affect the Set NO versus the measured NO value. For a 50% N₂O, 50% O₂ composition, the measured NO value will be approximately 7% less than the same Set NO value at 100% O₂. For example, at a Set NO value of 20 ppm, measured NO will be approximately 18 ppm.
- Similarly, the effect of two percent v/v Isoflurane will result in a high measured NO value of approximately three percent indicated for the same Set NO value at 100% O₂.
- Sudden changes in anesthetic agent concentration may cause brief transient changes in the measured NO and NO₂ values.

**Note:**

- With a circle anesthesia breathing circuit, the INOmax DSIR Plus will perform as specified in the technical specifications with fresh gas flow rates equal to or more than the patient minute volume.
- The breathing circuit between the sample tee and the patient should be between 150-300 mm (6-12 in.) long: greater than 150 mm (6 inches) to minimize the sampling of mixed inspired/ expired concentrations and less than 300 mm (12 inches) to help ensure correct patient NO₂ measurement.
- For OR ventilation systems with the inspiratory flow measurements at the inspiratory port of the absorber, place the injector module upstream of the inspiratory flow sensor.
Circle Anesthesia System

Figure 3-28 Example: Anesthesia System with Ventilator Circuit Diagram

1. Patient Gas Sample Line with Nafion
2. Patient Gas Sample Line Input Connection
3. INOmax DSFR Plus
4. Bellows Assembly
5. Ventilator
6. Ventilator Drive Gas
7. Absorber Expiratory Port
8. Absorber Inspiratory Port
9. Absorber
10. Injector Module
   a. Injector Module Input End
   b. Injector Module Output End
11. Inspiratory Tubing
12. 22M/15F X 22M/15F Adapter
13. Gas Sample Tee
14. Patient Wye
Dräger Babylog VN500/Infinity Acute Care System and Heinen & Löwenstein Leoni-plus Ventilator

**WARNING:** Always use a one-way valve to avoid high NO delivery.

**Note:** Validated for use outside of the United States.

1. Patient wye
2. Dräger Babylog VN500 / Leoni-plus Ventilator
3. Ventilator Expiratory Port
4. Ventilator Inspiratory Port
5. Patient Gas Sample Line Input Connection
6. INOmax DSIR Plus
7. NO/N₂ Injector Tube Front Panel Connection
8. Injector Module Electrical Cable Front Panel Connection
9. Injector Module
10. One-Way Valve
11. Humidifier Inlet
12. Humidifier
13. Humidifier Outlet
14. Patient Gas Sample Line with Nafion
15. Gas Sample Tee

Figure 3-29 Example: Dräger Babylog VN500 and Leoni-plus Circuit Diagram
Fisher & Paykel Healthcare Bubble CPAP

1. Oxygen Source
2. Oxygen Tubing
3. Bubble CPAP Pressure Manifold
4. 22F X 15M Adapter
5. 22M/15F X 22M/15F Adapter
6. Injector Module
7. Temperature Probe
8. Nasal Prong Infant Interface
9. Bubble CPAP Generator
10. F/P Inline Infant Nebulizer Kit (RT010) Adapter
11. Breathing Circuit
12. Humidifier
13. NO/N₂ Injector Tube
14. Injector Module Electrical Cable
15. INOmax DSIR Plus
16. Patient Gas Sample Line with Nafion

Figure 3-30 Example: Fisher & Paykel Healthcare Bubble CPAP System Circuit Diagram
Fisher & Paykel Healthcare Infant Circuit Nasal Cannula

1. Patient Gas Sample Line with Nafion
2. INOmax DS90 Plus
3. Oxygen Source
4. Oxygen Tubing
5. 22F X 15M Adapter
6. Injector Module
7. Pressure Relief Manifold
8. 22M/15F X 22M/15F Adapter
9. Injector Module Electrical Cable
10. NO/N2 Injector Tube
11. Humidifier
12. Breathing Circuit
13. Temperature Probe
14. Gas Sample Tee
15. Nasal Cannula

Figure 3-31 Example: Fisher & Paykel Healthcare Infant Circuit Nasal Cannula Diagram
Fisher & Paykel Healthcare Optiflow Breathing Circuit

1. Patient Gas Sample Line with Nafion
2. INOmax DSIR Plus
3. Oxygen Source
4. Breathing Circuit Hose
5. Injector Module
6. Injector Module Electrical Cable
7. NO/N₂ Injector Tube
8. 22F X 15M Adapter
9. Humidifier
10. Breathing Circuit
11. Temperature Probe
12. Gas Sample Tee
13. 22M/15F X 22M/15F Adapter
14. 22 mm ID X 22 mm ID Cuff Adapter
15. Optiflow Tracheostomy
16. Optiflow Nasal Cannula
17. Optiflow Mask

Figure 3-32 Example: Fisher & Paykel Healthcare Optiflow Breathing Circuit Diagram
Hamilton Arabella Nasal CPAP

Figure 3-33 Example: Hamilton Arabella Nasal CPAP Circuit Diagram

1. Arabella
2. Patient Gas Sample Line with Nafion
3. INOmax DSIR Plus
4. NO/N₂ Injector Tube
5. Injector Module Electrical Cable
6. Injector Module
7. 22F X 15M Adapter
8. Humidifier
9. Heated Delivery Circuit
10. Temperature Probe
11. Universal Generator
12. Arabella Sample Tee
13. 90 Degree Sample Port Adapter
ICU Ventilator Circuit

1. Patient Wye
2. Patient Gas Sample Line with Nafion
3. Ventilator
4. Ventilator Expiratory Port
5. Ventilator Inspiratory Port
6. Patient Gas Sample Line Input Connection
7. INOmax DSIR Plus
8. NO/N₂ Injector Tube Front Panel Connection
9. Injector Module Electrical Cable Front Panel Connection
10. 22M/15F X 22M/15F Adapter
11. Injector Module Electrical Cable Connection
12. Injector Module NO/N₂ Injector Tube Connection
13. 22F X 15M Adapter
14. Humidifier Inlet
15. Humidifier
16. Humidifier Outlet
17. Gas Sample Tee

Figure 3-34 Example: General Ventilator Diagram
INOblender use with the NeoPuff

1. Oxygen Source
2. NeoPuff
3. T-Piece Circuit (with Duckbill Port)
4. Patient Connection
5. Temperature Probe
6. Humidified Resuscitation System Circuit
7. Humidifier
8. Oxygen Tubing
9. INOblender
10. INOMAX Inlet

Figure 3-35 Example: INOblender use with the NeoPuff
Sensormedics 3100A/B High Frequency Oscillatory Ventilator with a Filtered Circuit

**WARNING:** Always use a one-way valve to avoid high NO delivery.

**Caution:** Use only parts provided in disposable package #50071, and tightly secure all connections.

![Diagram of High Frequency Oscillatory Ventilator](image)

1. Sensormedics 3100A/B Ventilator
2. Ventilator Outlet
3. 22M Adapter
4. Injector Module
5. Injector Module Electrical Cable Connection
6. INOmax DSIR Plus
7. NO/N2 Injector Tube
8. 8 mm Tubing X 15M Adapter
9. One-Way Valve
10. Paw Limit Valve Control
11. Filter
12. Humidifier Inlet
13. Humidifier Outlet
14. Bias Flow Tube
15. Patient Gas Sample Line with Nafion
16. 90 Degree Sample Port Adapter
17. Dump Valve Control
18. Paw Control Valve

Figure 3-36 Example: High Frequency Oscillatory Ventilator Diagram
Sensormedics 3100A/B High Frequency Oscillatory Ventilator with a Rigid or Flexible Circuit

**WARNING:** Always use a one-way valve to avoid high NO delivery.

Figure 3-37 Example: High Frequency Oscillatory Ventilator Diagram

1. Sensormedics 3100A/B Ventilator
2. Ventilator Outlet
3. Injector Module
4. INOmax DS$_{NO}$ Plus
5. NO/N$_2$ Injector Tube Connection
6. Injector Module Electrical Cable Connection
7. One-Way Valve
8. 22 mm ID X 22 mm ID Cuff Adapter
9. Humidifier Inlet
10. Humidifier Outlet
11. Patient Gas Sample Line with Nafion
12. 90 Degree Sample Port Adapter
13. Bias Flow Tube
SLE Life Support SLE5000

Note:
- Validated for use outside of the United States.
- A one-way valve is not required for use during high frequency ventilation mode.

Figure 3-38 Example: SLE Life Support SLE5000 Circuit Diagram
Spontaneously Breathing Patient on a Mask Circuit

1. O₂ Tubing
2. 15M X 4.5 mm Adapter
3. 22M/15F X 22M/15F Adapter
4. Breathing Circuit Tee
5. Breathing Circuit Bag
6. Injector Module
7. Breathing Circuit Hose
8. Gas Sample Tee
9. 22M/15F X 22M/15F Adapter
10. One-Way Valve
11. Sealed Face Mask
12. One-Way Valve
13. Patient Gas Sample Line with Nafion
14. NO/N₂ Injector Tube
15. INOmax DSIR Plus
16. Injector Module Electrical Cable
17. O₂ Flowmeter (wall outlet or cylinder)

Figure 3-39 Example: Spontaneously Breathing Patient Circuit Diagram
Spontaneously Breathing Patient on a Nasal Cannula

The INOmax DSIR Plus can be used with a nasal cannula to deliver INOMAX concentrations from 5-80 ppm with 800 ppm cylinders, and 5-40 ppm with 400 ppm cylinders and an oxygen flow rate as low as two L/min. Conditioning of the oxygen flow prior to delivery through the injector module will help ensure the most accurate flow measurement. Conditioning can be achieved by adding 300 mm of 22 mm hose between the oxygen tubing and the Injector Module.

Figure 3-40 Example: Spontaneously Breathing Nasal Cannula Patient Circuit Diagram
Teleflex Medical Comfort Flo Humidification System

1. Patient Gas Sample Line with Nafion
2. INOmax DSIR Plus
3. Injector Module
4. System Pressure Relief Valve
5. Air/Oxygen Blender or Oxygen Blender
6. Oxygen Tubing
7. Temperature Probe (Short Cable)
8. Angled 22 mm Connector
9. Patient Circuit
10. Temperature Probe Connector
11. Second Temperature Probe Connector
12. Comfort Flo Cannula
13. Injector Module Electrical Cable
14. NO/N₂ Injector Tube
15. 22F X 15M Adapter
16. ConchaTherm Heated Humidifier
17. Temperature Probe (Long Cable)
18. 90 Degree Sample Port Adapter

Figure 3-41 Example: Teleflex Comfort Flo Patient Circuit Diagram
Vapotherm 2000i

Figure 3-42 Example: Vapotherm 2000i Circuit Diagram

1. INOmax DSIR Plus
2. O₂ Flowmeter
3. O₂ Tubing
4. 15M x 4.5 mm Adapter
5. 22M/15F x 22M/15F Adapter
6. 300 mm of 22 mm Hose
7. 22M/15F x 22M/15F Adapter
8. Injector Module
9. 15M x 4.5 mm Adapter
10. Vapotherm 2000i
11. Patient Delivery Tube
12. O₂ Tubing Sample Tee
13. Patient Cannula
14. Patient Gas Sample Line with Nafion
15. NO/N₂ Injector Tube
16. Injector Module Electrical Cable
Vapotherm Precision Flow

- The INOmax DSIR Plus adds NO/N₂ gas flow to the breathing circuit flow in proportion to the NO setting (up to 10% at 80 ppm for a 800 ppm cylinder and 40 ppm for a 400 ppm cylinder) and subtracts gas from the breathing circuit via gas sampling at a nominal flow rate of 0.23 L/min.
- These effects impact the delivered gas flow rate when using the Vapotherm Precision Flow. It is recommended that after an NO setting change the user checks the delivered gas flow rate and adjusts the gas source flow rate as necessary.
- Follow all manufacturer instructions for connection to the Vapotherm Precision Flow.

Figure 3-43 Example: Vapotherm Precision Flow Circuit Diagram

1. Patient Gas Sample Line with Nafion
2. INOmax DSIR Plus
3. Precision Flow Unit
4. Injector Module
5. Patient Delivery Tube
6. Oxygen Tubing Sample Tee
7. Patient Cannula
8. Injector Module Electrical Cable
9. NO/N₂ Injector Tube
INOmax DS$ \text{IR}^\text{®}$ Plus

4/ Transport
INOmax DSIR® Plus

4/ Transport
4/ Transport

Caution:

- It is recommended that a second (backup) transport regulator cap assembly is available during all transports.
- It is recommended that a second (backup) cylinder of INOMAX is available during all transports.

Transport Options

A. When moving the INOmax DS$_{IR}$ Plus as a unit (cart and cylinders), (see Section A below).
B. When removing INOmax DS$_{IR}$ Plus and INOblender from the cart (see Section B below).
C. When using the INOblender as a stand-alone device (see Section C below).
D. When using a separate INOmax DS$_{IR}$ Plus and INOblender for transport (see Section D below).

A. Intrahospital transport (within the hospital) when moving the INOmax DS$_{IR}$ Plus as a unit (cart and cylinders)

1. Refer to cylinder duration chart as a guide to determine if there is enough drug to last through the transport, including unexpected delays. Bring additional cylinders as appropriate.
2. Connect the INOblender oxygen hose to a 3.5 bar (50 psig) portable oxygen source.
3. Manually ventilate the patient using the INOblender while configuring the INOmax DS$_{IR}$ Plus to the transport ventilator (see figure 4-13 or 4-14).
4. Upon return:
   a. Manually ventilate the patient while reattaching the INOmax DS$_{IR}$ Plus to the bedside ventilator.
   b. Confirm the operation of the INOmax DS$_{IR}$ Plus.
   c. Reconnect the INOblender oxygen hose to a 3.5 bar (50 psig) oxygen source.
B. Intrahospital transport (within the hospital) when removing the INOmax DSIR Plus and INOblender from the cart.

Check the INOMAX drug cylinders for the correct product identity, cylinder concentration, and expiration date.

1. Connect the high pressure transport regulator/cap assembly to the INOMAX transport cylinder, and tighten the fitting to the INOMAX cylinder.
2. Place the cap assembly over the INOhmter.
3. Open then close the INOMAX transport cylinder valve.

Note: Be sure to align the key way inside the Cap Assembly with the ibutton on the INOhmter.

4. Check for adequate cylinder pressure. Verify cylinder has at least 35 bar (500 psig).
5. Monitor pressure gauge for 30 seconds for any signs of pressure decrease. If no signs of pressure decrease is observed, the leak test is successful.

6a. Depressurize the transport cylinder regulator hose.
6b. Connect regulator hose to the available drug inlet.
7. Open INOMAX transport cylinder valve.

8a. Immediately close the valve for the cylinder on the cart and remove regulator hose.
8b. Depressurize the cylinder regulator hose using the purge port.

9a. Disconnect the cart infrared (IR) cable.
9b. Connect the IR cable from the transport regulator cap assembly to the back of the INOmax DS_{IR} Plus.

Note: Do not attempt to connect the transport regulator/cap assembly electrical plug to the INOblender outlet port. This will damage the connector plug electrical pins.
10. Disconnect the INOblender inlet hose from the back of the INOmax DSIR Plus, then loosen the clamp on the back of the INOmax DSIR Plus and remove from the cart. Position the INOmax DSIR Plus on the transport device and secure.

11. Loosen the clamp on the back of the INOblender, remove the INOblender from the cart and secure to the transport device.

12. Reconnect the INOblender inlet hose to the INOmax DSIR Plus and slide the quick-connect cover into place.

Upon return:

13. Return the INOblender to the cart and secure the clamp assembly.

14. Return the INOmax DSIR Plus to the cart and secure the clamp assembly.

15. Connect the INOblender inlet hose to the INOmax DSIR Plus INOblender outlet and slide the quick-connect cover into place.

Note: INOblender extension hose (PN 10014) will be required if the INOmax DSIR Plus and the INOblender are positioned more than two feet apart.
Once the devices are secure on the cart:

16a. Open/close INOMAX cylinder, then depressurize the regulator hose by pressing the regulator hose into the purge port.

16b. Open the INOMAX cylinder valve and insert the regulator hose into available INOMAX gas inlet.

16c. Close transport cylinder valve and remove the regulator hose from the INOMAX gas inlet and depressurize the regulator hose using the purge port.

16d. Disconnect the transport cylinder IR cable from the back of the INOmax DS$_{IR}$ Plus.

16e. Attach IR cable from the cart to the back of the INOmax DS$_{IR}$ Plus.

**Note:** Do not attempt to connect the transport regulator/cap assembly electrical plug to the INOblender outlet port. This will damage the connector plug electrical pins.
C. When using the INOblender as a stand-alone device.

Important: Read the INOblender Operation Manual PN 20181 before using the INOblender. Follow instructions and obey all Warnings and Cautions. Verify INOblender function using the INOblender test on page 4-9 or INOblender stand-alone pre-use checkout on page 4-10.

![Diagram](image1)

Figure 4-1

Normally the INOblender receives INOMAX from the INOmax DSIR Plus outlet (INOMAX cylinder supplies both devices; see Figure 4-1).

![Diagram](image2)

Figure 4-2

As a stand-alone device the INOMAX cylinder supplies INOMAX directly to the INOblender. (see Figure 4-2).

1. Disconnect INOMAX regulator hose from back of INOmax DSIR Plus.
2. Disconnect INOblender hose from back of INOmax DSIR Plus.
3. Connect INOMAX regulator hose to INOblender inlet hose.
4. Verify INOMAX cylinder valve is open.
5. Verify cylinder has at least 35 bar (500 psig).

Adjust Settings

6. Turn the INOblender setting dial to the desired concentration 5 to 80 ppm for an 800 ppm cylinder (5 to 40 ppm for a 400 ppm cylinder).
7. Turn the O₂ flowmeter to the desired flow rate (5 to 14 L/min).
8. Squeeze the manual resuscitator 3-4 times to purge the NO₂ from the system.

The INOblender is now ready for patient use.
INOblender Test Using the INOmax DSIR Plus to Analyze Output

**WARNING:**
- The purge procedure must be followed to help ensure NO₂ is purged from the system before the manual resuscitator bag is connected to the patient.
- The manual bag should be squeezed repeatedly during use to avoid NO₂ building up in the bag.
- If the bag is not squeezed repeatedly while delivering INOMAX, the bag should be removed from the patient and the bag purge procedure performed before continuing.
- The INOblender should be upright when setting the oxygen flowrate for accurate setting.
- Do not use pneumatically powered nebulizers with the INOblender. This will result in significant over delivery of INOMAX in excess of 80 parts per million (ppm) with 800 ppm cylinders and 40 parts per million with 400 ppm cylinders.
  - The INOblender outlet pressure has been validated for use up to 0.4 bar (5.8 psig) pressure. The amount of back-pressure generated by pneumatic nebulizers is significantly greater, 1.4 to 2.0 bar (20-30 psig), and will result in over delivery of INOMAX in excess of 80 ppm. The user adjusted dose setting on the INOblender will not correlate with, or have an effect on the actual delivered dose.
  - In addition, the INOblender flowmeter is not back-pressure compensated and will display a lower flow rate than actual when pressure is applied to the outlet.

**Caution:**
- When not in use, the oxygen flowmeter should be turned off.
- A user may determine that some clinical conditions may necessitate the use of an oxygen/air blender with the INOblender to achieve FiO₂ levels less than 100%.
- Delivered INOMAX dose from the INOblender is affected by varying oxygen concentrations (see table below):

<table>
<thead>
<tr>
<th>FiO₂</th>
<th>INOblender Accuracy Specification (at 50 psig)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.0</td>
<td>+/- 20% of set value or 2 ppm whichever is greater</td>
</tr>
<tr>
<td>0.21 to 0.95</td>
<td>+/- 30% of set value or 3 ppm whichever is greater</td>
</tr>
</tbody>
</table>
INOblender Test

Note:

- Confirm INOblender inlet hose is connected to the INOMAX regulator hose and the Quick-Connect cover is in place.
- Confirm 3.5 bar (50 psig) oxygen supply hose is connected to the \( O_2 \) inlet fitting on the back of the INOblender.

1. Connect the pre-use set up to the front of the INOblender.
2. Confirm the injector module is not attached to the pre-use set-up.
3. On the INOblender, set the INOMAX dose and flow to:

\[
\begin{array}{ccc}
\text{Cylinder Concentration (ppm)} & 800 & 400 \\
\text{INOblender Set Dose (ppm)} & 40 & 20 \\
\text{INOblender Flow (L/min)} & 10 & \\
\end{array}
\]

4. Verify NO value on the INOmax DSIR Plus using the table below.
5. Turn the dose and oxygen flow to zero.
6. Remove the pre-use set-up from the INOblender.

<table>
<thead>
<tr>
<th>Cylinder Concentration (ppm)</th>
<th>800</th>
<th>400</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acceptable NO Value (ppm)</td>
<td>32 - 48</td>
<td>16 - 24</td>
</tr>
</tbody>
</table>
INOblender Stand-Alone Pre-use Checkout

**Caution:** To help ensure proper operation, complete the pre-use checkout prior to each use.

### High-Pressure Leak Test

1. Make sure NO dose setting dial is turned to zero and flow meter is OFF.
2. Open and then close the INOMAX cylinder valve.
3. Check the INOMAX drug cylinders for the correct product identity, cylinder concentration, and expiration date. Verify cylinder has at least 35 bar (500 psig) and tighten the fitting to the INOMAX cylinder.
4. Monitor pressure gauge for 30 seconds for any signs of pressure decrease. If no pressure decrease is observed, high-pressure leak test successful, proceed to Delivery Confirmation and Purge.
5. If observed pressure decrease continues, see page 4-23, Cylinder Leak Check.
6. If leak cannot be traced, replace the INOblender.

### Delivery Confirmation and Purge

1. Set the INOblender to 40 ppm when using an 800 ppm cylinder (20 ppm for an 400 ppm cylinder). Confirm INOMAX cylinder valve is closed.
2. Set the oxygen flow on the INOblender flow meter to 10 L/min to begin purge.
3. Ensure the pressure gauge decreases approximately 14 bar (200 psig) in 10 seconds (± two seconds).
4. Continue purging until pressure gauge reads zero.

**Note:** If the pressure does not decrease, then the INOblender is not delivering NO and the INOblender should be replaced.
D. InterHospital Transport (Between Hospitals) when using a separate INOmax DSIR Plus and INOblender for transport

**WARNING:**
- If the INOmax DSIR Plus or INOblender is to be used in a transport vehicle, they should be affixed to the transport mounting post which is part of the transport mounting bracket assembly (part number 50041).
- The transport mounting post and/or the transport mounting bracket assembly should be secured to the transport isohlete/transport gurney in a manner which will secure the INOmax DSIR Plus/INOblender.

Prior to Leaving the Hospital
1. Complete the pre-use checkout for the transport INOmax DSIR Plus unit.
   a. The pre-use checkout is mandatory to ensure proper function of the INOmax DSIR Plus, INOMAX regulator, transport regulator/cap assembly and INOblender.
   b. Change the injector module and/or perform a high calibration if monitored values are out of range during the pre-use checkout.
2. Bring appropriate backup equipment in case of a malfunction during the transport (see caution above).

**Items recommended for interhospital transport:**
- INOmax DSIR Plus
- Transport regulator/cap assembly (two)
- D-size transport INOMAX cylinder (two)
- INOblender
- Injector module (two)
- Injector module cables (two)
- Transport mounting bracket assembly (optional)
- Disposables
  - NO/N₂ injector tube (two)
  - Patient gas sample line with Nafion (two)
  - Water separator cartridge (two)
- Properly secure the INOmax DSIR Plus, INOblender and INOMAX cylinders per hospital/air carrier protocols.

Transport equipment weight should be calculated to assure transport system meets weight allowance.

<table>
<thead>
<tr>
<th>Part Description</th>
<th>Weight</th>
<th>Dimensions</th>
</tr>
</thead>
<tbody>
<tr>
<td>INOmax DSIR Plus</td>
<td>5.3 kg / 11.7 lb</td>
<td>350 mm (W) X 220 mm (H) X 160 mm (D)</td>
</tr>
<tr>
<td>INOblender</td>
<td>1.5 kg / 3.3 lb</td>
<td>200 mm (W) X 120 mm (H) X 110 mm (D)</td>
</tr>
<tr>
<td>INOMAX D-size Cylinder</td>
<td>3.6 kg / 8.0 lb</td>
<td>111 mm (W) X 517 mm (H)</td>
</tr>
<tr>
<td>Transport Regulator/Cap Assembly</td>
<td>0.90 kg / 2.0 lb</td>
<td>N/A</td>
</tr>
<tr>
<td>Transport Mounting Bracket Assembly</td>
<td>0.97 kg / 2.14 lb (post only 0.29 kg / 0.64 lb)</td>
<td>N/A</td>
</tr>
<tr>
<td>INOblender Extension Hose</td>
<td>0.06 kg / 0.13 lb</td>
<td>N/A</td>
</tr>
</tbody>
</table>

Note: All sizes and weights are approximate and may vary slightly.
Duration Chart
INOMAX Cylinder Luxfer 2L-Size

For a Luxfer-Size *800 ppm and **400 ppm Cylinder Concentrations
***(Illustrative Only)

<table>
<thead>
<tr>
<th>INOMAX Dose (ppm)</th>
<th>Cylinder Conc. (ppm)</th>
<th>FLOW</th>
<th>5 L/min</th>
<th>10 L/min</th>
<th>20 L/min</th>
<th>40 L/min</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td><strong>800</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>2.5</td>
<td>6.1 Days</td>
<td>3.1 Days</td>
<td>1.5 Days</td>
<td>18.4 Hours</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>5</td>
<td>3.1 Days</td>
<td>1.5 Days</td>
<td>18.3 Hours</td>
<td>9.2 Hours</td>
<td></td>
</tr>
<tr>
<td>20</td>
<td>10</td>
<td>1.5 Days</td>
<td>18.1 Hours</td>
<td>9.0 Hours</td>
<td>4.5 Hours</td>
<td></td>
</tr>
<tr>
<td>40</td>
<td>20</td>
<td>17.6 Hours</td>
<td>8.8 Hours</td>
<td>4.4 Hours</td>
<td>2.2 Hours</td>
<td></td>
</tr>
<tr>
<td>80</td>
<td>40</td>
<td>8.3 Hours</td>
<td>4.2 Hours</td>
<td>2.1 Hours</td>
<td>1.0 Hours</td>
<td></td>
</tr>
</tbody>
</table>

This chart is representative of a range of doses available on the INOmax DSIR Plus. Doses higher than 20 ppm as a therapeutic dose are not recommended.

*** All calculations for the table above are based on a full cylinder of 155 bar (2248 psig), Luxfer 2 Liter cylinder, with a cylinder change at 14 bar (200 psig). The figures are calculated based on a total continuous breathing circuit gas flow and a cylinder conversion factor of 2.0 liters per bar (0.14 liters per psig).

• INOMAX flow = [Desired dose × total ventilator flow] ÷ [Cylinder concentration - desired dose]
• Cylinder volume = Cylinder conversion factor × cylinder pressure (bar/psig)
• Cylinder duration (hours) = (Cylinder volume ÷ INOMAX flow rate) ÷ 60

Calculations are considered estimates and may vary under clinical circumstances.
Duration Chart
INOMAX Cylinder D-Size

For a D-Size *800 ppm and **400 ppm Cylinder Concentrations
*** (Illustrative Only)

<table>
<thead>
<tr>
<th>INOMAX Dose (ppm)</th>
<th>Cylinder Conc.</th>
<th>5 L/min</th>
<th>10 L/min</th>
<th>20 L/min</th>
<th>40 L/min</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>*800</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>2.5</td>
<td>7.0 Days</td>
<td>3.5 Days</td>
<td>1.8 Days</td>
<td>21 Hours</td>
</tr>
<tr>
<td>10</td>
<td>5</td>
<td>3.5 Days</td>
<td>1.7 Days</td>
<td>21 Hours</td>
<td>10.5 Hours</td>
</tr>
<tr>
<td>20</td>
<td>10</td>
<td>1.7 Days</td>
<td>20.7 Hours</td>
<td>10.3 Hours</td>
<td>5.2 Hours</td>
</tr>
<tr>
<td>40</td>
<td>20</td>
<td>20 Hours</td>
<td>10 Hours</td>
<td>5 Hours</td>
<td>2.5 Hours</td>
</tr>
<tr>
<td>80</td>
<td>40</td>
<td>9.5 Hours</td>
<td>4.8 Hours</td>
<td>2.4 Hours</td>
<td>1.2 Hours</td>
</tr>
</tbody>
</table>

Typically used in transport

This chart is representative of a range of doses available on the INOmax DSIR Plus. Doses higher than 20 ppm as a therapeutic dose are not recommended.

*** All calculations in the table above are based on a full cylinder, 128 bar (1850 psig), “D” cylinder, with a cylinder change at 14 bar (200 psig). The figures are calculated based on a total continuous breathing circuit gas flow using a cylinder conversion factor of 2.8 liters per bar (0.19 liters per psig).

- INOMAX flow = [Desired dose × total ventilator flow] ÷ [Cylinder concentration - desired dose]
- Cylinder volume = Cylinder conversion factor × cylinder pressure (bar/psig)
- Cylinder duration (hours) = (Cylinder volume + INOMAX flow rate) ÷ 60

Calculations are considered estimates and may vary under clinical circumstances.
Transport Regulator/Cap Assembly Application

Note: Before leaving the bedside (intrahospital transport) or hospital (interhospital transport), check the INOMAX cylinder for the correct product identity labels, cylinder concentration and expiration date. Verify cylinder has at least 35 bar (500 psig) and tighten the fitting to the INOMAX cylinder.

1. Connect a high pressure regulator to an INOMAX cylinder and hand-tighten the fitting to the INOMAX cylinder (see Figure 4-3).

   Note: • For the CGA-type regulator, ensure the white plastic tip is in place on the regulator connector and not chipped or cracked. Remove and replace as necessary (see Replacing the CGA 626 tip on the INOMAX regulator, page 7-8).
   • For the ISO-type regulator connector, ensure that the O-ring is present and is not damaged (see Replacing the tip on the INOMAX regulator, page 7-9)

2. Connect the INOMAX regulator hose to one of the INOMAX inlets on the back of the INOmax DSIR Plus (see Figure 4-3).

Figure 4-3
3. Connect the infrared cable from the transport regulator/cap assembly to the back of the INOmax DS$_{IR}$ Plus (see Figure 4-4).

**Note:** Notice that the connector clicks to indicate that it is latched in place. Do not attempt to connect the transport regulator cap assembly electrical plug to the INOblender outlet port. This will damage the connector plug electrical pins.

**Caution:** When using the transport regulator/cap assembly (PN 10022, CGA or 10041, ISO) ensure the cap is fully seated and in place on the INOmeter and the infrared cable is connected and latched to the infrared connector port on the back of the INOmax DS$_{IR}$ Plus.

4. Place the cap assembly over the INOmeter (see Figure 4-5).

**Note:** Be sure to align the keyway inside the cap assembly with the iButton on the INOmeter (see Figure 4-5 and 4-6).

**Figure 4-4**

**Figure 4-5**

**Figure 4-6**

**Figure 4-7**

**Figure 4-8**

5. Grasp the cap assembly to open cylinder valve (see Figure 4-7 and 4-8).
Final Set-up Diagram

The following diagram (see Figure 4-9) and photo illustrates all of the components connected.

![Diagram showing INOMAX connection](image)

**Figure 4-9**

Communication will take place between the INOmax DS$_{IR}$ Plus and the INOmeter after the boot up phase of the INOmax DS$_{IR}$ Plus is complete.

**WARNING:** Loss of communication between the INOmax DS$_{IR}$ Plus and the INOMAX cylinder for more than one hour will result in interruption of INOMAX delivery.

**Note:**
- Cylinder icons are not visible and the NO delivery setpoint button will remain inactive until the INOmax DS$_{IR}$ Plus recognizes an INOMAX cylinder.
- When using the transport regulator/cap assembly only one cylinder will be displayed (see Figure 4-10).
Changing INOMAX Cylinders

Prepare new cylinder

1. Remove INOmeter wrapping (if applicable).
2. Remove INOmeter lock.
3. Attach spare transport regulator/cap assembly to new cylinder.
4. Open, then close cylinder valve.
5. Confirm cylinder pressure is at least 35 bar (500 psig).
6. Observe that gauge pressure is steady for 30 seconds.
7. Purge regulator using purge port on the back of the INOMax DSIR Plus.
8. Re-open new cylinder valve.
9. Connect new cylinder regulator hose to the available INOMAX inlet.
10. Remove previously used cylinder regulator hose from the INOMAX inlet.
11. Replace current IR cable with the IR cable from the spare transport regulator/cap assembly to back of the INOMax DSIR Plus.
12. Close previously used cylinder valve.
13. Purge regulator on previously used cylinder.
14. Insert INOmeter lock on previously used cylinder.
PN 10014 INOblender Extension Hose User Instructions

To mount the INOblender away from the INOmax DSIR Plus, an INOblender extension hose can be added between the INOmax DSIR Plus and the INOblender. To add the extension hose, refer to Figure 4-11 and continue with the following steps.

**WARNING:**
- Only use one length of extension hose between devices to minimize the risk of NO₂ formation within the hose.
- If the INOmax DSIR Plus or INOblender is used in a transport vehicle, the INOmax DSIR Plus and INOblender should be securely attached to the transport isobette/gurney. Secure each device to the transport mounting post, which is part of the transport mounting bracket assembly (part number 50041), to prevent possible injury. For detailed information, refer to the INOmax DSIR Plus Operation Manual for complete information.

1. Mount the INOmax DSIR Plus and INOblender per the INOmax DSIR Plus Operation Manual instructions.
2. Connect the male end of the extension hose to the INOblender inlet hose (1).
3. Connect the female end of the extension hose to the INOblender outlet (2) at the back of the INOmax DSIR Plus.
4. Slide the Quick-Connect covers into place (3).

Figure 4-11 Adding an INOblender Extension Hose between the INOmax DSIR Plus and INOblender during transport
As a stand-alone device, the INOblender is not connected to the INOmax DSIR Plus. The INOblender extension hose may be placed between the INOMAX regulator and the INOblender to provide additional length between the INOMAX cylinder and the INOblender. To add the extension hose, refer to Figure 4-12 and continue with the following steps:

1. Connect the male end of the extension hose to the INOblender inlet hose (1).
2. Connect the female end of the extension hose to the INOMAX regulator hose (2).
3. Slide the Quick-Connect covers into place (3).

Figure 4-12 Adding an INOblender Extension Hose between the INOMAX Regulator and INOblender during transport
Connection to a Dual-Limb Transport Ventilator Circuit

1. Patient Wye
2. Expiratory Breathing Circuit Hose
3. Patient Gas Sample Line with Nafion
4. Ventilator Expiratory Valve
5. Ventilator
6. INOmax DSIR Plus
7. Ventilator Inspiratory Port
8. 22M/15F X 22M/15F Adapter
9. Injector Module Electrical Cable
10. NO/N₂ Injector Tube
11. Injector Module
12. Inspiratory Breathing Circuit Hose
13. Gas Sample Tee

Figure 4-13 Example: Transport Ventilator Diagram
Connection to a Single-Limb Transport Ventilator Circuit

1. PEEP Valve
2. Patient Wye
3. Circuit Hose
4. Patient Gas Sample Line with Nafion
5. Ventilator
6. INOmax DSIR Plus
7. Ventilator Inspiratory Port
8. 22M/15F X 22M/15F Adapter
9. Injector Module Electrical Cable
10. NO/N₂ Injector Tube
11. Injector Module
12. Inspiratory Breathing Circuit Hose
13. Gas Sample Tee

Figure 4-14 Example: Single-Limb Transport Ventilator Diagram
**WARNING:** If the INOmax DSIR Plus is to be used in a transport vehicle, it should be affixed to the transport mounting post, see Figure 4-15.

Figure 4-15 Universal Mounting Post

The universal mounting post has a machined recess with an integrated cap to prevent twisting or accidental release of the device if the mounting clamp assembly becomes loose.

---

**Note:** The universal mounting post has a machined recess with an integrated cap to prevent twisting or accidental release of the device if the mounting clamp assembly becomes loose.

---

**Figure 4-16 Transport Mounting Bracket Assembly Attached to Transport Isolette**

1. Transport Isolette
2. Isolette Bar
3. INOmax DSIR Plus or INOblender Mounting Area
4. Universal Mounting Post
5. Isolette Handle
6. Transport Mounting Bracket Assembly (PN 50041, includes universal mounting post)

---

**Caution:**

- When using the transport regulator/cap assembly (PN 10022, CGA or 10041, ISO) ensure the cap is fully seated and in place on the INOmeter and the infrared cable is connected and latched to the infrared connector port on the back of the INOmax DSIR Plus (see page 4-17).
- It is recommended that a second transport regulator/cap assembly is available during all transports.

**Note:** Do not attempt to connect the transport regulator/cap assembly electrical plug to the INOblender outlet port. This will damage the connector plug electrical pins.

---

**Figure 4-17**
Cylinder Leak Check

If a leak is suspected during the high pressure leak test, the following steps can be taken to check for leaks (see Figure 4-18 for possible cylinder gas leak locations) in the INOMAX regulator or INOMAX cylinder.

1. Check the INOMAX drug cylinders for the correct product identity, cylinder concentration, and expiration date. Verify cylinder has at least 35 bar (500 psig), and tighten the fitting to the INOMAX cylinder.

2. Apply soapy water to points #1, #2, #3, #5 and #6 (see Figure 4-18); if bubbles form, there is a leak.

3. If there are no bubbles, the leak may be inside the INOMAX DSIR Plus and cannot be repaired. Replace the INOMAX DSIR Plus and contact Customer Support.

Recommended actions should a leak be detected:

1. A leak detected at points #1 and #2 may be corrected by tightening the INOMAX regulator hand wheel.
   a. If cylinder valve is open, close cylinder valve, relieve regulator pressure by purging, then tighten INOMAX regulator hand wheel.
   b. Open cylinder valve and reapply soapy water to points #1 and #2.
   c. If bubbles form, there is a leak.
   d. Remove INOMAX regulator and check for damage. For the CGA type regulator connector, check the white plastic tip on the INOMAX regulator for chips or cracks. Replace if necessary. For the ISO type regulator connector, check that the O-ring is present and not damaged. Replace if necessary (see replacing the tip/O-ring on the INOMAX regulator, Pages 7-8/7-9). Repeat step b.

2. If a leak is detected between the regulator body and regulator end cap (see point #3) replace INOMAX regulator and contact Customer Support.

3. A leak detected at the cylinder valve nut connection (see point #5) may not be repaired. Replace INOMAX cylinder and contact Customer Support.

4. A leak detected at the safety pressure release device (see point #6) may not be repaired. Replace INOMAX cylinder and contact Customer Support.

Note: Refer to hospital policies and procedures for dealing with leaking gas cylinders. Additional information regarding environmental effects can be found in the section 1/ General Information.
5/ Alarms and Troubleshooting
5/ Alarms and Troubleshooting
5/ Alarms and Troubleshooting

**WARNING:**
- Abrupt discontinuation of INOMAX may lead to worsening oxygenation and increasing pulmonary artery pressure, i.e., Rebound Pulmonary Hypertension Syndrome. To avoid abrupt discontinuation, utilize the INOblender or integrated pneumatic backup if necessary. If Rebound Pulmonary Hypertension occurs, reinstate INOMAX therapy immediately. (See the INOMAX prescribing Information for further details).
- If the high NO₂ alarm activates, the delivery system should be assessed for proper setup while maintaining INOMAX delivery. Adjust INOMAX and/or FiO₂ as appropriate. (See INOMAX Prescribing Information for further details on the effects of Nitrogen Dioxide, NO₂). If unable to determine the cause of the increased NO₂ levels, call Customer Support and do not discontinue therapy.

### Continuous Audible Tone

<table>
<thead>
<tr>
<th>Alarm</th>
<th>Possible Cause</th>
<th>Recommended Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>Continuous Audible Tone</td>
<td>A component within the INOmax DS₉R Plus has failed.</td>
<td>1. If the INOblender is available, manually ventilate the patient (see INOblender Operation Manual).</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. Replace the delivery system and remove from service.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3. Contact Customer Support.</td>
</tr>
</tbody>
</table>

### Alarm Help

#### High Priority Alarms

*All of these actions can be performed while delivering INOMAX to the patient:*

<table>
<thead>
<tr>
<th>Alarm</th>
<th>Possible Cause</th>
<th>Recommended Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. High NO</td>
<td>The High NO alarm level may be inappropriately set.</td>
<td>Confirm High NO alarm limit is set appropriately.</td>
</tr>
<tr>
<td></td>
<td>Circuit setup incorrect.</td>
<td>Check circuit for correct setup.</td>
</tr>
<tr>
<td></td>
<td>NO sensor may require calibration.</td>
<td>1. Perform low calibration.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. Perform high NO calibration.</td>
</tr>
<tr>
<td></td>
<td>Injector module may not be functioning properly.</td>
<td>1. Manually ventilate patient with the INOblender or turn integrated pneumatic backup delivery ON</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. Change injector module (INOMAX delivery will be interrupted).</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3. Change injector module cable (INOMAX delivery will be interrupted).</td>
</tr>
</tbody>
</table>
# High Priority Alarms

All of these actions can be performed while delivering INOMAX to the patient:

<table>
<thead>
<tr>
<th>Alarm</th>
<th>Possible Cause</th>
<th>Recommended Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>2. Low NO</td>
<td>The Low NO alarm setting may be inappropriately set.</td>
<td>Confirm Low NO alarm limit is set appropriately.</td>
</tr>
</tbody>
</table>
|                                 | Circuit setup incorrect. The patient gas sample line or NO/N₂ injector tube may be disconnected. | 1. Check circuit for correct setup.  
2. Confirm water bottle, water separator cartridge, NO/N₂ injector tube and patient gas sample line are in place. |
| Loss of NO delivery.            | If loss of NO delivery is suspected, manually ventilate patient with the INOblender or turn integrated pneumatic backup delivery ON | 1. Perform low calibration  
2. Perform high calibration.                                                      |
| NO sensor may require calibration. |                                                                              | 1. Manually ventilate patient with the INOblender  
or turn integrated pneumatic backup delivery ON  
2. Replace injector module (INOMAX delivery will be interrupted).  
3. Replace injector module cable (INOMAX delivery will be interrupted). |
| Injector module may not be functioning properly. |                                                                              | 1. Confirm the O-ring on the sensor is correctly seated and the sensor cover is fully closed.  
2. Contact Customer Support.                                                       |
## High Priority Alarms

All of these actions can be performed while delivering INOMAX to the patient:

<table>
<thead>
<tr>
<th>Alarm</th>
<th>Possible Cause</th>
<th>Recommended Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>3. High NO₂</td>
<td>The High NO₂ alarm setting may be inappropriately set.</td>
<td>Confirm High NO₂ alarm limit is set appropriately.</td>
</tr>
<tr>
<td></td>
<td>The patient circuit setup may be incorrect.</td>
<td>Check circuit for correct setup.</td>
</tr>
<tr>
<td></td>
<td>Incomplete system purge.</td>
<td>Repeat the purge procedure with the injector module out of the patient breathing circuit. Use INOblender if necessary.</td>
</tr>
<tr>
<td></td>
<td>Monitored NO₂ value is too high.</td>
<td>Consider reducing the INOMAX dose according to INOMAX prescribing information.</td>
</tr>
<tr>
<td></td>
<td>NO₂ sensor may require calibration.</td>
<td>1. Perform low calibration.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. Perform high NO₂ calibration.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3. Contact Customer Support.</td>
</tr>
<tr>
<td>4. High O₂</td>
<td>The High O₂ alarm setting may be inappropriately set.</td>
<td>Confirm High O₂ alarm limit is set appropriately.</td>
</tr>
<tr>
<td></td>
<td>The patient breathing circuit setup may be incorrect.</td>
<td>Check circuit for correct setup.</td>
</tr>
<tr>
<td></td>
<td>O₂ sensor may require calibration.</td>
<td>1. Perform low calibration.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. Perform high O₂ calibration.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3. Contact Customer Support.</td>
</tr>
<tr>
<td>5. Low O₂</td>
<td>The Low O₂ alarm setting may be inappropriately set.</td>
<td>Confirm Low O₂ alarm limit is set appropriately.</td>
</tr>
<tr>
<td></td>
<td>The patient breathing circuit setup may be incorrect.</td>
<td>Check circuit for correct setup. Ensure patient gas sample line connections are secure.</td>
</tr>
<tr>
<td></td>
<td>O₂ sensor may require calibration.</td>
<td>1. Perform low calibration.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. Perform high O₂ calibration.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3. Contact Customer Support.</td>
</tr>
</tbody>
</table>

INOMAX can dilute O₂ concentration set at the ventilator by up to 10%.
# High Priority Alarms

All of these actions can be performed while delivering INOMAX to the patient:

<table>
<thead>
<tr>
<th>Alarm</th>
<th>Possible Cause</th>
<th>Recommended Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>6. Cylinder Not Detected</td>
<td>INOmax DS&lt;sub&gt;IR&lt;/sub&gt; Plus infrared cart cable is not connected or has failed.</td>
<td>Confirm infrared cart cable is connected to infrared connector on back of INOmax DS&lt;sub&gt;IR&lt;/sub&gt; Plus.</td>
</tr>
<tr>
<td></td>
<td>IR interference.</td>
<td>1. Reposition/rotate INOmax DS&lt;sub&gt;IR&lt;/sub&gt; Plus cart.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. Remove any obstacle between INOmeter and cart.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3. Move the interfering light source.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>4. Move the INOmax DS&lt;sub&gt;IR&lt;/sub&gt; Plus cart to reduce the high intensity light in the area of the INOmeter.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5. Shield the INOmeter from the suspect light source.</td>
</tr>
<tr>
<td></td>
<td>INOmeter may have failed.</td>
<td>Replace INOMAX cylinder on cart.</td>
</tr>
<tr>
<td></td>
<td>Transport Cap not connected to the INOmeter (if utilizing a transport regulator/cap assembly)</td>
<td>1. Connect transport regulator/cap assembly cable to infrared connector on back of INOmax DS&lt;sub&gt;IR&lt;/sub&gt; Plus.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. Confirm transport cap assembly is over INOmeter.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3. Replace transport regulator/cap assembly and properly align.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>4. Replace INOMAX cylinder.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5. Contact Customer Support.</td>
</tr>
<tr>
<td>7. Cylinder Valve Closed</td>
<td>INOMAX cylinder valve is closed.</td>
<td>Confirm INOMAX cylinder valve is fully open.</td>
</tr>
<tr>
<td></td>
<td>When two cylinders are present on the cart, open cylinder may not be visible to the IR system due to interference or obstruction.</td>
<td>1. Reposition INOmax DS&lt;sub&gt;IR&lt;/sub&gt; Plus cart.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. Remove any obstacle between INOmeter and cart.</td>
</tr>
<tr>
<td></td>
<td>INOmeter may have failed.</td>
<td>1. Replace INOMAX cylinder.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. Contact Customer Support.</td>
</tr>
</tbody>
</table>
### High Priority Alarms

All of these actions can be performed while delivering INOMAX to the patient:

<table>
<thead>
<tr>
<th>Alarm</th>
<th>Possible Cause</th>
<th>Recommended Action</th>
</tr>
</thead>
</table>
| 8. Delivery Failure          | Over-delivery of INOMAX or an internal error has been detected.               | 1. Manually ventilate patient with the INOblender or turn integrated pneumatic backup delivery ON  
2. Turn INOmax DSIR Plus to standby, then restart.  
   Once device is ready to restart therapy:  
3. Turn integrated back up off (if used) and then set the INOMAX dose.  
4. Confirm delivery and check alarms.  
5. Contact Customer Support. |
| 9. Delivery Stopped          | MONITORED NO > 100 ppm for at least 12 seconds                                | Manually ventilate patient with the INOblender or turn integrated pneumatic backup delivery ON |
|                              | Drug past expiry date.                                                        | Replace INOMAX cylinder.                                                             |
|                              | Drug concentration mismatch.                                                  | Replace INOMAX cylinder.                                                             |
|                              | INOmeter may have failed.                                                     | Replace INOMAX cylinder.                                                             |
|                              | INOMAX cylinder valve is closed.                                              | Open INOMAX cylinder valve.                                                          |
|                              | INOMAX cylinder is not detected.                                              | 1. Replace INOMAX cylinder.                                                          
2. Contact Customer Support. |
2. Remove expired INOMAX cylinder from INOmax DSIR Plus cart.  
3. Connect an INOMAX cylinder with a valid expiration date.  

**WARNING:** Delivery Stopped will occur two minutes from point when Drug Past Expiry Date alarm is activated.
## High Priority Alarms

All of these actions can be performed while delivering INOMAX to the patient:

<table>
<thead>
<tr>
<th>Alarm</th>
<th>Possible Cause</th>
<th>Recommended Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>11. Drug Concentration Mismatch</td>
<td>INOMAX cylinder is the wrong concentration.</td>
<td>1. Close mismatched cylinder valve.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. Remove wrong concentration INOMAX cylinder from INOmax DS&lt;sub&gt;IR&lt;/sub&gt; Plus cart.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3. Connect an INOMAX cylinder with a valid concentration to INOmax DS&lt;sub&gt;IR&lt;/sub&gt; Plus.</td>
</tr>
</tbody>
</table>

**WARNING:** Delivery Stopped will occur two minutes from point when Cylinder Concentration Mismatch alarm is activated.

| 12. Injector Module Fail       | The injector module electrical cable may be disconnected.            | Disconnect and reconnect both ends of the injector module cable.                     |
|                                | The injector module may have failed.                                | Replace the injector module (INOMAX delivery will be interrupted).                   |
|                                | The injector module electrical cable may have failed.               | 1. Replace the injector module cable (INOMAX delivery will be interrupted).          |
|                                |                                                                      | 2. Contact Customer Support.                                                         |

1. Manually ventilate patient with the INOblender or turn integrated pneumatic backup delivery ON.
## High Priority Alarms

All of these actions can be performed while delivering INOMAX to the patient:

<table>
<thead>
<tr>
<th>Alarm</th>
<th>Possible Cause</th>
<th>Recommended Action</th>
</tr>
</thead>
</table>
| 13. Low Battery | Battery is running low. | Up to 30 minutes of battery life remains following low battery alarm activation.  
1. Check AC power connection at wall and/or back of INOmax DSIR Plus (green indicator light should be illuminated).  
2. Contact Customer Support. |
| 14. Low Cylinder Pressure | Cylinder valve is closed. | Confirm INOMAX cylinder valve is fully open.  
1. The regulator hose may not be connected. | Confirm correct INOMAX regulator hose is connected.  
2. The NO cylinder supply may be low. | Check the INOMAX regulator gauge pressure, replace the cylinder if necessary.  
3. INOmax DS<sub>IR</sub> Plus has developed an internal leak. | 1. Bypass the INOmax DS<sub>IR</sub> Plus and connect the INOblender directly to the INOMAX regulator.  
2. Contact Customer Support. |
| 15. Service Required | The INOmax DS<sub>IR</sub> Plus has failed. | Manually ventilate patient with the INOblender or turn integrated pneumatic backup delivery ON.  
1. Remove from service.  
2. Contact Customer Support. |

### Manual Delivery Available

Utilize the INOblender or Integrated pneumatic backup delivery.
# Low Priority Alarms

All of these actions can be performed while delivering INOMAX to the patient:

<table>
<thead>
<tr>
<th>Alarm</th>
<th>Possible Cause</th>
<th>Recommended Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>16. Backup On</td>
<td>The backup mode has been turned ON.</td>
<td>1. Correct the reason for initiating integrated pneumatic backup delivery.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. Turn integrated pneumatic backup delivery OFF and confirm set NO dose and monitor</td>
</tr>
<tr>
<td></td>
<td></td>
<td>alarm settings have been restored.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3. Contact Customer Support.</td>
</tr>
<tr>
<td>17. Under Delivery</td>
<td>INOMAX delivery is less than 50% of set dose.</td>
<td>1. Check calculated delivery graph on the settings screen.</td>
</tr>
<tr>
<td></td>
<td>[CALCULATED dose is &lt;50% of set dose] AND (CALCULATED dose is &lt; set dose - 5 PPM)] for 12</td>
<td>2. Check circuit flow rate is 2-120 L/min.</td>
</tr>
<tr>
<td></td>
<td>consecutive seconds.</td>
<td>3. Check if dose is &gt;60 ppm with breathing circuit flow &gt;60 L/min.</td>
</tr>
<tr>
<td>18. Failed NO Sensor</td>
<td>NO calibration may have drifted out of specification.</td>
<td>1. Perform low calibration.</td>
</tr>
<tr>
<td></td>
<td>Delivery of INOMAX continues during this alarm.</td>
<td>2. Perform high NO calibration.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3. Contact Customer Support.</td>
</tr>
<tr>
<td>19. Failed NO₂ Sensor</td>
<td>NO₂ calibration may have drifted out of specification.</td>
<td>1. Perform low calibration.</td>
</tr>
<tr>
<td></td>
<td>Delivery of INOMAX continues during this alarm.</td>
<td>2. Perform high NO₂ calibration.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3. Contact Customer Support.</td>
</tr>
<tr>
<td>20. Failed O₂ Sensor</td>
<td>O₂ calibration may have drifted out of specification.</td>
<td>1. Perform low calibration.</td>
</tr>
<tr>
<td></td>
<td>Delivery of INOMAX continues during this alarm.</td>
<td>2. Perform high O₂ calibration.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3. Contact Customer Support.</td>
</tr>
</tbody>
</table>
# Low Priority Alarms

All of these actions can be performed while delivering INOMAX to the patient:

<table>
<thead>
<tr>
<th>Alarm</th>
<th>Possible Cause</th>
<th>Recommended Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>21. Monitoring Failure</td>
<td>Delivery of INOMAX continues during this alarm.</td>
<td>1. Remove from service.</td>
</tr>
<tr>
<td></td>
<td>Monitoring (sample) system has failed.</td>
<td>2. Contact Customer Support.</td>
</tr>
<tr>
<td></td>
<td>The sample line may be blocked.</td>
<td>Confirm patient gas sample line is not occluded, replace if necessary.</td>
</tr>
<tr>
<td></td>
<td>The water separator cartridge may be blocked.</td>
<td>Replace water separator cartridge.</td>
</tr>
<tr>
<td></td>
<td>The 1.0 micron disk filter may be blocked.</td>
<td>1. Replace the 1.0 micron disk filter.</td>
</tr>
<tr>
<td></td>
<td>Two cylinder valves are open.</td>
<td>2. Contact Customer Support.</td>
</tr>
<tr>
<td>23. Two Cylinders Open</td>
<td>Two cylinder valves are open.</td>
<td>1. Close one INOMAX cylinder valve and depressurize regulator.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. Contact Customer Support.</td>
</tr>
<tr>
<td>24. Water Bottle Full</td>
<td>The water bottle is full.</td>
<td>Empty water bottle.</td>
</tr>
<tr>
<td></td>
<td>Water bottle is empty but the message remains in the alarm message box.</td>
<td>1. Remove water bottle and wipe off optical sensor.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. Contact Customer Support.</td>
</tr>
<tr>
<td>25. Low Calibration Failed</td>
<td>Zeroing valve has failed.</td>
<td>1. Repeat manual low calibration.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Wait for low calibration to complete (approximately three minutes).</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. Contact Customer Support.</td>
</tr>
<tr>
<td>Indicator</td>
<td>Possible Cause</td>
<td>Recommended Action</td>
</tr>
<tr>
<td>-----------</td>
<td>---------------</td>
<td>--------------------</td>
</tr>
<tr>
<td>27. Set Dose is Zero, Please Close Cylinder Valve</td>
<td>The set dose has been set to zero and the INOMAX cylinder valve is still open.</td>
<td>Close the INOMAX cylinder valve and depressurize regulator.</td>
</tr>
</tbody>
</table>
| 28. NO Delivery Button Inactive | Device does not recognize an INOMAX cylinder, the dose knob will be greyed out and it will not allow the user to set an initial INOMAX dose. | 1. Load INOMAX cylinder on to the cart.  
2. Remove any obstruction between the INOmeter and the INOmax DSIR Plus cart cover.  
3. Move the interfering light source.  
4. Move the INOmax DSIR Plus cart to reduce the high intensity light in the area of the INOmeter.  
5. Shield the INOmeter from the suspect light source.  
6. Replace cylinder. |
| \( IR \text{ cable is not connected to the back of the INOmax DS}_{\text{IR}} \text{ Plus.} \) | Verify the IR cable is connected to the back of the INOmax DS\(_{\text{IR}}\) Plus. |  |
| \( \text{Integrated pneumatic backup switch is ON.} \) | 1. Correct the reason for initiating integrated pneumatic backup delivery.  
2. Turn integrated pneumatic backup delivery OFF and confirm set NO dose has been restored. |  |
If the system fails to operate properly:
1. Check the patient condition and take appropriate action.
2. Use the INOblender (see INOblender Operation Manual) or backup if necessary.
3. Verify that the system is set up as detailed in 3 Patient Application.
4. Find a symptom or alarm condition in the troubleshooting table which best describes the problem and follow the recommended actions to resolve the problem.

If the problem cannot be corrected:
Contact the Authorized Representative listed on the back cover of the operation manual.

If the INOmax DSIR Plus must be returned for servicing:
Contact the Authorized Representative listed on the back cover of the operation manual.

**WARNING:**
- If an alarm occurs, safeguard the patient first before troubleshooting or repair procedures.
- Use caution when troubleshooting the INOmax DSIR Plus while in use for a patient.

To activate on-screen alarm help, press the Alarm Help Button next to the Alarm Silence button.
**WARNING:** Set the INOmax DS\textsubscript{IR} Plus alarm thresholds for the current patient conditions to monitor any inadvertent changes in treatment.

**Caution:**
- Any alarm setpoint adjustments made will not be maintained when system power is cycled.
- Default values will be used following a complete power loss (no AC main power and depleted battery).

**General alarm information**
A listing of alarm messages is provided at the end of this section.
All alarms have audible tones and visual messages.

| In the event of a total power failure or a main alarm speaker failure, a secondary audible alarm circuit activates, providing a continuous buzzing tone that cannot be silenced (see page 5-1, "Continuous Audible Tone"). |

**Note:** Status information will not be displayed during alarm conditions. Once the alarm clears the status information will be displayed.

**High and low-priority alarms**
The INOmax DS\textsubscript{IR} Plus has both high and low-alarm priorities.
High-priority alarms are accompanied with a red flashing Alarm Silence button.
Low-priority alarm conditions will display a continuous yellow Alarm Silence button.
High and low-priority alarm messages are displayed in fields one and two (see Figure 5-1) with the most recent message shown in field one.
Field 2 is used for status information such as “Running on Battery” and “Patient Info. Incomplete”.

![Figure 5-1 Text Message Area Showing Fields one and two.](image)

The following table provides the audible alarm tone information for high and low-priority alarms.

<table>
<thead>
<tr>
<th>Frequency</th>
<th>Description</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>High Priority</td>
<td>400 Hz</td>
<td>10-pulse group</td>
</tr>
<tr>
<td>Low Priority</td>
<td>400 Hz</td>
<td>1 pulse</td>
</tr>
</tbody>
</table>

**Alarm silencing**
Pushing the Alarm Silence button will silence high-priority alarms for 120 seconds (time will count down to zero). When a new alarm condition occurs, the audible alarm becomes active again.

A low-priority alarm event is permanently silenced when the Alarm Silence button is pressed. When a new low-alarm condition occurs, the audible alarm becomes active again.

Alarm messages remain displayed during the alarm silence period as long as the alarm condition is active.

![Figure 5-2 Alarm Active Screen](image)

1. Alarm Silence Button
2. Alarm Help Button
3. Violated Monitored Value Limit

Figure 5-2 Alarm Active Screen

To activate on-screen alarm help, press the Alarm Help Button next to the Alarm Silence button.
User adjustable monitor alarms

Caution: Do not set upper and lower alarm limits to extreme values, as this could reduce the effectiveness of the monitoring alarm system.

Monitor alarm delay active indicator

Monitor alarms for O₂, NO₂, and NO will be inactive anytime the Monitor Alarm Delay Active indicator is displayed. This delay only affects the monitor alarms, all other alarms remain active.

The Monitor Alarm Delay Active indicator will be displayed for two minutes:

- Upon exit from the calibration screen (whether or not a calibration was actually performed)
- Following an automatic low calibration
- Following completion of an auto purge

The O₂, NO₂, and NO monitors have user adjustable alarm settings that are displayed to the side of the monitored value.

- The top button is the high-level alarm setting, and the lower button is the low-level alarm setting (see Figure 5-2).
- A low-alarm limit cannot be set above the high-limit setting.

When an alarm occurs for a monitored value, the violated alarm setting button flashes red (see #3 in Figure 5-2).

- To adjust an alarm level to a new value, press the selected alarm level button on the touch screen, rotate the control wheel to adjust to the new level and then confirm by pushing the control wheel or the selected alarm level button again.
- If the new alarm level is not confirmed within 20 seconds, the alarm level defaults back to its previous value.

The adjustment ranges for these alarm settings are shown in the table below.

<table>
<thead>
<tr>
<th>Alarm</th>
<th>Adjustment</th>
<th>Increments</th>
<th>Default</th>
<th>Priority</th>
</tr>
</thead>
<tbody>
<tr>
<td>High NO (ppm)</td>
<td>1 to 100</td>
<td>NO*0.0-1.0 by 0.1 ppm *1-99 by 1 ppm</td>
<td>Initially 90, then 50% above the initial set dose*</td>
<td>High</td>
</tr>
<tr>
<td>Low NO (ppm)</td>
<td>0 to 99</td>
<td></td>
<td>OFF (--) then 50% below the initial set dose ±</td>
<td>High</td>
</tr>
<tr>
<td>High NO₂ (ppm)</td>
<td>0 to 5</td>
<td>NO₂ by 0.1 ppm</td>
<td>3</td>
<td>High</td>
</tr>
<tr>
<td>High O₂ (% v/v)</td>
<td>21 to 100 Then OFF (--)</td>
<td>O₂ by 1%</td>
<td>-- (OFF)</td>
<td>High</td>
</tr>
<tr>
<td>Low O₂ (% v/v)</td>
<td>18 to 99</td>
<td></td>
<td>21 %</td>
<td>High</td>
</tr>
</tbody>
</table>

The first time a dose is set from zero, the upper and lower NO alarm limits are set 50% above and 50% below the set dose.

* Dose settings < 3 ppm will result in a high alarm setting of 5 ppm; otherwise rounded up to the nearest ppm and limited to 90 ppm maximum.

± Rounded down to the nearest ppm.
Alarm History

When an alarm condition has been resolved, the alarm message is no longer displayed on the main screen.

The recent alarms can be seen by pressing the Recent Alarms button.

The Recent Alarms button is present and displayed as a “double-bell” when there are no active alarms and any previously resolved alarms have not been cleared.

The alarms are displayed in chronological order, with most recent at the top (for example, Figure 5-4 the recent alarm conditions that have occurred).

Note:
- Alarm history dates and times are displayed per user-set off-set time (see Time Adjust setting, page 1-14).
- Alarm conditions lasting less than one second may not display in the alarm area of the user screen, but will post to the alarm log and alarm history.

Press the clear Recent Alarms button to clear the recent alarms and return to the main screen.

To return to the main screen without clearing the recent alarm history, press the EXIT or the Return to Previous Menu button.

If no action is taken, the system will return to the main screen after 30 seconds.
A complete list of all alarms that have occurred since the INOmax DSIR Plus has been turned ON can be viewed by pressing the Alarm History button on the menu screen (see figure 5-5).

**Priority**
A yellow dot signifies a low priority alarm, and a red dot signifies a high priority alarm.

Press the EXIT button or the Return to Previous Menu button to return to the menu screen.
(Intentionally left blank)
INOmax DS<sub>IR</sub> Plus

6/ Calibration
6/ Calibration
6/ Calibration

**WARNING:** Do not change any sensor while delivering NO to a patient.

**Note:** During any calibration process, all other alarms remain active while monitoring alarms are disabled.

To access the calibration menu:
Press the menu button on the main screen to enter the menu screen (second menu level).

The lower part of the screen displays the low calibration (Low Cal) and high calibration (High Cal) buttons.

- Select the Low Cal button to start a low calibration. The date and time the most recent low calibration occurred is displayed below the Low Cal button.
- Select the High Cal button to enter the high calibration screen. The earliest sensor high calibration due date is displayed above the High Cal button.

If the date is flashing, it signifies a calibration is past due.

**Note:** To return to the main screen, press the return to the previous level button in the top right of the screen.

Instructions for completing a low and high calibration are on the following pages.
Low Calibration

The low calibration of the monitor sensors uses room air to calibrate all three sensors at the same time. The system automatically draws in room air from an inlet port behind the water bottle, not the sample line. A low calibration is completed automatically when the INOmax DSIR Plus is turned ON and during the following conditions:

- At 3, 6, and 12 hour intervals following each dose change.
- Every 12 hours as long as the dose is not changed.
- Every 24 hours when the INOmax DSIR Plus is turned ON and the dose is set to zero.
- If the low calibration is cancelled after boot-up, the device will reattempt again every 15 minutes until successful.
- If an automatic low calibration fails, it will reattempt the calibration a second time. If it fails the second time, a Low Calibration failed alarm is raised. The next automatic calibration will occur at the next interval. For example, if the three hour calibration was just completed, the next calibration will occur in six hours.

**Note:** A fifteen minute period of time with no user screen interactions is required before an automatic low calibration will initiate.

1. From the menu screen (second menu level), press the Low Cal button to initiate a low calibration.

2. The calibration will take approximately three minutes, during which a progress bar for each sensor indicates the progress. A Low Calibration indicator will display in the alarm area of the screen during the low calibration. To cancel a low calibration, press the CANCEL LOW CAL button.
If the low calibration was unsuccessful, the INOmax DSIR Plus will automatically attempt another low calibration. If the second low calibration attempt fails, the alarm area will display a Low Calibration Failed alarm.

- Attempt a manual low calibration.

If a sensor has failed, the display will indicate the failed sensor symbol in the monitoring area of that sensor. (Press the Alarm Help button for on-screen alarm help or see Section 5/ Alarms and Troubleshooting.)
Oxygen Sensor High Calibration

**Caution:** Never connect the sample line directly to a high pressure gas source (greater than 150 cmH₂O); this could damage the sampling system.

**Note:** Complete a low calibration (see Low Calibration section) prior to completing the high calibration.

The oxygen high calibration requires a user supplied source of 100% oxygen.

1. From the high calibration screen (third menu level), press the 100% O₂ button to initiate the O₂ high calibration wizard.

   ![High Calibration Screen]

   ![High Calibration Wizard]

   **Note:** If the date is flashing, it signifies the calibration is past due.

2. Assemble connectors into a calibration setup and set oxygen flow to five L/min.

   (1) 100% O₂ Source
   (2) O₂ Tubing
   (3) 15M X 4.5 mm I.D Adapter
   (4) Gas Sample Tee
   (5) Patient Gas Sample Line with Nafion

3a. To continue the high calibration wizard press the NEXT button.

3b. To start the high calibration without using the wizard, press the START CAL button.

3c. To exit the O₂ high calibration wizard, press the CANCEL button.

   **Note:** Cap off sample tee if patient gas sample line is removed from patient breathing circuit.
4. After reaching step 5 of the calibration wizard, the calibration will take approximately three minutes, during which a progress bar for the O₂ sensor indicates progress.

**Note:** If the CANCEL button is pressed during the high O₂ calibration process, the calibration will be cancelled and the user will be returned to the high calibration screen.

5. When the O₂ high calibration is successful, a single tone will be heard and the progress bar will turn green. Press the NEXT button to return to the high calibration screen.

**Note:** The monitor displays should indicate approximately 100% O₂, 0.0 ppm NO₂ and 0.0 ppm NO.

Disconnect the sample line from the calibration setup and turn OFF the O₂ flowmeter.

**Note:** Reconnect patient gas sample line if previously removed from patient breathing circuit.

A Monitor Alarm Delay Active indicator will occur anytime the user exits the calibration screen, even if a calibration was not initiated. All other alarms are still active (see Section 5/ Alarms and Troubleshooting for additional information).

When using the high calibration wizard, if the BACK button is pressed during the O₂ high calibration process, the progress bar will turn yellow, signifying a cancelled calibration.

- Press the NEXT button to start the high calibration wizard.
- Press the START CAL button to start the calibration again.
- Press the CANCEL button to exit the O₂ high calibration screen.
If the O₂ sensor has failed, the display indicates the failed sensor symbol in the monitoring area of that sensor.

Repeat calibration, (press the Alarm Help button for on-screen alarm help or see Section 5/ Alarms and Troubleshooting.)

If the calibration was unsuccessful, the O₂ progress bar will turn red.

- Attempt another calibration.

Note: To repeat the O₂ high calibration, press the START CAL button at the bottom of the screen.
NO Sensor High Calibration

**Caution:**
- When performing a high calibration, make sure to select the correct calibration gas (INOcal NO, 45 ppm, P/N BOM-COM-0150) and confirm the expiration date before using.
- Never connect the sample line directly to a high pressure gas source (greater than 150 cmH₂O); this could damage the sampling system.

**Note:** Complete a low calibration (see Low Calibration section) prior to completing the high calibration.

**INOcal calibration gas kit sample tubing**
When using the calibration tubing kit (P/N 50107), which is supplied with the INOcal regulator kit (P/N 10036), ensure that the beige one-way valve supplied with the tubing is installed and oriented as indicated in Figure 6-1.

**Caution:** An incorrectly installed one-way valve can lead to over-pressurization of the sampling system. A leak in the calibration tubing kit (PN 50107) attached to the calibration cylinder regulator can result in displayed NO values greater than the set dose value after passing a low and high calibration successfully. This can be caused by aging of the calibration tubing. The calibration tubing kit should be replaced under the following circumstances:
- The tubing is discolored or stiff.
- There is a crack or break in the tubing.

1. Remove cylinder cap and inspect for damaged threads and contaminants.
2. Check seal on regulator. Verify it is correctly in place and undamaged.
3. Attach regulator to cylinder (Turn regulator nut counter-clockwise to tighten).
4. **NOTE:** Tubing adapter will be required if regulator outlet diameter measures 0.24 in. (6.10 mm).
5. Attach tubing kit to regulator outlet.

**NOTE:** Confirm water bottle, water separator cartridge and patient gas sample line are in place.
6. Open the INOcal cylinder valve (turn counterclockwise) to flow gas.

6a. If the pressure is in the red or black zone (0-25 psig) select another INOcal cylinder.

7. Attach tubing kit to patient gas sample line.

Note: Cap off sample tee if patient gas sample line is removed from patient breathing circuit.

8. From the high calibration screen (third menu level), press the 45 ppm NO button to initiate the NO high calibration.

Note: If the date is flashing on the high calibration screen, the calibration is past due.

9a. To continue the high calibration wizard, press the NEXT button.

9b. To start the high calibration without using the wizard, press the START CAL button.

9c. To exit the NO high calibration, press the CANCEL button.
When the NO high calibration is successful, a single tone will be heard and the progress bar will turn green. Press the NEXT button to return to the high calibration screen.

Note: The monitor displays should indicate approximately 0.0% O₂, 0.0 ppm NO₂ and 45 ppm NO.

After reaching step 7 of the high calibration wizard, the calibration will take approximately three minutes, during which a progress bar for the sensor indicates progress.

Note: If the CANCEL button is pressed during the NO high calibration process, the calibration will be cancelled and the user will be returned to the high calibration screen.

When the NO high calibration is successful, a single tone will be heard and the progress bar will turn green. Press the NEXT button to return to the high calibration screen.

Note: If the CANCEL button is pressed during the NO high calibration process, the calibration will be cancelled and the user will be returned to the high calibration screen.

10. Disconnect the patient gas sample line from the calibration tubing and close the INOcal cylinder valve.

Note: Reconnect patient gas sample line if previously removed from patient breathing circuit.

A Monitor Alarm Delay Active indicator will occur anytime the user exits the calibration screen, even if a calibration was not initiated. All other alarms are still active (see Section 5/ Alarms and Troubleshooting for additional information).
If the NO sensor has failed, the display indicates the failed sensor symbol in the monitoring area of that sensor.

Repeat calibration. (Press the Alarm Help button for on-screen alarm help or see Section 5/ Alarms and Troubleshooting.)
NO₂ Sensor High Calibration

**Caution:**
- When performing a high calibration, make sure to select the correct calibration gas (INOcal NO₂, 10 ppm, P/N BOM-COM-0162) and confirm the expiration date before using.
- Never connect the sample line directly to a high pressure gas source (greater than 150 cm H₂O); this could damage the sampling system.

**Note:** Complete a low calibration (see Low Calibration section) prior to completing the high calibration.

**INOcal calibration gas kit sample tubing**

When using the calibration tubing kit (P/N 50107), which is supplied with the INOcal regulator kit (P/N 10036), ensure that the beige one-way valve supplied with the tubing is installed and oriented as indicated in Figure 6-2.

**Caution:**
An incorrectly installed one-way valve can lead to overpressurization of the sampling system. A leak in the calibration tubing kit (P/N 50107) attached to the calibration cylinder regulator can result in displayed NO₂ values greater than the set dose value after passing a low and high calibration successfully. This can be caused by aging of the calibration tubing. The calibration tubing kit should be replaced under the following circumstances:
- The tubing is discolored or stiff.
- There is a crack or break in the tubing.

1. Remove cylinder cap and inspect for damaged threads and contaminants.
2. Check seal on regulator. Verify it is correctly in place and undamaged.
3. Attach regulator to cylinder (Turn regulator nut counter-clockwise to tighten).
4. NOTE: Tubing adapter will be required if regulator outlet diameter measures 0.24 in. (6.10 mm).
5. Attach tubing kit to regulator outlet.

**Note:** Confirm water bottle, water separator cartridge and patient gas sample line are in place.

---

**Figure 6-2**

**Diagram:**
- One-way Valve
- Gas Flow
- Flow Arrows
- Cylinder Cap
- Regulator
- Water Bottle
- Water Separator Cartridge
- Patient Gas Sample Line
6. Open the INOcal cylinder valve (turn counter-clockwise) to flow gas.

6a. If the pressure is in the red or black zone (0-25 psig) select another INOcal cylinder.

7. Attach tubing kit to patient gas sample line.

Note: Cap off sample tee if patient gas sample line is removed from patient breathing circuit.

8. From the high calibration screen (third menu level), press the 10 ppm NO₂ button to initiate the NO₂ high calibration.

Note: If the date is flashing on the high calibration screen, the calibration is past due.

9a. To continue the high calibration wizard, press the NEXT button.

9b. To start the high calibration without using the wizard, press the START CAL button.

9c. To exit the NO₂ high calibration wizard, press the CANCEL button.
After reaching step 7 of the high calibration wizard, the calibration will take approximately three minutes, during which a progress bar for the sensor indicates progress.

**Note:** If the CANCEL button is pressed during the NO₂ high calibration process, the calibration will be cancelled and the user will be returned to the high calibration screen.

When the NO₂ high calibration is successful, a single tone will be heard and the progress bar will turn green. Press the NEXT button to return to the high calibration screen.

**Note:** The monitor displays should indicate approximately 21% O₂, 10 ppm NO₂ and 0.0 ppm NO.

10. Disconnect the patient gas sample line from the calibration tubing and close the INOcal cylinder valve.

**Note:** Reconnect patient gas sample line if previously removed from patient breathing circuit.

A Monitor Alarm Delay Active indicator will occur anytime the user exits the calibration screen, even if a calibration was not initiated. All other alarms are still active (see Section 5/ Alarms and Troubleshooting for additional information).
If the calibration was unsuccessful, the NO₂ progress bar will turn red.

- Attempt another calibration.

Note: To repeat the NO₂ high calibration, press the START CAL button at the bottom of the screen.

When using the high calibration wizard, if the BACK button is pressed during the NO₂ high calibration process, the progress bar will turn yellow, signifying a cancelled calibration.

- Press the NEXT button to start the high calibration wizard.
- Press the START CAL button to start the calibration again.
- Press the CANCEL button to exit the NO₂ high calibration screen.

If the NO₂ sensor has failed the display indicates the failed sensor symbol in the monitoring area of that sensor.

Repeat calibration. (Press the alarm help button for on-screen alarm help or see Section 5/ Alarms and Troubleshooting.)
INOmax DS$_{IR}$ Plus

7/ Maintenance
INOMax DS_{IR}^{®} Plus

7/ Maintenance
7/ Maintenance

Unpacking the INOmax DSIR Plus

Note: Following unpacking and prior to the first use:
- Remove any protective caps from the connectors and ports on the INOmax DSIR Plus.
- Ensure the INOmax DSIR Plus is on a flat surface or is fixed securely to a cart or transport sled.

Caution: Do not sterilize or disinfect with the power connected.

Note: The INOmax DSIR Plus does not contain any user repairable parts.

User Maintenance Schedule

<table>
<thead>
<tr>
<th>Frequency</th>
<th>Maintenance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Daily (during patient use)</td>
<td>1. Check the INOMAX cylinder pressure. A cylinder with less than 14 bar (200 psig) should be replaced.</td>
</tr>
<tr>
<td></td>
<td>2. Empty the water bottle as needed.</td>
</tr>
<tr>
<td>Start of each patient</td>
<td>Must perform the Pre-Use Procedure.</td>
</tr>
<tr>
<td>Between each patient</td>
<td>1. Sterilize and/or disinfect the injector module.</td>
</tr>
<tr>
<td></td>
<td>2. Clean water bottle.</td>
</tr>
<tr>
<td></td>
<td>3. Replace the single patient-use items.</td>
</tr>
<tr>
<td></td>
<td>4. Make sure that the delivery system power cord is always plugged into an emergency-power-backed electrical outlet.</td>
</tr>
<tr>
<td></td>
<td>5. Make sure the connectors, hoses and cables are in good condition.</td>
</tr>
<tr>
<td>Monthly</td>
<td>1. Perform a low and a high calibration of NO, NO₂ and O₂.</td>
</tr>
<tr>
<td></td>
<td><strong>Note:</strong> A flashing date above the high sensor calibration button signifies a high calibration is past due.</td>
</tr>
<tr>
<td></td>
<td>2. Check INOMAX regulators for leaks.</td>
</tr>
</tbody>
</table>
Cleaning the INOmax DS$_{IR}$ Plus

**Caution:**
- Do not autoclave or gas sterilize the INOmax DS$_{IR}$ Plus.
- Do not clean with the power connected and the INOmax DS$_{IR}$ Plus turned ON.
- Be sure that the INOmax DS$_{IR}$ Plus is completely dry before using.
- Do not saturate the INOmax DS$_{IR}$ Plus with excessive solution. Liquid may flow into the system and damage internal components.
- Do not use organic, petroleum based solvents, glass cleaners, acetone or other harsh cleaning agents.
- Do not use abrasive cleaning agents (such as steel wool, silver polish or cleaner).
- Do not touch or rub the display panel with abrasive cleaning compounds or anything which can scratch the panel.
- Do not use organic solvents to clean the display panel.

**Cleaning Procedure**

**Caution:** Apply cleaning agent to a cloth before application; do not spray directly on the delivery system to prevent pooling and direct contact with electrical connections which can cause damage over time.

**External surfaces and the Display panel**
- Disconnect the power cord from the wall outlet and turn the INOmax DS$_{IR}$ Plus OFF before cleaning.
- Clean the outer surface of the INOmax DS$_{IR}$ Plus with a soft cloth dampened in a mild soap and water solution, isopropyl alcohol (70%) or with one of the following cleaning agents while following the manufacturer’s recommendations.

<table>
<thead>
<tr>
<th>Cleaning Agent</th>
<th>Active Ingredients</th>
</tr>
</thead>
</table>
| **Precise Hospital Foam Cleaner Disinfectant** by Caltech Industries | o-Phenylphenol < 0.37%  
Other ingredients 99.63% |
| **Pure Green 24** by Pure Green, LLC | SDC – silver ions 0.003%  
Citric acid 4.84%  
Other ingredients 95.157% |
<table>
<thead>
<tr>
<th>Cleaning Agent</th>
<th>Active Ingredients</th>
</tr>
</thead>
</table>
| **PDI Super Sani Cloth** by PDI | n-Alkyl (68% C12, 32% C14) dimethyl ethylbenzyl ammonium chlorides 0.25%  
                                    n-Alkyl (60% C14, 30% C16, 5% C12, 5% C18) dimethyl benzyl ammonium chlorides 0.25%  
                                    Isopropyl alcohol 55%  
                                    Inert ingredients 44.50% |
| **Sani Cloth HB** by PDI       | n-Alkyl (60% C14, 30% C16, 5% C12, 5% C18) dimethyl benzyl ammonium chlorides 0.07%  
                                    n-Alkyl (68% C12, 32% C14) dimethyl ethylbenzyl ammonium chlorides 0.07%  
                                    Inert ingredients 99.86% |
| **Asepti-HB** by Ecolab Inc.   | n-Alkyl (60% C14, 30% C16, 5% C12, 5% C18) dimethyl benzyl ammonium chlorides 0.07%  
                                    n-Alkyl (68% C12, 32% C14) dimethyl ethylbenzyl ammonium chlorides 0.07%  
                                    Inert ingredients 99.86% |
| **Cavicide and CaviWipes** by Metrex | Diisobutylphenoxyethoxyethyl dimethyl benzyl ammonium chloride 0.28%  
                                      Isopropyl alcohol 17.2%  
                                      Inert ingredients 82.52% |

**Cleaning Water Bottle**

**Caution:** If alcohol is used to clean water bottle, make sure alcohol is completely evaporated before placing back onto sample block.

- Alcohol vapors will cause NO₂ sensor to read high (as much as six ppm) and NO sensor to read low (approximately 0.5 to one ppm).
- This is a transient response and will stop once alcohol vapors dissipate (bottle dries out).

**Procedure**

- Clean water bottle with a soft cloth dampened in a mild soap and water solution or with isopropyl alcohol (70%).
- Allow water bottle to air dry.

**Bioquell Hydrogen Peroxide Sterilant**

Bioquell Hydrogen Peroxide Sterilant and hydrogen peroxide vapor generators are regulated by the US Environmental Protection Agency (EPA) as pesticide chemicals in accordance with Federal Insecticide, Fungicide, and Rodenticide Act (FIFRA). Use of these products in cleaning/disinfection processes have not been validated with the INOmax DSIR Plus. Do not use these products to decontaminate the INOmax DSIR Plus or any ancillary products used with the INOmax DSIR Plus.
Cleaning the INOmeter

**Caution:**

- Apply cleaning agent to a cloth before application; do not spray directly on the INOmeter. It is important to prevent pooling and direct contact with electrical connections, which can cause damage over time.
- Do not autoclave or gas sterilize the INOmeter.
- Be sure that the INOmeter is completely dry before using.
- Do not saturate the INOmeter with excessive solution. Liquid may flow into the device and damage internal components.
- Do not use organic, petroleum-based solvents, glass cleaners, acetone or other harsh cleaning agents.
- Do not use abrasive cleaning agents (such as steel wool, silver polish or cleaner).

External surfaces and the Display

- Clean the outer surface of the INOmeter with a soft cloth dampened in a mild soap and water solution, isopropyl alcohol (70%) or with one of the cleaning agents (see cleaning agent list above) while following the manufacturer's recommendations.
Injector Module Sterilizing and/or Disinfecting

**WARNING:** If the injector module was used in the wet/humidified part of the breathing circuit, it should be sterilized between each patient use.

**Caution:** Remove the injector module cable prior to sterilizing or disinfecting the injector module.

If the injector module has been used in the dry part of the breathing circuit, the injector module should be sterilized and/or disinfected in 70% ethyl alcohol after each patient use.

**Autoclave Sterilizing the Injector Module**

1. *Disconnect the electrical cable and the injector tube before autoclaving.*
2. Autoclave the injector module at 134° C for three minutes at 1.85 bar (27 psig).
3. After sterilization, examine the parts.
4. Replace any broken, worn, distorted or discolored parts.

**Disinfecting the Injector Module**

1. Fill a container with 70% ethyl alcohol.
2. Totally submerge the injector module in the 70% ethyl alcohol for at least 30 minutes. If debris is noticed on the hot wire sensor, gently agitate the module in the alcohol bath.
3. Remove the injector module from the liquid and drain the excess alcohol from the module’s electrical connector, injector port and inside flowmeter.
   
   **Note:** If rinsing is required, use a separate bath filled with distilled water.
4. Allow liquid to evaporate completely before using the injector module.

   **Note:**
   - Do not insert anything into the injector module throat in an effort to remove contamination or to dry.
   - If lint fibers remain wrapped around the hot wire sensor after drying, do not use the module. Remove it from service and contact Customer Support.

**Note:** Patient circuit adapters, patient gas sample line, injector module tubing and water separator cartridge are single-patient use items. Do not sterilize them. Dispose of all single-patient use items in accordance with universal precautions for contamination.
Replacing the O₂, NO and NO₂ Sensors

**WARNING:** Handle and dispose of sensors according to facility biohazard policies. Do not incinerate.

1. Remove the rear sensor cover by turning the two screws counterclockwise until loose (see Figure 7-1).

2. Grasp the sensor to be replaced on both sides and gently pull it from its socket (see Figure 7-2).

   - The shorting wire must be removed from the NO₂ sensor before replacing (see Figure 7-6).
   - Make sure all of the sensor O-rings are present and seated properly.

3. To install replacement NO or NO₂ sensor, align the pins with the socket and press it into place (see Figure 7-3).

4. To install O₂ sensor, remove the shorting button (see Figure 7-4) and insert the contact end (open end with three gold rings) into recess until it seats (no specific orientation is necessary).
5. Replace the sensor cover and tighten the two screws clockwise (see Figure 7-5).
6. Perform a low and high calibration for the sensor before returning the system to use.

<table>
<thead>
<tr>
<th>Note:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Newly Installed Sensor</td>
</tr>
<tr>
<td>O₂ and NO₂</td>
</tr>
<tr>
<td>NO</td>
</tr>
</tbody>
</table>

Insufficient conditioning will result in inaccurate gas readings.

Figure 7-5
Figure 7-6
NO₂ Shorting Wire
Replacing the Water Separator Cartridge

The disposable water separator cartridge on the rear of the water bottle housing protects the monitoring system from moisture and other contaminants.

To replace the Water Separator Cartridge:

1. Grasp the cartridge on the back and top edge and gently pull it up and out of the dovetail slot in the sampling block (see Figure 7-7).

2. Discard the used cartridge in a receptacle designated for medical wastes.

3. To replace the cartridge, line it up with the dovetail slot and push it into place until it seats properly.

4. Check for leaks by running the system, occluding the sample line until the sample line occlusion alarm message appears.

Replacing the CGA 626 tip on the INOMAX regulator

1. Disconnect the regulator from the INOMAX drug cylinder.

Note: Depressurize the INOMAX regulator by using the purge port on the back of the INOmax DSrPlus prior to removing from the cylinder valve.

2. Remove the old CGA 626 (for ISO see figure 7-9) tip by pulling on the tip and turning it counterclockwise (see Figure 7-8).

3. Ensure the threads are clean on the regulator tip (if required, use a lint free cloth).

4. Install the new tip:

   Flex the four prongs by squeezing two prongs at a time using only your fingers. This will help start the new tip into the threads. Turn the tip clockwise when threading the tip. When the tip is fully inserted, it should turn freely.
Replacing the O-ring on the ISO 5145 INOMAX regulator fitting

1. Disconnect the regulator from the INOMAX therapy gas cylinder.

Note: Depressurize the INOMAX regulator by using the purge port on the back of the INOmax DSIR Plus prior to removing from the cylinder valve.

2. Remove the old ISO O-ring by rolling it off its groove (see Figure 7-9).
3. Clean the connector tip (if required, using a lint free cloth).
4. Roll the new O-ring into its groove. When correctly installed, it should not be removable by turning it.

Figure 7-9

Caution: Do not use hard objects to remove the O-ring as they may damage the metal gland and cause a leak.
Cylinder Leak Check

If a leak is suspected during the high pressure leak test (see Section 2/ Automated Pre-Use Checkout; High Pressure Leak Test), the following steps can be taken to check for leaks (see Figure 7-10 for possible cylinder gas leak locations) in the INOMAX regulator or INOMAX cylinder.

1. Confirm that INOMAX regulator is connected to cylinder valve outlet (hand tighten only), cylinder valve is open and that the cylinder has more than 200 psig.

2. Apply soapy water to points #1, #2, #3, #5 and #6 (see Figure 7-10); if bubbles form, there is a leak.

3. If there are no bubbles, the leak may be inside the INOMAX DSIR Plus and cannot be repaired. Replace the INOMAX DSIR Plus and contact Customer Support.

**Recommended actions should a leak be detected:**

1. A leak detected at points #1 and #2 may be corrected by tightening the INOMAX regulator hand wheel.
   a. If cylinder valve is open, close cylinder valve and tighten INOMAX regulator hand wheel.
   b. Open cylinder valve and reapply soapy water to points #1 and #2.
   c. If bubbles form, there is a leak.
   d. Remove INOMAX regulator and check for damage. For the CGA type regulator connector, check the white plastic tip on INOMAX regulator for chips or cracks. Replace if necessary. For the ISO type regulator connector, check that the O-ring is present and is not damaged. Replace if necessary (see replacing the tip/O-ring on the INOMAX regulator, Pages 7-8/7-9). Repeat step b.

2. If a leak is detected between the regulator body and regulator end cap (see point #3) replace INOMAX regulator and contact Customer Support.

3. A leak detected at the cylinder valve nut connection (see point #5) may not be repaired. Replace INOMAX cylinder and contact Customer Support.

4. A leak detected at the safety pressure release device (see point #6) may not be repaired. Replace INOMAX cylinder and contact Customer Support.

**Note:** Refer to hospital policies and procedures for dealing with leaking gas cylinders. Additional information regarding environmental effects can be found in the Section 1/ General Information.

---

**Figure 7-10**

1. Cylinder Valve Regulator Connection
2. INOMAX Regulator Hand Wheel Connection
3. Regulator End Cap Connection
4. Tamper Evident Tape
5. Valve Nut
6. Safety Pressure Release Device
Preventative Maintenance

**Perform the following maintenance task every year:**
- Replace O₂ and NO sensors.

**Perform the following maintenance task every two years:**
- Check battery.
- Check internal tubing.
- Replace sample system tubing and filters.
- Replace NO₂ sensor.

Equipotential grounding is the bonding of all conductive surfaces in the room together and to earth. This can be implemented in the patient care environment if it is crucial to keep all conductive surfaces at the same electrical potential or on the same ground plane.

If an equipotential grounding system is installed, the ground system should be tested per chapter four of NFPA 99.
## Parts and Accessories

**WARNING:** Only use parts/accessories designated for use with this system.

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<tr>
<th>Parts/Accessories</th>
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*Straight Shown.
Images for reference only.
INOmax DS<sub>IR</sub>® Plus

8/ Product Specifications
INOmax DSIR® Plus

8/ Product Specifications
8/ Product Specifications

**WARNING:**

- In the United States, the approved patient population for the INOmax DS\textsubscript{IR} Plus, as specified in the drug labeling for INOMAX\textsuperscript{®} (nitric oxide) for inhalation, is limited to term and near-term neonates with hypoxic respiratory failure. The INOmax DS\textsubscript{IR} Plus is not intended to be used in other patient populations.

- Outside of the United States, use of the INOmax DS\textsubscript{IR} Plus is limited to the use in accordance with INOMAX or INOflo, nitric oxide for inhalation prescribing information as established with the national health authority.

The INOmax DS\textsubscript{IR} Plus is designed to function in the parameter ranges listed in this section. Use outside of these ranges is not recommended.
## Ventilator Compatibility

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<td>Respiratory Rate:</td>
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The INOmax DSIR Plus is compatible with the ventilators listed below:

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<th>Pediatric/ Adult</th>
<th>Transport</th>
<th>High Frequency</th>
<th>Anesthesia</th>
<th>Nasal Continuous Positive Airway Pressure (CPAP)</th>
<th>High Flow Nasal Cannula</th>
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## Ventilators/Breathing Systems validated for use in the United States

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<th>High Frequency</th>
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<td>Anesthesia</td>
<td>Nasal Continuous Positive Airway Pressure (CPAP)</td>
<td>High Flow Nasal Cannula</td>
</tr>
<tr>
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</tr>
<tr>
<td>InfraSonics</td>
<td>Infant Star 500</td>
<td>•</td>
<td></td>
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<tr>
<td>InfraSonics</td>
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<tr>
<td>Maquet (formerly Siemens)</td>
<td>Servo i</td>
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<td>Nasal Cannula</td>
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<td>Newport</td>
<td>E360</td>
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<tr>
<td>Puritan Bennett</td>
<td>7200</td>
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<td>Puritan Bennett</td>
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<td>Respironics</td>
<td>Esprit</td>
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<td>Sichrist</td>
<td>IV-100B</td>
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<tr>
<td>Sensormedics</td>
<td>3100 A (standard and filtered circuits)</td>
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<tr>
<td>Sensormedics</td>
<td>3100B (standard and filtered circuits)</td>
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<tr>
<td>Smiths Medical</td>
<td>babyPAC 100</td>
<td>•</td>
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<tr>
<td>Smiths Medical</td>
<td>paraPAC Medic 200D</td>
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<tr>
<td>Smiths Medical</td>
<td>ventiPAC 200D</td>
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<tr>
<td>Teleflex Medical</td>
<td>Comfort Flo Humidification System</td>
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<tr>
<td>Vapotherm</td>
<td>2000i</td>
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<tr>
<td>Vapotherm</td>
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<tr>
<td>Manufacturer</td>
<td>Model</td>
<td>Neonatal</td>
<td>Pediatric/Adult</td>
<td>Transport</td>
<td>High Frequency</td>
<td>Anesthesia</td>
<td>Nasal Continuous Positive Airway Pressure (CPAP)</td>
<td>High Flow Nasal Cannula</td>
</tr>
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<tr>
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<td>Fabian +nCPAP Evolution</td>
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<td>●</td>
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<tr>
<td>Acutronic Medical Systems AG</td>
<td>Fabian HFO</td>
<td>●</td>
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<tr>
<td>Dräger</td>
<td>Evita Babylog VN500</td>
<td>●</td>
<td></td>
<td></td>
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<td>●</td>
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</tr>
<tr>
<td>Dräger</td>
<td>Zeus</td>
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<td></td>
<td></td>
<td>●</td>
</tr>
<tr>
<td>Heinen &amp; Löwenstein</td>
<td>Leoni+</td>
<td>●</td>
<td></td>
<td>●</td>
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</tr>
<tr>
<td>Infrasonics</td>
<td>Infant Star 950</td>
<td>●</td>
<td></td>
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<td>●</td>
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<tr>
<td>Metran</td>
<td>Humming HMX</td>
<td>●</td>
<td></td>
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<td>●</td>
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<tr>
<td>SLE Life Support</td>
<td>SLE 5000</td>
<td>●</td>
<td></td>
<td></td>
<td></td>
<td>●</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
NO Delivery

Set NO Range: 0.1 - 80 ppm (800 ppm cylinder)
0.1 - 40 ppm (400 ppm cylinder)
Set NO Resolution: 0.1 ppm from 0 to 1 ppm
1 ppm from 1 to 40 ppm
2 ppm from 40 to 80 ppm
Accuracy @ 20°C: ± 20% or 2 ppm, whichever is the greater
NO Inlet Pressure: 1.7 to 2.4 Bar (25 to 35 psig)
Maximum NO Supply Pressure: 2.4 Bar (35 psig)
NO Low Pressure Alarm: 1.6 Bar (23 psig) (nominal)
Max Circuit Pressure: 1.4 Bar (20 psig)
Breathing Circuit Gas Composition: Air / O₂ mixtures

Injector Module

Conical Connectors: Inlet, 22 mm female.
Outlet, 22 mm male and 15 mm female.
Autoclavability: Autoclavable at 134°C for 3 minutes at 1.85 bar (27 psig).
Maximum Pressure Drop: 1.5 cmH₂O at 60 L/min

Gas Monitoring

<table>
<thead>
<tr>
<th>Gas</th>
<th>Range</th>
<th>Resolution</th>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nitric Oxide</td>
<td>0 - 10 ppm</td>
<td>0.1</td>
<td>± (20% of reading + 0.5 ppm)</td>
</tr>
<tr>
<td></td>
<td>10 - 100 ppm</td>
<td>1</td>
<td>± (10% of reading + 0.5 ppm)</td>
</tr>
<tr>
<td>Nitrogen Dioxide</td>
<td>0 - 10 ppm</td>
<td>0.1</td>
<td>± (20% of reading or 0.5 ppm whichever is greater)</td>
</tr>
<tr>
<td>Oxygen</td>
<td>18 - 100 % v/v</td>
<td>1</td>
<td>± 3% v/v</td>
</tr>
</tbody>
</table>

Max Breathing Circuit Pressure: 150 cmH₂O
Calibration: Daily zero; span when needed
Rise Time: 30 seconds (10 - 90 %)
Sample Flow: 230 mL/min

Integrated Pneumatic Backup Delivery

Integrated pneumatic backup delivery = 250 mL/min Fixed Flow of NO/N₂

Physical

<table>
<thead>
<tr>
<th>Delivery system</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Max. Weight:</td>
<td>5.3 kg</td>
</tr>
<tr>
<td>Max. Width and Depth:</td>
<td>350 mm W x 160 mm D</td>
</tr>
<tr>
<td>Max. Height:</td>
<td>220 mm</td>
</tr>
</tbody>
</table>
Environmental

<table>
<thead>
<tr>
<th>Operating:</th>
<th>Transport/Storage:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperature:</td>
<td>5 to 40°C</td>
</tr>
<tr>
<td>Humidity:</td>
<td>15 to 95% RH</td>
</tr>
<tr>
<td>Ambient Pressure:</td>
<td>57 to 110 kPa</td>
</tr>
<tr>
<td>Water Ingress Protection:</td>
<td>IPX1</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Transport/Storage:</th>
<th>-20 to + 60°C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Humidity:</td>
<td>15 to 95% RH</td>
</tr>
<tr>
<td>Ambient Pressure:</td>
<td>57 to 110 kPa</td>
</tr>
<tr>
<td>Water Ingress Protection:</td>
<td>IPX1</td>
</tr>
</tbody>
</table>

INOMAX Regulator

- Inlet Pressure: 14 to 155 Bar (200 to 2,248 psig)
- Outlet Pressure: 1.7 to 2.4 Bar (25 to 35 psig)
- Cylinder Valve Connector: CGA 626 (ISO 5145)

Electrical

**Important:** Disconnect main power cord to isolate equipment from main power.

- Input Voltage: 100-240 V AC @ 50 / 60 Hz
- Input Power: 110 VA max
- Input Fuse: 3 A
- Classification: Class I, Type B
- Standards: CSA certified to meet the following for medical electrical equipment:
  - UL 60601-1: 2003 edition 2
  - ANSI/AAMI 60601-1: 2005 edition 3
  - IEC 60601-1: 2005 edition 3
- Battery Backup: A sealed lithium ion rechargeable battery provides power backup to operate the system for up to six hours when fully charged.
  - Connect the system to an electrical outlet for at least ten hours to charge the battery.
  - When the low battery alarm occurs, there are 30 minutes until battery depletion.
  - Dispose of used batteries according to local regulations.
- USB Port: Not used. Not for use when patient is connected.
- Ethernet Port: For service only. Not for use when patient is connected.
- RS 232: Enables serial communications for use with hospital electronic health record (EHR) system.
- Infrared Port: Infrared communication with the INOMAX cylinder.

Alarm Log

The alarm history is deleted when device is turned off. However, the service log, which is accessible by service personnel is maintained (including alarm log) when power is cycled and/ or when total power loss occurs.
RS 232 Data Output

Enables serial communications for use with hospital electronic health record (EHR) system. Must be connected to the manufacturer-specified third-party hardware (to be determined).

**WARNING:**
- INOmax DSIR Plus should only be connected to RS 232 ports that have:
  - Four kV input to output isolation
  - Four kV input to mains isolation and
  - an internal “reference voltage” “U” (as defined in section 20.3 of IEC60601-1 edition two) of less than or equal to 50 VDC or 50 VRMS and dielectric isolation certified in accordance with IEC 60601-1. Interface cabling must not go outside of the room (e.g., into walls where potential isolation issues could exist). Adherence to the above provide compliance to clause 20.3 “Value of test Voltage” in edition two and clause(s) 8.5.4 “Working Voltage” and Clause 8.8.3 “Dielectric Strength” in edition three.
- RS 232 cables must be shielded. The RS 232 cable shield shall have a minimum of 90% coverage. The shield shall only be connected at one end of the cable to minimize noise induced by ground currents.

**Note:**
- Connector retention jack posts can be found at the INOmax DSIR Plus connector. The RS 232 interface cable/connector should be constructed to include cable retention fasteners to help ensure a robust connection.
- This serial communication protocol requires INOmax DSIR Plus software revision 2.1 or higher to function. The software revision of the device can be accessed by pressing the Menu Button on the Main Screen and then the Settings Button (see Figure 1-6.).

**Definitions**

<table>
<thead>
<tr>
<th>Acronym/Definition</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>CRC</td>
<td>Cyclic Redundancy Check</td>
</tr>
<tr>
<td>RS 232</td>
<td>RS 232 (Recommended Standard 232) is the traditional name for a series of standards for serial binary single ended data and control signals connecting between a DTE (Data Terminal Equipment) and a DCE (Data Circuit-terminating Equipment).</td>
</tr>
<tr>
<td>ASCII</td>
<td>American Standard Code for Information Interchange</td>
</tr>
</tbody>
</table>

**RS 232 Port:**
- Nine pin female DSUB connector
- Pin two - received data, Pin three - transmitted data, Pin five - ground (isolated), Pin seven - RTS (unused), Pin eight - CTS (unused) and Pins one, four, six and nine - no connection
- 38,400 baud, one start bit, eight ASCII data bits, one stop bit, no parity, and no flow control
- Messages are output at a minimum rate of once per second, terminated with a checksum and carriage return
**Data output includes:**

- **Device information**
  - Model number, device generated identifier, software revision and user generated patient identifier
- **Monitored values**
  - Monitored O₂, NO₂ and NO
- **Settings**
  - Dose setpoint
  - Alarm setpoints
    - High O₂, low O₂, high NO₂, high NO and low NO
- **Alarm messages**
- **Device status**
- **INOMAX cylinder serial number and open/closed status**

**Note:** A detailed document regarding output data format is available upon request.
# Electromagnetic Compatibility Information

<table>
<thead>
<tr>
<th>Emissions Test</th>
<th>Compliance</th>
<th>Electromagnetic Environment - Guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF radiated emissions per</td>
<td>Group 1</td>
<td>The INOmax DS\textsubscript{IR} Plus system uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment. The INOmax DS\textsubscript{IR} Plus system 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>CISPR 11 Ed. 5.1b:2010</td>
<td>Class B</td>
<td></td>
</tr>
<tr>
<td>RF conducted emissions per</td>
<td>Class B</td>
<td></td>
</tr>
<tr>
<td>CISPR 11 Ed. 5.1b:2010</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Harmonic emissions</td>
<td>Class B</td>
<td></td>
</tr>
<tr>
<td>IEC 61000-3-2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Voltage fluctuations/flicker</td>
<td>Complies</td>
<td></td>
</tr>
<tr>
<td>emissions</td>
<td></td>
<td></td>
</tr>
<tr>
<td>IEC 61000-3-3</td>
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</tr>
</tbody>
</table>
**Guidance and Manufacturer's Declaration – Electromagnetic Immunity**

The INOmax DSIR Plus system is intended for use in the electromagnetic environment specified below. The user of the INOmax DSIR Plus system 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>± 0.5kV, ± 1.0kV and ± 2.0kV for power supply lines</td>
<td>± 0.5kV, ± 1.0kV and ± 2.0kV for power supply lines</td>
<td>Mains power quality should be that of a typical commercial or hospital environment.</td>
</tr>
<tr>
<td>IEC 61000-4-4</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Surge</td>
<td>Line-to-line: ± 0.5 kV, ± 1 kV</td>
<td>Line-to-line: ± 0.5 kV, ± 1 kV</td>
<td>Mains power quality should be that of a typical commercial and/or hospital environment.</td>
</tr>
<tr>
<td>IEC 61000-4-5</td>
<td>Line-to-earth: ± 0.5 kV, ± 1 kV, ± 2 kV</td>
<td>Line-to-earth: ± 0.5 kV, ± 1 kV, ± 2 kV</td>
<td></td>
</tr>
<tr>
<td>Voltage dips, short interruptions and voltage variations on power supply input lines</td>
<td>&lt;5 % $U_T$ ( &gt;95 % dip in $U_T$) for 0.5 cycle</td>
<td>&lt;5 % $U_T$ ( &gt;95 % dip in $U_T$) for 0.5 cycle</td>
<td>Mains power quality should be that of a typical commercial and/or hospital environment. If the user of the system requires continued operation during power mains interruptions, it is recommended that the system be powered from an uninterruptible power supply or a battery.</td>
</tr>
<tr>
<td>IEC 61000-4-11</td>
<td>40 % $U_T$ (60 % dip in $U_T$) for 5 cycles</td>
<td>40% $U_T$ (60% dip in $U_T$) for 5 cycles</td>
<td></td>
</tr>
<tr>
<td></td>
<td>70 % $U_T$ (30 % dip in $U_T$) for 25 cycles</td>
<td>70% $U_T$ (30% dip in $U_T$) for 25 cycles</td>
<td></td>
</tr>
<tr>
<td></td>
<td>&lt;5 % $U_T$ ( &gt;95 % dip in $U_T$) for 5 s</td>
<td>&lt;5 % $U_T$ ( &gt;95 % dip in $U_T$) for 5 sec.</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** $U_T$ is the AC mains voltage prior to application of the test level.
### Guidance and Manufacturer’s declaration - Electromagnetic Immunity

INOmax DS_{IR} Plus system is intended for use in the electromagnetic environment specified below. The user of the INOmax DS_{IR} Plus system should assure that they are 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>IEC 61000-4-6</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>3 Vrms</td>
<td>3 Vrms (V1)</td>
<td>Portable and mobile RF communications equipment, including cables, should be used no closer to any part of the INOmax DS_{IR} Plus system than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter. Recommended separation distance (d = 1.2\sqrt{P})</td>
</tr>
<tr>
<td></td>
<td>10 Vrms</td>
<td>10 Vrms (V2)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>150 kHz to 80 MHz</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>outside ISM bands(^a)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Radiated RF</td>
<td>IEC 61000-4-3</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>10 V/m</td>
<td>10 V/m</td>
<td>Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey(^c), should be less than the compliance level in each frequency range.(^d)</td>
</tr>
<tr>
<td></td>
<td>26 MHz to 1 GHz</td>
<td>26 MHz to 1 GHz</td>
<td>Interference may occur in the vicinity of equipment marked with the following symbol:</td>
</tr>
<tr>
<td></td>
<td>3 V/m</td>
<td>3 V/m</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1 GHz to 2.5 GHz</td>
<td>1 GHz to 2.5 GHz</td>
<td></td>
</tr>
</tbody>
</table>

\(^a\) The ISM (industrial, scientific and medical) bands between 150 kHz and 80 MHz are 6.765 MHz to 6.795 MHz; 13.553 MHz to 13.567 MHz; 26.957 MHz to 27.283 MHz; and 40.66 MHz to 40.70 MHz.

\(^b\) The compliance levels in the ISM frequency bands between 150 kHz and 80 MHz and in the frequency range 80 MHz to 2.5 GHz are intended to decrease the likelihood that a portable communications device could cause interference if it is inadvertently brought into patient areas. For this reason, an additional factor of \(10/3\) is used in calculating the recommended separation distance for transmitters in these frequency ranges.

\(^c\) 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 INOmax DS_{IR} Plus system is used exceeds the applicable RF compliance level above, the INOmax DS_{IR} Plus system should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as reorienting or relocating the INOmax DS_{IR} Plus system.

\(^d\) Over the frequency range 150 kHz to 80 MHz, field strengths should be less than 3 V/m.
The INOmax DSIR Plus system is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The user of the INOmax DSIR Plus system can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the INOmax DSIR Plus system 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 Outside ISM bands\ 150 kHz to 80 MHz In ISM bands</td>
</tr>
<tr>
<td>0.01</td>
<td>0.12</td>
</tr>
<tr>
<td>0.1</td>
<td>0.38</td>
</tr>
<tr>
<td>1</td>
<td>1.2</td>
</tr>
<tr>
<td>10</td>
<td>3.8</td>
</tr>
<tr>
<td>100</td>
<td>12</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.

NOTE 3: An additional factor of 10/3 is used in calculating the recommended separation distance for transmitters in the ISM frequency bands between 150 kHz and 80 MHz and in the frequency range 80 MHz to 2.5 GHz to decrease the likelihood that mobile/portable communications equipment could cause interference if it is inadvertently brought into patient areas.

NOTE 4: These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects, and people.
INOmax DSIR® Plus

9/ Appendix
INOmax DS_{IR} Plus

9/ Appendix
9/ Appendix

Manual Pre-Use Checkout

The following instructions are provided for when the on-screen pre-use wizard is not used.

1. Turn device ON, low calibration will begin and complete (Continue with steps 2-4 while calibration completes)

2. Initial Connections:
   Confirm attachment of the following:
   a. Water separator cartridge, water bottle, and patient gas sample line in place
   b. Injector module cable and tubing are connected
   c. Plug in power cord and verify AC power light is ON
   d. Regulator to INOMAX cylinder
   e. Regulator hose to INOmax DSIR inlet
   f. INOblender hose connected and white lock in place
   g. Oxygen source (50 psig) to back of INOblender
   h. IR cable in place

3. Assemble pre-use set-up connectors (see Figure 9-1). Do Not turn on O₂ flowmeter yet.

4. High Pressure Leak Test:
   Open/close INOMAX cylinder valve
   a. Verify, at least 34.5 bar (500 psig) cylinder pressure
   b. Verify, no decrease in cylinder pressure for 30 seconds
5. **Manual Purge/Alarm Verification:**
   a. Press CANCEL to exit pre-use wizard (low calibration should be complete to continue).
   b. Verify INOMAX cylinder valve is closed.
   c. Set O₂ flowmeter to 10 L/min
   d. Purge INOmax DSIR
      • Set the INOMAX dose based on cylinder concentration:

<table>
<thead>
<tr>
<th>Cylinder Concentration (ppm)</th>
<th>800</th>
<th>400</th>
</tr>
</thead>
<tbody>
<tr>
<td>Set Dose (ppm)</td>
<td>40</td>
<td>20</td>
</tr>
</tbody>
</table>

   • “Cylinder Valve Closed” alarm will occur.
   • Continue until cylinder gauge pressure drops to 0 psig.
   • Measured NO₂ will increase and then decrease as NO₂ is purged from the system.
   • “Low Cylinder Pressure” alarm will occur.
   e. Turn INOMAX dose to zero.
   f. Open INOMAX cylinder valve.

6. **Integrated Pneumatic Backup Test:**
   a. Verify pre-use assembly flowmeter set to 10 L/min
   b. Turn INOmax DSIR backup switch ON
   c. Allow monitored values to stabilize
   d. Verify measured values based on cylinder concentration

<table>
<thead>
<tr>
<th>Cylinder Concentration (ppm)</th>
<th>800</th>
<th>400</th>
</tr>
</thead>
<tbody>
<tr>
<td>NO (ppm)</td>
<td>14 - 26</td>
<td>7 - 13</td>
</tr>
<tr>
<td>NO₂ (ppm)</td>
<td>≤ 1.0</td>
<td>≤ 1.0</td>
</tr>
</tbody>
</table>

e. Turn backup switch OFF
7. **Performance Test:**
   
a. Verify $O_2$ flowmeter is set to 10 L/min
   
b. Set INOMAX dose based on cylinder concentration:

<table>
<thead>
<tr>
<th>Cylinder Concentration (ppm)</th>
<th>800</th>
<th>400</th>
</tr>
</thead>
<tbody>
<tr>
<td>Set Dose (ppm)</td>
<td>40</td>
<td>20</td>
</tr>
</tbody>
</table>

   c. Verify monitored values

<table>
<thead>
<tr>
<th>Cylinder Concentration (ppm)</th>
<th>800 ppm</th>
<th>400 ppm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Set Dose (ppm)</td>
<td>40</td>
<td>20</td>
</tr>
<tr>
<td>Acceptable NO Value (ppm)</td>
<td>35 - 45</td>
<td>17 - 23</td>
</tr>
<tr>
<td>Acceptable NO$_2$ Value (ppm)</td>
<td>&lt; 1.5</td>
<td>&lt; 1</td>
</tr>
<tr>
<td>Acceptable FiO$_2$ (%)</td>
<td>95 ± 3</td>
<td>95 ± 3</td>
</tr>
</tbody>
</table>

d. Set INOMAX dose to 0 ppm
   
   • "Set Dose is Zero, Please Close Cylinder Valve" reminder will appear- DO NOT close cylinder valve at this time, dismiss reminder.

e. Turn oxygen flowmeter OFF

8. **INOblender Test:**
   
a. Remove injector module from pre-use assembly and reconnect tubing

b. Remove $O_2$ tubing from flowmeter and attach to INOblender outlet

c. Set INOblender flow to 10 L/min, INOMAX dose to:

<table>
<thead>
<tr>
<th>Cylinder Concentration (ppm)</th>
<th>800</th>
<th>400</th>
</tr>
</thead>
<tbody>
<tr>
<td>Set Dose (ppm)</td>
<td>40</td>
<td>20</td>
</tr>
</tbody>
</table>

d. Verify monitored values on the INOmax DS$_{IR}$ Plus

<table>
<thead>
<tr>
<th>Cylinder Concentration (ppm)</th>
<th>800</th>
<th>400</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acceptable NO Value (ppm)</td>
<td>32 - 48</td>
<td>16 - 24</td>
</tr>
</tbody>
</table>

e. Set INOblender dose and flow to 0
Pre-Use Assembly

1. O₂ Flowmeter
2. O₂ Tubing
3. 15M x 4.5 mm Adapter
4. 22M / 15F x 22M / 15F Adapter
5. Injector Module
6. 300 mm of 22 mm hose
7. Patient Gas Sample Line with Nafion
8. Gas Sample Tee
9. Injector Module Electrical Cable
10. NO/N₂ Injector Tube

Figure 9-1

Additional Dose Setting Information

Each click on the control knob corresponds to a known change in dose. The incremental dose per click corresponds to a value dependent upon the dose range in which the change is made, as illustrated in the table below.

<table>
<thead>
<tr>
<th>Dose Setting Range</th>
<th>Dose Change Per Click 400 ppm</th>
<th>Dose Change Per Click 800 ppm</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 to 40 ppm</td>
<td>1 ppm</td>
<td>1 ppm</td>
</tr>
<tr>
<td>40 to 80 ppm</td>
<td>NA</td>
<td>2 ppm</td>
</tr>
</tbody>
</table>