



HVT 2.0 High Velocity Therapy System

Instructions for Use





The screen images shown in this IFU may not match the layout of the screen on your device due to configurable parameters the user may have selected during set-up or the software version of the device

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HVT 2.0

Section 1: Indications, Warnings & Precautions

The HVT 2.0 high velocity therapy system consists of the HVT 2.0 device and a Disposable Patient Circuit (DPC).

The HVT 2.0 system is equipped with an internal air blower. The addition of an external oxygen source (wall, tank, or oxygen concentrator) enables FiO_2 delivery from 21% to 100%, dependent on the oxygen source.

The HVT 2.0 system is intended to be used by qualified medical professionals, such as physicians, nurses, respiratory therapists.

Accessories may not be available in all countries. Contact the local Vapotherm representative for more information.

If there is a serious incidence with the device, it should be reported to Vapotherm Technical Support at the number specified on the final page of the IFU or to your Authorized Vapotherm Representative. Follow local regulations and report the incidence to the competent authority or regulating agency.

Indications / Intended Use

The HVT 2.0 system provides high velocity nasal insufflation (HVNI) with simultaneous warmed and humidified respiratory gas delivery to augment breathing of spontaneously breathing adult and pediatric patients (5 kg and up) suffering from respiratory distress and/or hypoxemia in the hospital setting, via small bore nasal cannula. HVT 2.0 is not intended to provide total ventilatory requirements of the patient and not for use during field transport. The flow rates may be from 5 to 45 liters per minute (BTPS). The device is intended to be used in hospital, skilled nursing facilities, and sub-acute facilities.

Contraindications

- Not appropriate for patients who are not spontaneously breathing, are unable to protect their airway, or have anatomic or injury-induced blockage of the nasal pathway to the nasopharyngeal space.
- Not for treating OSA and snoring.
- HVT 2.0 is not for field transport.
- HVT 2.0 is MRI unsafe. Do not use it in an MR environment.
- Not for use with an Oxygen Concentrator (when treating respiratory distress).

Warnings & Precautions

Please take the time to review and become familiar with the warnings, precautions, and notes listed in this Instructions for Use document. They cover safety considerations, special requirements, and regulations. Warnings and precautions must be understood by the user to prevent adverse events to the patient or to the operator due to electromagnetic disturbances.

The user of this product shall have sole responsibility for any malfunction due to operation or maintenance errors by unauthorized/untrained personnel. Federal Law (U.S.) restricts the sale of this device to or by order of a physician. This device should be used only by a trained operator.

A Warning indicates that a situation may occur which is potentially harmful to the patient or user.	A Precaution indicates a condition that may lead to equipment damage, malfunction, or inaccurate operation.	A Note indicates a point of emphasis to make operation more efficient or convenient.

A General Warnings

- HVT 2.0 is not a Continuous Positive Airway Pressure (CPAP) device. There are no controls to deliver or monitor airway pressure. HVT 2.0 should not be used to deliver pressure in a closed system.
- Patients receiving supplemental oxygen are often acutely ill and appropriate clinical vigilance should be observed by the care team. Additional patient monitoring, including pulse oximetry, is necessary if the HVT 2.0 is used to give supplementary oxygen.
- Use only the accessories, transducers, and cables specified or provided by the manufacturer of this equipment. Use of other accessories, transducers, and cables other than those specified or provided by the manufacturer of this equipment could result in increased electromagnetic emissions or decreased electromagnetic immunity of this equipment and result in improper operation.
- Do not add any attachments or accessories to the HVT 2.0 system that are not listed in this Instructions for Use. The HVT 2.0 device might not function correctly, which could affect the quality of the therapy or could cause harm to the patient.

- Oxygen supports combustion. This device should not be used near open flames, flammables, oil, or grease. The device is not intended for use in oxygenrich environments.
- Do not use the device in or around water (with the exception of the water bag that feeds the system).
- The oxygen must be clean, dry, medicalgrade gas to prevent harm to the patient and to prevent damage to the device.
- Improperly sizing the cannula, specifically complete occlusion of the nares by the nasal prongs, may lead to a risk of pneumothorax.
- Cannula prongs should not obstruct more than 50% of the nares of the patient.
- Change nasal cannulas when soiled. Replace cannulas according to clinical judgment and hospital policy but not to exceed 30 days continuous use.
- If used with an oxygen concentrator, the maximum oxygen percentage will be limited depending on the type of concentrator used and total set flow.
- An Air Compressor is not to be used with the device.

A General Warnings (continued)

- Only use sterile water. Failure to utilize sterile water supply or clean oxygen source may increase the risk of bacterial contamination.
- Always follow aseptic technique (including proper hand washing and avoiding direct hand contact with connection points) when setting up the HVT 2.0 device and use Standard Precautions when placing on a patient.
- Use with magnetic resonance imaging (MRI) and radio frequency (RF) equipment may cause patient injury.
- Medical electrical equipment needs special precautions regarding radio frequency (RF) electromagnetic radiation. Portable and mobile RF communications equipment such as base stations for cordless telephones and land mobile radios, amateur radio and AM and FM radios, can affect medical equipment and should not be used near the device.
- If the HVT 2.0 system is placed in close proximity to RF emitters, an indication that HVT 2.0 performance is being affected by the emitters is false alarms and front panel display showing values that are out of specification. In certain circumstances, HVT 2.0 may affect or be affected by nearby equipment due to electromagnetic interference. If this should happen, try moving the HVT 2.0 device or moving the device causing interference or consult with the manufacturer.
- Portable RF communications equipment (including peripherals such as antenna cables and external antennas) should be used no closer than 30 cm (12 inches) to any part of the HVT 2.0 system, including specified cables. Otherwise, degradation of the performance of this equipment could result.
- Use of this equipment adjacent to or stacked with other equipment should be avoided because it could result in improper operation. If such use is necessary, this equipment and the other equipment should be observed to verify that they are operating correctly.

- To avoid electric shock;
 - Use only the power supply cord that was provided with the device. Do not use any other cord. Do not use extension cords.
 - Do not operate the device if the power supply cord is damaged.
 - The power supply cord can be disconnected to isolate the product from mains. It is recommended that the device be unplugged when not in use to prevent hazards occurring when unattended.
 - Do not modify this equipment without authorization from the manufacturer.
 - Before cleaning and disinfecting, unplug the device from line power.
 - Do not use near or in water.
 - Do not use if device is damaged.
- HVT 2.0 is MR Unsafe and is not intended for use in MRI environments. Keep away from magnetic resonance imaging (MRI) equipment.
- Do not connect any device, system, or accessory that has not been approved by Vapotherm.
- Do not use the HVT 2.0 device at an altitude above 3000m or outside a temperature of 18 to 30°C. Using the device outside of this temperature range or above this altitude can affect the quality of the therapy or harm the patient.
- To prevent disconnection of the tubing or tubing system during use, especially during ambulatory use, only tubes in compliance with ISO 5367 or ISO 80601-2-74 should be used.
- The device should not be turned on and left unattended when not connected to a patient.
- General Fault alarms are failures in the control or measurement systems. Depending on the cause of the failure, gas delivery may or may not be interrupted. If a General Fault alarm occurs, disconnect the patient, and shut off the device. The device must be repaired by trained service personnel.
- To reduce the risk of strangulation from patient tubing, use the provided tubing clip to secure the patient tubing.

General Warnings (continued)

- The internal safety battery is designed for temporary use only when AC or external DC power to the unit has been interrupted, and no transfer battery is present. When the HVT 2.0 device is running on the internal safety battery, there is no heat or humidity provided with the set flow and FiO, and the humidity level may drop below safe limits. After the internal safety battery is fully discharged, the device will not operate, and patient gas flow will cease. When fully charged, the internal safety battery provides at least 15 minutes of power. The internal safety battery is not intended for patient transport.
- All disposable components are labeled as "single patient use only" and must be replaced after 30 days of use on a single patient. Cannulas should be replaced according to clinical use, but not to exceed 30 days. Do not attempt to sterilize or reuse any of these components, and

follow all local and federal regulations for disposal. Outside the U.S., follow national or international regulations.

- Do not use disposables on more than one patient. Multi-patient use may lead to patient injury from infection and/or delivery of therapy outside of published specifications.
- To reduce the risk that the patient may aspirate condensed water from the breathing circuit, regularly observe the patient and output of the patient interface for excess water, and if detected remove patient interface from the patient. Water in the center lumen can result from condensation or due to a leak from the outer lumens that surround the breathing circuit. Condensation in the cannula may also occur in certain ambient conditions and low flow rates. If minimal condensation occurs after confirming no leaks, it is recommended to select a lower temperature setpoint.

A General Precautions

- HVT 2.0 will not operate without the internal safety battery in place. Have an internal safety battery on hand to ensure the continued availability of the use of the device. To ensure safe and reliable operation, use only the Vapotherm specified replacement battery.
- Do not
 - cover the device (blocking the vent may damage the device)
 - immerse the device in water
 - steam or gas sterilize the device.
- Even a fully charged battery will lose its charge over a period of weeks when the device is not connected to line power. It is recommended that the device be connected to line power at least 2 hours a month to maintain the battery charge. The internal safety battery should only be accessed or replaced by trained service personnel.
- Do not use bleach, organic solvents, or abrasive cleaners. Hypochlorite solutions liberate toxic gases such as chlorine when acidified or heated. The reaction with ammonia or with substances that can generate ammonia can produce chloramines which are also toxic and have explosive potential.

NOTE: Flexible sterile water supply is recommended. If rigid or semi-rigid bottles are used, a Vapotherm-approved venting bottle cap spike must be used.

NOTE: HVT 2.0 may be operated with a limited performance at oxygen inlet pressures as low as 4 psi (28 kPa). However, for the full specified range of gas flows and oxygen percentages, appropriate for treating respiratory distress, oxygen inlet pressures must be 40 psi (276 kPa) or above (Caution: Not to exceed 87 psi).

HVT 2.0

Section 2: Overview of the HVT 2.0 System

The HVT 2.0 high velocity therapy system consists of the HVT 2.0 device and a Disposable Patient Circuit (DPC).

While the HVT 2.0 device can be used over again on another patient, each patient will require a Disposable Patient Circuit (DPC) and nasal cannula, which are attached to the HVT 2.0 device and enable the delivery of high velocity therapy to the patient. The ProSoft nasal cannula is proprietary to Vapotherm and the HVT 2.0. device will not work correctly unless using ProSoft cannulas.

Other accessories validated for use with the device are the roll stand, the Transfer Upgrade Kit, and the HVT 2.0 Nurse Call Interface Cable and the HVT 2.0 EMR Link Cable.



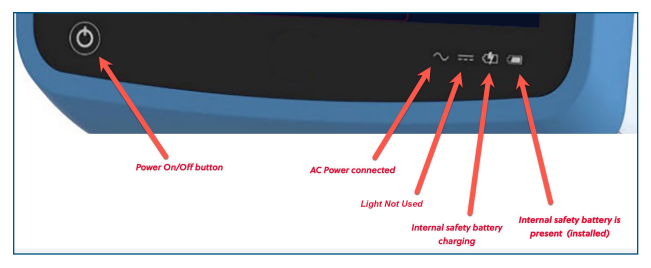


Figure 1: HVT 2.0 Device (above) and LED indicators on front of device

Features of the HVT 2.0 System

- Flow range from 5 to 45 L/min BTPS.
- Oxygen percentage is fully adjustable when connected to a 40 psi (276 kPa) oxygen gas source (non-oxygen concentrator).
- Temperature can be adjusted from 33 to 39°C.
- Built-in oxygen/air blender.
- All internal sensors are self-calibrating and self-monitoring.
- Flow, oxygen, and temperature settings are adjusted via touch screen scroll bars.
- Built-in electronic flow meters and controllers.
- Minimal downtime between patients: less than ten minutes to change disposables and disinfect.
- Warm-up time less than five minutes.
- Preheat feature circulates water and warms the circuit water to 33°C.
- EMR and Nurse Call connectivity capable of indicating an alarm condition on a hospital Nurse Call system and interfacing with Electronic Medical Record technologies.
- The Disposable Patient Circuit (DPC) is detachable and disposable: no disinfection necessary.
- A single DPC enables the full system flow range (5 to 45 L/min). The DPC is fully assembled and ready to use out of the packaging.
- Universal power requirements allow use anywhere with only a change of power cord.
- Internal safety battery maintains flow and oxygen percentage for at least 15 minutes if AC power is cut off. Safety battery recharges in 2 hours.



Principles of Operations

The HVT 2.0 system utilizes an integrated internal blower to deliver warmed and humidified breathing gas at flows up to 45 L/min to spontaneously breathing patients, without the need for wall air or any pressured air source. The device incorporates a proportional valve and flow sensors that allow the oxygen percentage and total gas flow to be set independently.

The HVT 2.0 system consists of two parts: the HVT 2.0 device and the Disposable Patient Circuit (DPC). A validated patient interface (e.g., Vapotherm ProSoft nasal cannula) is required to deliver therapy to the patient. In addition, there are optional accessories that can be added to the therapy based on a patient's needs.

HVT 2.0 Device

The HVT 2.0 device contains all the electrical and electronic components, including the proportional valve and flow controllers and remote sensors to monitor the disposable water path. The device has no water pathways, and the gas pathway contains only dry gas at room temperature, and consequently does not need internal cleaning or disinfection.

The flow of oxygen and air is measured by mass flow sensors. The operating software calculates the required flow of each needed to reach the target flow and oxygen percentage set by the operator. The system controls gas flows accordingly by adjusting proportional solenoid valves on the gas line.

Firmware running in the device uses sensors to monitor gas pressure and water temperature. Alarms are activated if any parameters are outside the normal range. Troubleshooting instructions for the alarms can be immediately displayed on the screen. Other indicators show a low charge in the internal safety battery.

After a two-hour charging period, the internal safety battery will maintain the set flow and oxygen blend for at least 15 minutes should AC power be interrupted.

WARNING: The internal safety battery is designed for temporary use only when AC power to the unit has been interrupted, and no transfer battery is present. When the HVT 2.0 device is running on the internal safety battery, there is no heat or humidity provided with the set flow and FiO₂ and the humidity level may drop below safe limits. After the internal safety battery is fully discharged, the device will not operate, and patient gas flow will cease. When fully charged, the internal safety battery provides at least 15 minutes of power. The internal safety battery is not intended for patient transport.

Disposable Components

The following disposable components are compatible with HVT 2.0:

- **Disposable Patient Circuit** (DPC) [**REQUIRED**] Single disposable patient circuit that enables delivery of high velocity therapy for adult, pediatric, and infant patients, for flows 5 to 45 L/min. The DPC is provided fully assembled and ready to use.
- **ProSoft® Nasal Cannula** [**REQUIRED**] Various sizes are available so that a cannula can be selected to fit the patient comfortably. The cannula is connected to the DPC.
- **Tubing Adapter** (optional) Available should the therapy need to be delivered through a trach mask or t-piece instead of the nasal cannula.

All disposable components are labeled as "single patient use only" and must be replaced after 30 days of use on a single patient. Cannulas should be replaced according to clinical use, but not to exceed 30 days. Do not attempt to sterilize or reuse any of these components, and follow all local and federal regulations for disposal. Outside the U.S., follow national or international regulations.



WARNING: All disposable components are labeled as "single patient use only" and must be replaced after 30 days of use on a single patient. Cannulas should be replaced according to clinical use, but not to exceed 30 days. Do not attempt to sterilize or reuse any of these components, and follow all local and federal regulations for disposal. Outside the U.S., follow national or international regulations.



WARNING: Do not use disposables on more than one patient. Multi-patient use may lead to patient injury from infection and/or delivery of therapy outside of published specifications.



Disposables	Patient Weight (kg) / Age			Flow Range (L/min)		
	5 - 10 kg ≥1 mo	10 - 20 kg ≥ 1 mo - 6 yrs	20 - 40 kg ≥ 6 - 18 yrs	> 40 kg > 12 yrs	>100 kg > 12 yrs	
Disposable Patient Ci	rcuit (DPC)	·	,	·		
Air/O ₂ (Standard)	Х	Х	Х	Х	Х	5-45
Cannulas	1		1	1		
ProSoft Adult Long					Х	5-45
ProSoft Adult				Х		5-45
ProSoft Adult Small/ Pediatric			Х			5-45
Unicorn Adult			х	Х	Х	5-45
ProSoft Pediatric Small		Х				5-20
Unicorn Pediatric		Х				5-20
ProSoft Intermediate Infant	×					5-8
ProSoft Infant	Х					5-8
Unicorn Infant	Х					5-8
Optional add-on	Optional add-on					
Tubing Adapter	Х	х	X	Х	Х	8-45



WARNING: Cannula prongs should not obstruct more than 50% of the nares of the patient.

NOTE: Cannula prongs should not obstruct more than 50% of the total combined area of the nares of the patient.



NOTE: When delivering therapy to pediatric patients (via a nasal cannula or the tubing adapter), the recommended guidance for setting starting flow rate is 2 L/min/kg. See <u>"Performance" on page C-2</u> for humidification output at specific flow rates.

NOTE: The Tubing Adapter is not intended to be connected directly to a trach collar. An open system must be maintained to ensure gas egress. To facilitate humidification via a bypassed upper airway, connect to a tracheostomy mask or T-piece.

Accessories

- Roll Stand rolling stand to hold the HVT 2.0 device.
- **Transfer Upgrade Kit** to allow for moving the patient from one location to another within the hospital, includes:
 - Transfer Battery 1-hour Lithium-ion battery (VTBP-2.0, 14.4Vdc; 2 x6800mAh; 2 x97.9 Wh)
 - Oxygen manifold
 - Oxygen hoses (U.S. Only)
 - Adjustable oxygen tank holder (Use only E-cylinder size)
- HVT 2.0 Nurse Call Interface Cable (2.9m) to allow connectivity to the hospital Nurse Call System.
- HVT 2.0 EMR Link Cable (2.9m) to allow interface with the hospital Electronic Medical Record System.



WARNING: Use only the accessories, transducers, and cables specified or provided by the manufacturer of this equipment. Use of other accessories, transducers, and cables other than those specified or provided by the manufacturer of this equipment could result in increased electromagnetic emissions or decreased electromagnetic immunity of this equipment and result in improper operation.



WARNING: Do not add any attachments or accessories to the HVT 2.0 system that are not listed in this Instructions for Use. The HVT 2.0 device might not function correctly, which could affect the quality of the therapy or could cause harm to the patient.



HVT 2.0

Section 3: Setting Up the HVT 2.0 System

The following steps must be taken in preparation for using the HVT 2.0 System:

- 1. Assemble the HVT 2.0 device for use (including the Transfer Upgrade Kit, if applicable). (See details below).
- 2. Complete the Initial Setup of device settings. (See <u>"HVT 2.0 Initial Device Setup</u> <u>Process"</u> details below).
- 3. Insert the Disposable Patient Circuit (DPC) unit. **Note**: The DPC is provided fully assembled and ready to use.
- 4. Select the patient and interface type.
- 5. Connect the HVT 2.0 device to the patient.

Assemble the HVT 2.0 Device for Use

- Attach the HVT 2.0 device securely to the sturdy roll stand or place it on a tabletop. See <u>"Appendix C – Technical Specifications"</u> for the roll stand dimensions.
- 2. Visually check that the patient air filter is installed. (The HVT 2.0 device comes with one pre-installed, and there is a replacement in the user kit).
- 3. [**Optional**] Install the Transfer Upgrade Kit. For step-by-step instructions, see the <u>"Set up the Transfer Upgrade Kit"</u> section below.
- 4. Insert power supply cord into a facility-approved wall outlet.

WARNING: Do not operate the device if the power supply cord is damaged.
WARNING: The power supply cord can be disconnected to isolate the product from mains. It is recommended that the device be unplugged when not in use to prevent hazards occurring when unattended.
WARNING: Use only the power supply cord that was provided with the device. Do not use any other cord. Do not use extension cords.

- 5. Connect the oxygen hose to the oxygen inlet at the back of the device or tubing if using an oxygen concentrator.
- 6. If applicable, connect the Nurse Call cables to the appropriate port on the back of the HVT 2.0 device.

Once connected, the Nurse Call System will be enabled.

See <u>"Appendix A – Nurse Call System Installations"</u> for complete instructions for use on connection and use of Nurse Call.

If available, connect the EMR system. See <u>"Appendix B – Electronic</u> <u>Medical Records (EMR) Integration</u>" for more information.

Set up the Transfer Upgrade Kit

Setting up the Transfer Upgrade Kit involves four (4) steps:

- 1. Insert the Transfer Battery into the HVT 2.0 device.
- 2. Attach the oxygen manifold to the Roll Stand.
- 3. Connect the oxygen hoses to their appropriate locations per the labels on each hose.
- 4. Attach the tank holder to the Roll Stand.

For more information on transferring patients, see <u>"Intra-Hospital Patient Transfer" on page 42.</u>

NOTE: The maximum weight capacity of the roll stand when equipped with the transfer kit is 34.5 kg. The basket has a maximum weight capacity of 4.4 kg.

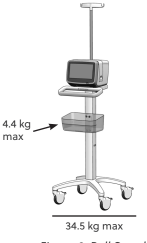


Figure 2: Roll Stand



HVT 2.0 Initial Device Setup Process

There is a five-step process for initial setup for the HVT 2.0 device. **Note**: Initial setup screens will only appear when powering on the dervice for the first time or after Factory Reset is selected (from the Admin Settings menu).

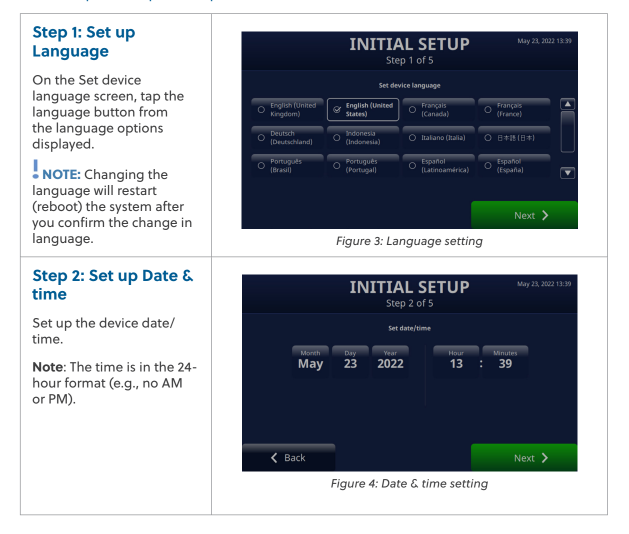
<u>"Step 1: Set up Language"</u>

"Step 2: Set up Date & time"

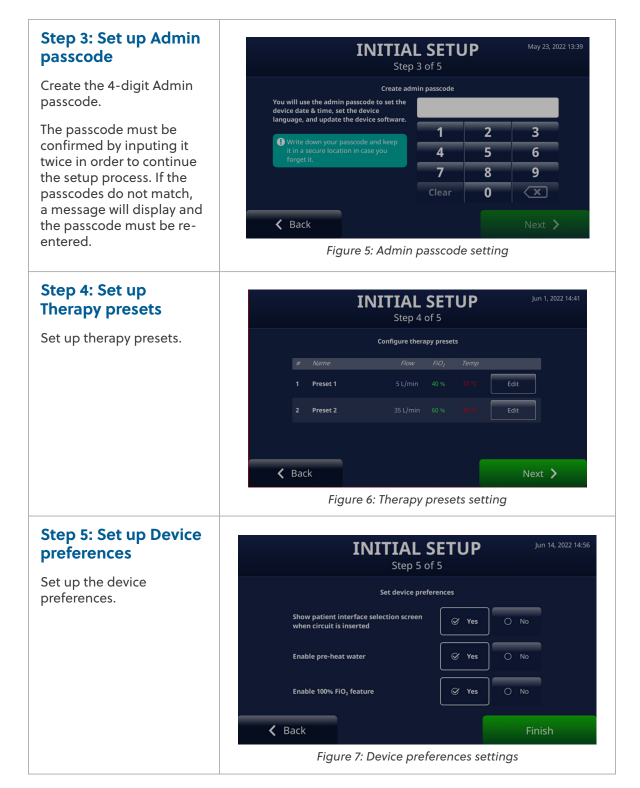
"Step 3: Set up Admin passcode"

"Step 4: Set up Therapy presets"

"Step 5: Set up Device preferences"







HVT 2.0 Device Settings

The HVT 2.0 device has a number of settings that can be accessed and adjusted, if necessary. These settings can be adjusted from the Settings menu.

To access the Settings menu:

1. Tap the [**Unlock**] button at the top left corner of the main screen.

The [**Unlock**] button will be replaced by a [**Lock**] button, and a [**Menu**] button will display next to it.

2. Next, tap the [**Menu**] button to display the Settings menu.

Lock Menu	NO CIRCUIT Tap value to adjust target	Jun 1, 2022 14:53
Flow	FiO ₂	Temp
8	21	37
L / min	%	°C
Alarm silence		🕻 Dim display

Figure 8: Tap the [**Unlock**] button to access Settings [**Menu**] button

The [**Menu**] button enables users to access the General Settings and Admin Settings in order to establish settings for the device in both Standby and Run modes.

The following settings can be defined from the General Settings menu:

- **Circuit details** Select the patient interface (trach adapter or cannula size). See<u>"Circuit Details" on page 23</u> for more details on this feature.
- Screen brightness Set the desired screen brightness level.
- Audio volume Set the volume level of the alarms.
- **Event log** Displays all events captured on the device, including event details, date and time. See<u>"Event Log" on page 23</u> for more details on this feature.
- Oxygen Source Select the source of the oxygen that will be connected to the device (e.g., Wall/tank or Concentrator). See<u>"Oxygen Source" on page 24</u> for more details on this feature.
- **Software details** Displays Controller board and GUI application software versions.
- Admin settings access Enables user to open the Admin Settings menu.

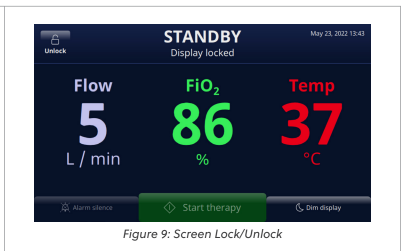
The following settings can be defined from the Admin Settings menu:

- Preferences (includes Patient Type selection and Preheat)
- **Preheat water** Enable the Preheat water mode, which circulates water and warms the circuit water to 33°C. See<u>"Preheat Water" on page 25</u> for more details on this feature.

- Therapy presets Add or edit therapy preset configurations for Flow, Oxygen, and Temperature. See<u>"Therapy Presets" on page 27</u> for more details on this feature.
- Date & time Set the format for the device date and time.
- Language Select the screen language.
- **Calibrate water level** Activate water level calibration, only if advised by Vapotherm to do so.
- Admin Passcode The user is able to change the admin passcode.
- Software update Update Controller board and GUI application software, when directed by Vapotherm personnel. See<u>"Appendix D: Software Update Process"</u> for more instructions on how to udate the device software
- Factory reset The user is able to restore the device to factory settings from the Admin. Settings menu when not in therapy. Therapy must be stopped to reset the device. Select "Factory Reset" from the Admin Settings Menu. The Factory reset will reboot the device and display the <u>"HVT 2.0 Initial Device Setup Process"</u>.

NOTE on Screen

The screen will lock automatically after 3 minutes of non-activity. To unlock the screen, tap the [**Unlock**] button in the top left of the screen.



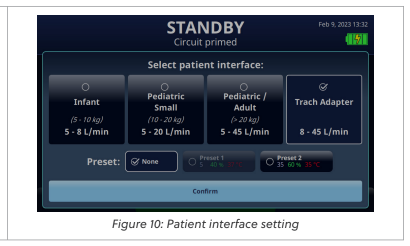


Circuit Details

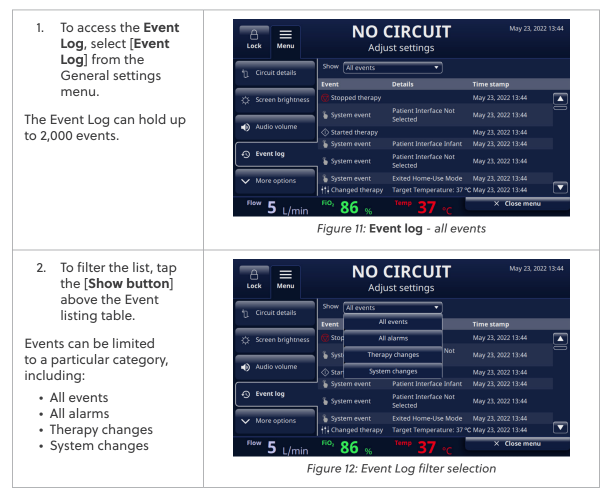
Select patient interface

When you select **Circuit Details** from the General settings menu, the Select patient interface screen displays.

From the screen, select the correct cannula size or select Trach adapter for the patient.



Event Log





Oxygen Source

Oxygen Source can be selected from the General settings menu.

 Select the source of oxygen for the patient by tapping either the Wall/tank option or the Concentrator option.

May 23, 2022 13:41 **NO CIRCUIT** 8 Lock Menu Adjust settings O₂ Oxygen source 🧭 🛛 Wall / tank O Concentrator Oxygen source (i) Software details {ဂ်ိ} Admin settings Back to top FIO2 21 × Close menu Flow 8 L/min

Figure 13: Therapy presets setting

• NOTE: When using an oxygen concentrator, refer to the Concentrator Flow Rate table in <u>"Appendix C – Technical Specifications"</u>- in the "Use with Oxygen Concentrator" section, for the estimated oxygen concentration.

- 2. If **Concentrator** is selected, a confirmation message <Option is not for respiratory distress will be displayed>. User acknowledgement of the message is required to continue.
- !Note: The device will always default to Wall/ Tank oxygen source upon startup.

If concentrator selection is confirmed, the current O percentage and maximum flow rate will display. To make a change, tap on O, concentration to select a new oxygen concentration percentage. To set the maximum flow rate, tap either 5 L/Min or 10 L/Min. A checkmark will display next to the selected flow rate.



Figure 14: Oxygen source: Concentrator selected

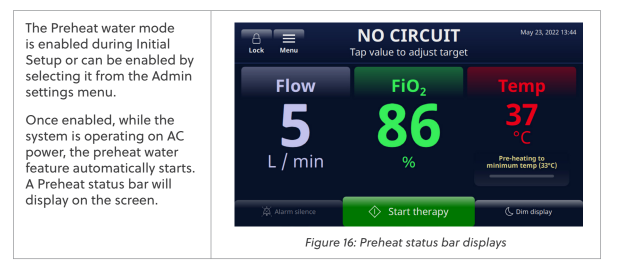


3. The system will May 23, 2022 13:43 **NO CIRCUIT** default to 92% oxygen Lock Menu Adjust settings concentration. If you need to change the O, Oxygen source O Wall / tank ♂ Concentrator oxygen concentration, Oxygen source: tap the 92 and select (i) Software details O₂ concentration Max flow rate the new percentage 96 𝗭 5 L/min from the scroll bar. 4. Tap the [Confirm] O₂ conc. 91 92 93 94 95 96 button below the scroll bar to save the changes. Confirm Figure 15: Concentrator - Scroll bar to change O, changes

Preheat Water

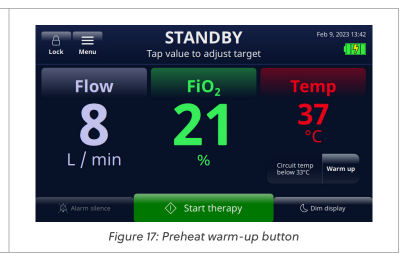
Enabling the Preheat water mode circulates water and warms the circuit water to 33°C. When the system is operating from AC power, preheat automatically starts after circuit priming completes or therapy is stopped.

Preheat Water is part of the Preferences option from the Admin settings menu.





Note that when the system operating power transitions from AC power to Transfer battery power, preheating the water can be activated via the [**Warm-up**] button that will display on the screen. Press the button to begin the pre-heat process.

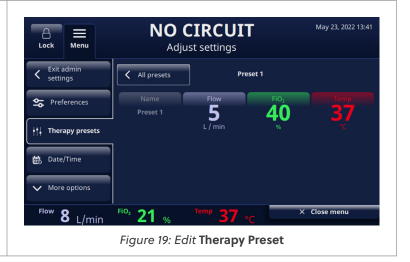


Therapy Presets

- From the Admin settings menu, tap the [Therapy Presets] button to display the screen.
- The Therapy Presets settings allow you to set up two different therapy settings that are ready for quick initiation of therapy. To create a preset, tap the [Edit] button to the right of the preset.
- To select your preset values, select the Flow, FiO₂, or Temp by tapping them.



Figure 18: Therapy presets





 When the parameter is tapped, a scroll bar will display below it. Use the scroll bar to select your target value(s) for each parameter: Flow, FiO₂, or Temp.

> Tap [**Confirm**] to save the value as a part of the preset.



Software Update

From the Admin Settings menu, tap the **[Software update**] button to display the screen.

The screen will display the latest software versions and dates for the Controller board and Graphic User Interface (GUI) application.

• NOTE: The button will only be enabled when a USB is placed in the back of the device. See <u>"Appendix D:</u> <u>Software Update Process"</u> for sofware update instructions.





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HVT 2.0

Section 4: Using the HVT 2.0 system

Modes of Operation overview

HVT 2.0 has three modes of operation: Sleep, Standby, and Run. The mode displays at the top of the screen.



Figure 22: Modes of operation

Sleep: The device is plugged in but is not turned on. No therapy is being delivered.

Standby: The device is on, and the screen is illuminated. No therapy is being delivered. This mode is indicated by either "**NO CIRCUIT**" (DPC is not installed) or "**STANDBY**" (DPC installed but therapy not started). To start therapy, tap the [**Start therapy**] button at the bottom of the screen.

Run: The device is on and delivering therapy according to the parameter's setup. This mode is indicated by "**RUNNING**" being displayed at the top of the screen and the [**Stop therapy**] button at the bottom of the screen.

Start-Up Preparation

Before assembling the device for use, make sure the following items are available:

- Disposable Patient Circuit (DPC)
- Sterile water supply
- Validated patient interface (ProSoft nasal cannula or trach adapter)
- Adequate oxygen source and connectors
- 1. Visually inspect the power supply cord to verify that it is not damaged or kinked. Then, plug the power supply cord into a facility-approved wall outlet.
- 2. Connect the oxygen hose to the facility-approved oxygen wall outlet, oxygen tank, or an oxygen concentrator.

NOTE: When using an oxygen concentrator, a 1/4" barb nipple is required.

3. Press the () [**Power**] button on the device. The screen will illuminate and display the software revision number and will automatically conduct the alarm sound test.



Starting Therapy

WARNING: The device should not be turned on and left unattended when not connected to a patient.

If the device has been exposed to very hot or very cold temperatures, allow it to reach operating temperature before use. After being exposed to extreme temperatures, the device can take up to 12 hours to reach room temperature.

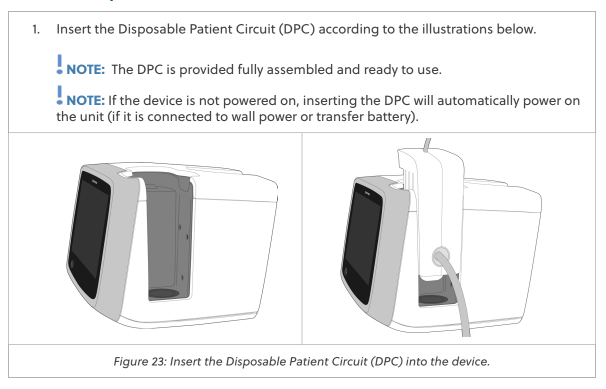
The Start Therapy procedures have been divided into the following sections:

- 1. <u>"Insert the Disposable Patient Circuit (DPC)"</u>
- 2. <u>"Select Patient Interface Type"</u>
- 3. <u>"Select Therapy Parameters"</u>
- 4. "Start Therapy"

NOTE: Optional Therapy Presets are available to select your therapy parameters.

Initiate and Start Therapy

Insert the Disposable Patient Circuit (DPC)





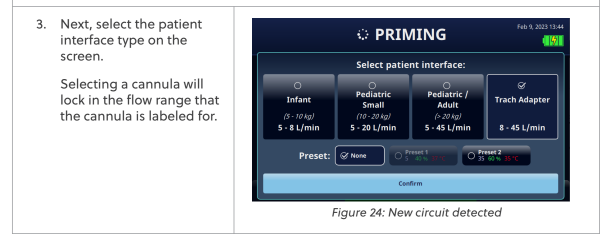
2. Hang the sterile water supply from the Roll Stand and connect it to the DPC via the water inlet tube. Spike the water supply and unclip the tubing, if necessary, to begin the flow of water to the device.

NOTE: When using some water bottles, it may be necessary to use a hook to lower and hang the water bottle (included in the user kit) to facilitate connection with the spike tube.

Flexible sterile water supply are recommended. If rigid or semi-rigid bottles are used, a Vapotherm-approved venting bottle cap must be used.



WARNING: Only use sterile. Failure to utilize sterile water supply or clean oxygen source may increase the risk of bacterial contamination.



Select Patient Interface Type

1. Select the desired patient interface type on the screen. If selecting a cannula, place the cannula on the patient to allow the cannula to warm to the patient's skin temperature. This helps minimize condensation during therapy delivery.

NOTE: The Infant cannula selection includes both the infant and intermediate-sized cannulas. The Pediatric Small is limited to the pediatric small-sized cannula. And, the Pediatric/Adult cannula selection includes the Pediatric/Adult Small, Adult, and Adult Long-sized cannulas. [For more information on cannula sizes, see <u>"HVT 2.0 Disposable Components" on page 15.</u>].



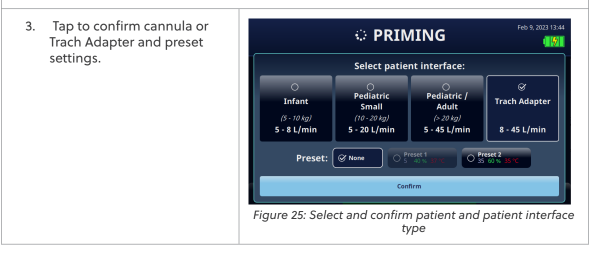
WARNING: Cannula prongs should not obstruct more than 50% of the nares of the patient.

NOTE: If the flow rate was set prior to selecting the cannula, the device will verify that the flow rate is within the range permitted for the selected cannula. If it is not within the appropriate range, a message will display indicating either the flow rate needs to be lowered or a different patient interface type should be selected.

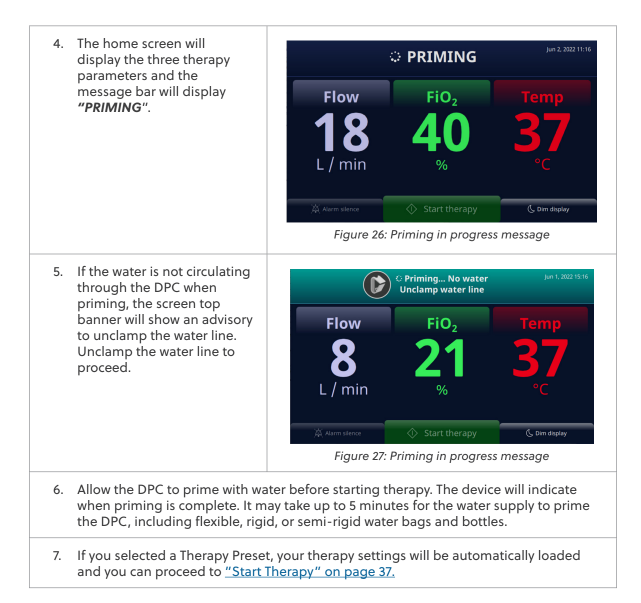
Sample screen message: Current flow (35 L/min) higher than selected cannula limit (20 L/ min). Selecting this cannula will decrease flow to 20 L/min.

2. Select an optional therapy preset by tapping [Preset 1 or Preset 2].

NOTE: Select "none" to manually-adjust therapy parameters.

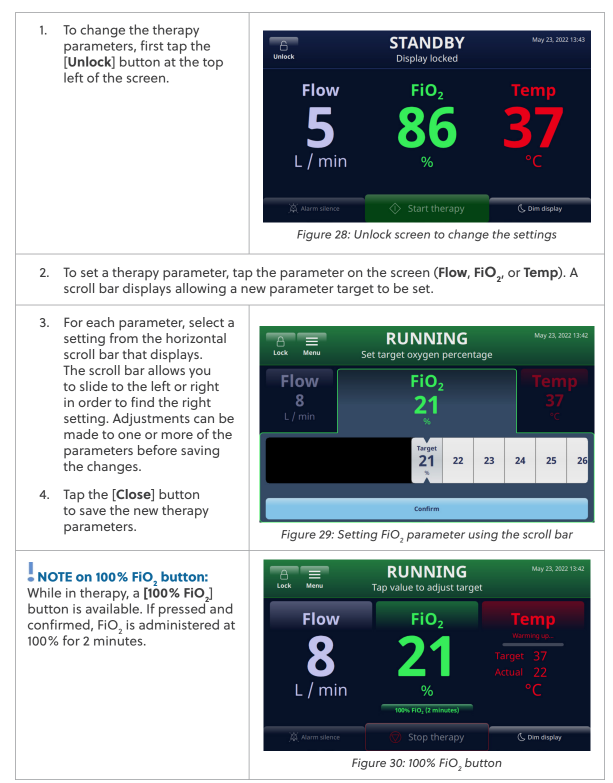




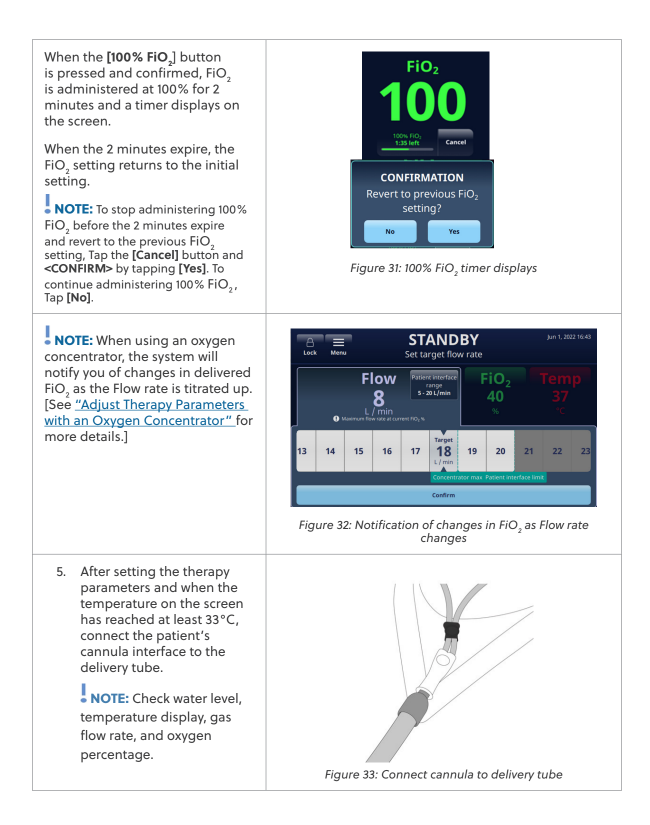


Select Therapy Parameters

If you did not select a therapy preset, follow the steps below to manually adjust the therapy settings.







WARNING: Always follow aseptic technique (including proper hand washing and avoiding direct hand contact with connection points) when setting up the HVT 2.0 device and use Standard Precautions when placing on a patient.
WARNING: Cannula prongs should not obstruct more than 50% of the nares of the patient.
WARNING: Change nasal cannulas when soiled. Replace cannulas according to clinical judgment and hospital policy but not to exceed 30 days continuous use.
WARNING: Improperly sizing the cannula, specifically complete occlusion of the nares by the nasal prongs, may lead to a risk of pneumothorax.
WARNING: To reduce the risk that the patient may aspirate condensed water from the breathing circuit, regularly observe the patient and output of the patient interface for excess water, and if detected remove patient interface from the patient. Water in the center lumen can result from condensation or due to a leak from the outer lumens that surround the breathing circuit. Condensation in the cannula may also occur in certain ambient conditions and low flow rates. If minimal condensation occurs after confirming no leaks, it is recommended to select a lower temperature setpoint.

NOTE: Droplets of condensation may appear at the end of patient delivery tube while unit is warming up. This is normal and will stop within a few minutes when set temperature is reached and the cannula is fitted to the patient.

NOTE: Some condensation around the nose is possible. In addition, a high moisture level may mobilize mucus from nose and sinuses. Make sure patient has a supply of facial tissues.

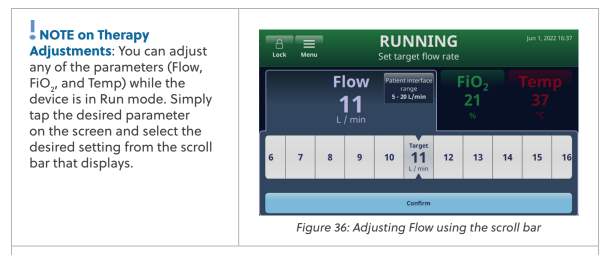
NOTE: Take precautions to minimize cooling of the unheated cannula by trying to maintain contact with the patient's skin and insulating the exposed portion of the cannula with bedding.



Start Therapy

 Start therapy by tapping the [Start therapy] button at the bottom 	A = Lock Menu	STANDBY Tap value to adjust target	May 23, 2022 13:43
of the screen to begin delivering flow.	Flow	FiO ₂	Temp
	5 L / min	86 %	37
)黨 Alarm silence	🚸 Start therapy	C Dim display
	Figur	re 34: [Start therapy] bu	utton
2. After tapping the [Start therapy] button, the device is in Run mode. This is indicated by	C Unlock	RUNNING Display locked	Jun 2, 2022 11:18
the word " RUNNING " displayed at the top of the screen.	Flow	FiO ₂	Temp Warming up Target 37
NOTE: If the target temperature has not been reached, the screen will also indicate the Actual versus	L / min	40 %	Actual 22
Target temperatures as it monitors the status of the temperature.	☆ Alarm silence Figu	Stop therapy or stop therapy or stop in Run me	C Dim display





Adjustments can be made among any or all of the three parameters (Flow, $FiO_{2^{\prime}}$ and Temp) before saving the changes. When adjustments are made, tap the [**Close**] button at the bottom of the screen to save the setting(s). The system automatically adjusts to the new setting with no interruption to the therapy delivery.

Adjust Therapy Parameters with an Oxygen Concentrator

To adjust therapy parameters for a system that is set up to work with an oxygen concentrator, do the following:

- Tap the [Unlock] button at the top left of the screen. The [Lock] and [Menu] buttons will display, and the instruction "Tap value to adjust target" displays under the 'READY' label at the top of the screen.
- To set a therapy parameter, tap the parameter on the screen (Flow, FiO₂, and Temp). A scroll bar will display, allowing a new parameter target to be selected.



3. Tap the **Flow** parameter on the screen to set a Jun 1, 2022 16:43 **STANDBY** target flow rate. The target Set target flow rate rate will be highlighted Patient interface on the scroll bar. If this is Flow FiO₂ a maximum rate for the range 5 - 20 L/min **8** L / min concentrator, a message will display under the scroll bar stating "Concentrator Targe Max." Initially, the slider 13 14 15 16 17 18 19 20 21 22 23 will not go past the maximum limit, and the system will beep. Confirm Figure 38: Setting target flow rate parameter To select a higher flow rate (above the maximum), slide the scroll bar a second time or directly select a value above the limit. The system will update the target to the selected value. 4. Once a higher than the **STANDBY** maximum target has 8 Lock Menu Set target flow rate been selected, the system automatically adjusts the Flow Patient interface FiO₂ oxygen percentage, and range 8 - 45 L/min **35** 31 a message to that effect displays on the screen 🕕 Fi (see the message "FiO, Target lowered to 56% due to 30 31 32 33 34 35 36 37 38 39 40 high flow rate" on the screen below). Confirm Figure 39: System resets the oxygen percentage based on high flow rate



- 5. In order to provide therapy adjustment guidelines, the system displays the following messages above and below the scroll bar (see the screen):
 - FiO₂ lowered to XX% due to high flow rate (displays to indicate the system adjusted oxygen percentage based on high flow rate selected)
 - **Concentrator Max** (displays to indicate the maximum flow rate for the oxygen concentrator).
 - **Cannula limit** (displays to indicate the upper flow rate limit for the cannula selected)



Figure 40: Screen messages providing high flow rate guidelines

6. When adjustments are completed, tap the [**Confirm**] button at the bottom of the screen.



Stop Therapy

Once the device is in Run mode, tap the [**Stop therapy**] button to do any of the following:

- 1. Remove the cannula from the patient.
- 2. Discontinue therapy.
- 3. Shut down the device.

When you tap the [Stop therapy] button, a Confirmation pop-up screen displays:

Lock Menu	RUNNING Tap value to adjust target	May 23, 2022 13:44
Flow 5 L / min	CONFIRMATION Are you sure you want to stop therapy? No, continue	Temp Warming up Target 37 Actual 22 °C
	👽 Stop therapy	🕓 Dim display

Figure 41: [Stop therapy] Confirmation pop-up screen

Discontinue Therapy

When a patient is ready to discontinue use of the HVT 2.0 device, follow the instructions below to stop therapy:

- 1. Remove the cannula from the patient.
- 2. Remove the DPC unit from the HVT 2.0 device.
- 3. Dispose of the cannula, the DPC, and the water supply according to hospital waste management policies.
- 4. Press the () [**Power**] button to place the device in Sleep mode.
- 5. Clean and disinfect the device according to the instructions in <u>"Cleaning &</u> <u>Disinfection"</u> in <u>"Section 7: Maintenance & Disinfection"</u>.



Shut Down

To fully power down the device, proceed as follows:

- 1. First, ensure that no therapy is being delivered to the patient.
- 2. Disconnect the power supply cord from the wall outlet.
- 3. Press the \bigcup [**Power**] button on the device to power down. The [**Power**] button must be pressed, otherwise the internal safety battery becomes active.



Caution: Even a fully charged battery will lose its charge over a period of weeks when the device is not connected to line power. It is recommended that the device be connected to line power at least 2 hours a month to maintain the battery charge. The internal safety battery should only be accessed or replaced by trained service personnel.

Intra-Hospital Patient Transfer

NOTE: This section is only applicable if the HVT 2.0 device is equipped with the HVT 2.0 Transfer Upgrade Kit.

The HVT 2.0 system with the transfer upgrade supports the transfer of patients and patient ambulation within the hospital environment, including from the Emergency Department to inpatient care areas, to and from tests and procedures (non-MRI), and to rehabilitation activities. In mobile use, the HVT 2.0 device is powered by an optional transfer battery pack.

Before using the device to transfer patients, confirm the following:

- 1. The tanks contain adequate oxygen supplies. See Appendix G for estimated tank run time based on set Flow and Fi02 rates.
- 2. The transfer battery is fully charged and installed into the HVT 2.0.
- 3. There is no leaking from the manifold.

The chart below provides estimates of the transfer battery runtime based on flow rates at 21°C ambient temperature and at 37°C temperature setpoint.

Flow rate (L/min)	Run time
25	75 minutes
45	35 minutes



How to Transfer a Patient

- 1. Insert up to two oxygen tanks into the tank holder.
- 2. If tanks do not have a built-in regulator, attach the gas regulator. Connect the tanks to the gas manifold using the hoses.
- 3. Disconnect the wall oxygen hose.
- 4. Unplug the HVT 2.0 device from the wall outlet. (**Note**: Keep the power cord with the device.) The device will now run on the transfer battery without interruption to deliver therapy. The transfer battery indicator at the top of the screen will indicate how much transfer battery life is left.
- 5. Continuously monitor the oxygen tank and transfer battery life throughout the transfer process to ensure that there is no interruption in therapy delivery.

NOTE: When connected to AC power, charging of the transfer battery may be slow or paused if operating the device in environments outside of normal operating temperatures.

NOTE: The status of the transfer battery charge displays on the upper right of the device screen, as follows:

9	4 green	bars	plus	border	(full)
---	---------	------	------	--------	--------

4 green bars (almost full)

- 3 green bars (~ 75% charge)
- 📶 2 yellow bars (~ 50% charge)
- 1 red bar (~ 25% charge)
- Empty battery icon (empty)
- 6. Once the destination is reached, plug the device's power supply cord into a facility-approved wall outlet.
- 7. Connect the wall oxygen hose to the wall oxygen source. Close the oxygen cylinders. The manifold will automatically switch to the wall source.

Section 5: Alarms & Advisories, Hospital/ Clinician Use

Alarms & Advisories Overview

The essential performance of the device consists of proper humidification at high flow rates, heating of water to physiologic levels, and delivery of appropriate FiO₂. The user needs to appropriately respond to alarms and perform the required maintenance to ensure that the essential performance of the device is maintained. All alarms and advisories within the system are visible on the display as they occur. The steps to address the alarm or advisory will also be visible on the display via the [More information] button. Refer to the section "How to get Information about an Alarm" on page 46 for more details.

HVT 2.0 employs two levels of alarms, depending on the urgency of the alarm.

Medium Priority Alarms: The alarm tone is a series of three beeps, repeated every 5 seconds. Medium priority alarms take priority over and are sounded at a higher volume than all low priority alarms. The alarm banner is displayed in yellow and flashes. In this state, the system is not delivering the prescribed therapy to the patient.

Low Priority Alarms: The alarm tone is a series of two beeps, repeated every 20 seconds. The alarm banner is displayed in blue and does not flash. In this state, the system may be delivering the prescribed therapy to the patient.

Advisory: Advisory messages are only visible when no alarms are present. Only one advisory is visible at a time. The advisory tone is one beep.

Keep in mind that:

- All alarms interface to the Nurse Call system if it is enabled.
- Multiple simultaneous alarms will display on the alarm list on the screen.

NOTE: During startup, the device will automatically perform a self-test to ensure proper operation. If any issues are detected, the system will alarm and prevent the starting of therapy.

What to do if an Alarm or Alert Occurs

Alarms will be audible and visual. A tone will sound, indicating the level of priority (Low or Medium), and the warning message will display on the screen. Tap the [Alarm silence] button to silence the alarm for 2 minutes. [Note: To cancel the alarm silencing, tap the [Alarm silence] button again]. For more information about the specific alarm and what action to take, refer to the lists of alarms: <u>Table: Medium Priority Alarms</u> and <u>Table: Low Priority Alarms</u>.



Figure 42: Sample of caution displaying on the screen



WARNING: General Fault alarms are failures in the control or measurement systems. Depending on the cause of the failure, gas delivery may or may not be interrupted. If a General Fault alarm occurs, disconnect the patient, and shut off the device. The device must be repaired by trained service personnel.

NOTE on Operator's Position: The alarm priorities have been designed for an operator positioned within 3 meters of the device.

Alarm System Functionality Verification

To test Alarm system functionality, insert a DPC into the unit and enter Run mode (parameter settings can be variable). Do not connect a cannula to the end of the delivery tube. While the unit is in Run mode, place your gloved thumb over the end of the delivery tube (see illustrations below). Verify that the Occluded/Blocked Tube alarm message displays on the screen and an audible alarm occurs. After verifying alarm functionality, wipe down the delivery tube outlet connector using aseptic technique.

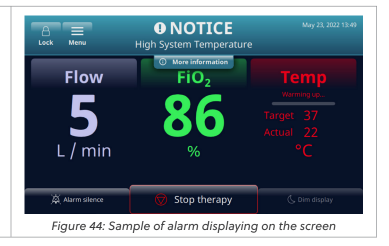


Figure 43: Verifying alarm system functionality

How to get Information about an Alarm

When an alarm sounds and flashes a banner on the screen, the device will display the name of the alarm. A [More information] button is available just below the banner to provide more details about the alarm.

NOTE: All alarms can be found on the <u>"Event Log"</u>.





After tapping the [**More information**] button, details about the alarm and what action to take will display. Tap the [**Hide info**] button when you are done in order to return to the previous screen.

High Sys	NOTICE tem Temperatur	re	
Check that the vents at the back and the bottom of the device are not obstructed.			
If the error persists, call your Vapotherm service provider for help. (error code 62)			
Flow 5 L/min Floz 86 % Temp 37 °C			
	at the back	at the back and the bottom call your Vapotherm service	

Multiple Alarms - Medium Priority and Low Priority

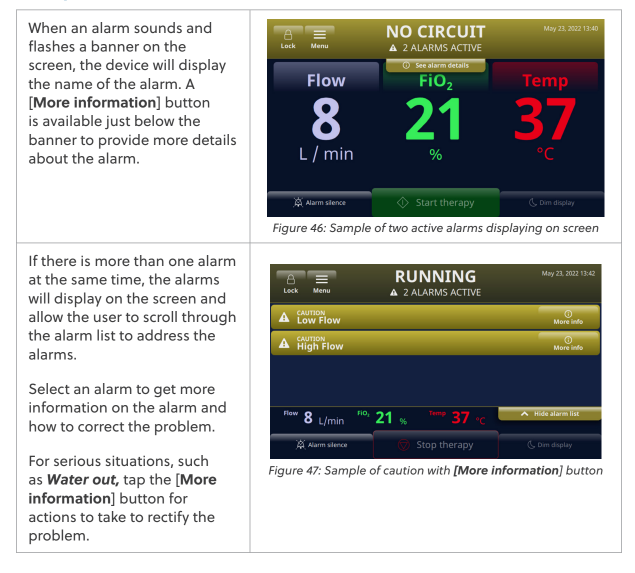


Table: Medium Priority Alarms

Message	Description and Actions Required
Low Flow	Detected flow does not match user setting. Check cannula selection.
	If the alarm re-occurs, call your Vapotherm service provider for help.
High Flow	Detected flow does not match user setting.
	If the alarm re-occurs, call your Vapotherm service provider for help.
Low Gas	Delivered gas temperature low.
Temperature	If the alarm re-occurs, call your Vapotherm service provider for help.
Low Gas	Delivered gas temperature low.
Temperature	Max temperature when using transfer battery is 37°C. To clear fault condition, reduce temperature setpoint
High Gas	Delivered gas temperature high.
Temperature	If the alarm re-occurs, call your Vapotherm service provider for help.
High Gas	Gas temperature exceeded limit.
Temperature - Therapy Stopped	If the alarm re-occurs, call your Vapotherm service provider for help.
FiO ₂ Low	Detected FiO ₂ does not match user setting. Check oxygen source.
	If the alarm re-occurs, call your Vapotherm service provider for help.
FiO ₂ High	Detected FiO_2 does not match user setting.
	If the alarm re-occurs, call your Vapotherm service provider for help.
Water Out	Water empty.
	Replace supply.
Occluded / Occluded delivery tube/cannula. Blocked Tube	
	Remove any kinks, clamps, or obstructions in delivery tube and cannula.
No Power Connected	Device running on safety battery. Therapy is being provided with unheated gas.
	Connect device to power.
Oxygen Input Overpressure	Oxygen source input pressure too high.

Table: Medium Priority Alarms (Continued)

Message	Description and Actions Required
Circuit Removed	Circuit removed while therapy running. Therapy stopped.
During Therapy	To continue therapy, re-insert circuit.
FiO ₂ Error -	Detected FiO ₂ does not match user setting.
Therapy Stopped	If the alarm re-occurs, call your Vapotherm service provider for help.
Safety Battery	Safety battery failed or not installed.
Failure	Call your Vapotherm service provider for help.
System Error	If the error persists, remove the oxygen source from the device. Call your Vapotherm service provider for help.
Oxygen	Low oxygen concentration from concentrator detected.
Concentrator	Check that the selected value in the Oxygen source menu matches
Error	the connected oxygen concentrator.
System Error - Therapy Stopped	A device fault has occurred. Therapy stopped and cannot restart. To clear fault condition: power off the device, then power it on again . If alarm re-occurs, call your Vapotherm service provider for help.
Circuit Error -	Circuit error. Therapy stopped.
Therapy Stopped	If alarm re-occurs, call your Vapotherm service provider for help.
System Error - Therapy Stopped	Device overheated. Therapy stopped and cannot restart. To clear the fault condition: power off the device, then power it on again. Call your Vapotherm service provider for help. (Specific error code will be shown).

Table: Low Priority Alarms

Message	Description and Actions Required
Water Low	Water running low. Replace supply.
No Power	Device running on safety battery.
Connected	Connect device to power.
High System	Check that the vents on the back and the bottom of the device are not obstructed.
Temperature	If the error persists, call your Vapotherm service provider for help.

Table: Information Advisories

Message	Description and Actions Required
Water Low	Water running low.
	Replace supply.
Water Out	Water empty.
	Replace supply.
Safety Battery Low	Safety battery charge is low.
Transfer Battery Active	Device running on Transfer Battery.
Transfer Battery	Transfer battery charge is low.
Low	Connect device to power outlet to power or replace transfer battery with charged battery.
O ₂ Configuration Mismatch	Detected O ₂ input pressure (wall/tank) does not match selected O ₂ source (concentrator).
	Confirm oxygen source selection in settings menu.
O ₂ Configuration Mismatch	Detected O ₂ input pressure (concentrator) does not match selected O ₂ source (wall/tank).
	Confirm oxygen source selection in settings menu.

Water Level Sensing

The HVT 2.0 device is equipped with water level sensing to notify clinicians of the water level and indicate when water must be added. It provides notifications in the following intervals:

Alarm Notification	Water Level
Low Priority Alarm – Message: Water Supply Low. Replace water supply.	First Alarm: Water supply running low.
Medium Priority Alarm – Message: Water Supply Empty. Replace water supply.	Second Alarm: Out of water.

Table: Recommended water change interval based on flow rate and operation at 37°C.

Flow Rate	Average water usage per day	Recommended Change Interval
5-10 L/min	650 ml	500 ml / 12 h
10-20 L/min	1300 ml	500 ml / 8 h
20-30 L/min	2000 ml	1000 ml / 12 h
30-45 L/min	2600 ml	1000 ml / 8 h



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Section 6: Training

See your Vapotherm Service Provider or contact Vapotherm for detailed product training.

Additional training can be found at the Vapotherm Academy website (<u>https://academy.vapotherm.com/</u>) accessible from the <u>Vapotherm website</u>.

Section 7: Maintenance & Disinfection

All HVT 2.0 Disposables (patient circuits, cannulas, and accessories) are validated for single patient use for up to 30 days.

Preventive Maintenance

Part Number/Item Description	Maintenance required (Replace)	Requires return to Vapotherm Service Center
Patient Air Intake Filter	 Replace the filter every 6 months or as recommended per your institution's policy. (See instructions below, <u>"Replace the Patient Air</u> <u>Intake Filter")</u> 	No
Internal Safety Battery	• Replace every 5 years (See instructions below, <u>"Replace the Internal</u> <u>Safety Battery")</u>	No
Internal Blower	Replace every 5 years	Yes

The table below outlines the anticipated schedule of routine maintenance.

Inspect the Power Supply Cord

Visually inspect the power supply cord whenever you connect the device to line power to verify that the power supply cord is not damaged or kinked.

Inspect the Chassis Vent and Cooling Fan

Visually inspect the chassis vent (below the device) and cooling fan (on the device's back panel) to verify that they are not obstructed and allow the free flow of air.

Replace the Patient Air Intake Filter

The patient Air Intake Filter should be inspected frequently for particulate contamination and replaced every 6 months or as recommended per your institution's policy.

To replace, insert the patient air intake filter into the back of the HVT 2.0 device.



Figure 48: Insert patient air intake filter in back of device

Replace the Internal Safety Battery

Caution: HVT 2.0 will not operate without the internal safety battery in place. Have an internal safety battery on hand to ensure the continued availability of the use of the device. To ensure safe and reliable operation, use only the Vapotherm specified replacement battery.

After five years of use, the internal safety battery must be replaced. The internal safety battery is located in the same compartment as the Transfer Battery. Remove the Transfer Battery access door, then remove the four screws holding the internal safety battery cover. Use the pull tab to slide out the internal safety battery. Slide the new internal safety battery until it rests on the bottom of the compartment. Replace the cover and screws to assure that the internal safety battery is secured in position.

/!\



Cleaning & Disinfection

WARNING: Before cleanin

/!\

WARNING: Before cleaning and disinfecting, unplug the device from line power.

Caution: Do not use bleach, organic solvents, or abrasive cleaners. Hypochlorite solutions liberate toxic gases such as chlorine when acidified or heated. The reaction with ammonia or with substances that can generate ammonia can produce chloramines which are also toxic and have explosive potential.

The entire disposable patient circuit (DPC) is disposable, and no disinfection is required. The main unit, including the docking station for the disposable water path, should be wiped down with Super Sani-Cloth® or another approved cleaner (see list below). **Unplug the HVT 2.0 while cleaning and disinfecting.** The HVT 2.0 device must always be cleaned and disinfected between patients. Follow the steps below to ensure a clean and disinfected device.

- 1. Wipe down the main unit with Super Sani-Cloth® or another approved cleaner.
- 2. Visually inspect for visible soil. If visible soil is present, repeat step 1. A brush (e.g., Spectrum M16 brush) may be used in addition to wiping down the unit.
- 3. Wet the main unit with another Super Sani-Cloth[®]. Keep the surface wet for at least six minutes. Use additional Super Sani-Cloth[®], if needed.

The following detergent wipes can be used to remove any soil from the device:

- 70 to 90% isopropyl alcohol wipe
- 2% (maximum) Chlorine cleaning solution
- 6% (maximum) Hydrogen Peroxide
- Metrex CaviWipes®
- PDI Healthcare Sani-Cloth® AF3 Germicidal
- EcoLab Incidin® OxyWipe
- BODE Chemie GmbH Bacillol® 30 Tissues
- GAMA Healthcare LTD. Clinell[®] Alcohol Wipes
- Vernacare Tuffie Disinfectant Wipes



WARNING: All disposable components are labeled as "single patient use only" and must be replaced after 30 days of use on a single patient. Cannulas should be replaced according to clinical use, but not to exceed 30 days. Do not attempt to sterilize or reuse any of these components, and follow all local and federal regulations for disposal. Outside the U.S., follow national or international regulations.

NOTE: The HVT 2.0 device should be cleaned according to the above instruction after each patient.

Disposal Instructions

HVT 2.0 Device Disposal

The HVT 2.0 device contains electronics. Do not discard with regular waste. Instead, return the device to Vapotherm or dispose of it according to local guidelines for disposing of electronics.

For the European Union, dispose of the device according to the Waste Electrical and Electronic Equipment (WEEE) directive.

Patient Circuit and Accessories Disposal

At the end of use, place the cannula, the Disposable Patient Circuit (DPC), and any other Vapotherm consumable accessories that were used into a waste supply. No disassembly of the disposable is needed prior to disposal. Discard all disposables according to hospital guidelines and local regulations for the safe disposal of medical single-use items.

Service

Service on the device should only be performed by qualified, certified service technicians. To prevent injury, do not attempt to do any service to the device while the device is connected to a patient.

Do not use the device if it is damaged or not working properly. If it is damaged or not working properly, contact Vapotherm or your authorized Vapotherm representative.

Vapotherm provides 24/7* Technical Support: 1 (888) 320-4506.

* Available in the U.S. only. Customers outside the U.S. should contact their distributor or local authorized Vapotherm Service Center. If you do not know who your service center or distributor is, contact Vapotherm.

Software Updates

Software updates can be made locally via the USB port. Software updates will be made available to customers upon release.



Appendix A – Nurse Call System Installations

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Installation Verification Procedure	

Nurse Call Installation Instructions

Introduction

The information below describes the Nurse Call communication interface of the HVT 2.0 device. This information is intended for use by hospital IT, biomedical engineering, or other experts who wish to connect the HVT 2.0 into a Nurse Call System.

Scope

This information applies to the current design and embedded firmware version of the HVT 2.0 device.

Nurse Call Hardware Interface Description

The Nurse Call Communication Cable is available in four variants:

Cable Type
NO contact
NO + 10k contact
NC contact
NC + 10k contact

Please order the cable type compatible with your nurse call system.

Installation Verification Procedure

Verify that the complete system is functioning by creating a Test Alarm and checking that the correct result has been received.

- 1. Connect and turn on the HVT 2.0 device. (Refer to the <u>"Starting Therapy"</u> section of the HVT 2.0 Instruction for Use).
- 2. Force an alarm occurrence by placing your thumb over the distal end of the delivery tube to simulate a blocked tube condition.
- Confirm that you receive the result you expect in the system according to the hospital standard for that alarm, such as a warning light is turned on or an audio signal is received.
- 4. Release your thumb from over the distal end of the delivery tube to clear the blocked tube condition and confirm that the nurse call alarm condition clears.

After the test has been successfully concluded, the Nurse Call is ready for use.

Appendix B – Electronic Medical Records (EMR) Integration

The HVT 2.0 provides an isolated RS-232 serial interface to support hospital integration of HVT 2.0 operational data with an electronic medical record. Information for the hardware interface and data format are available from Vapotherm upon request.

The HVT 2.0 data stream is a transmit only communication protocol. When enabled, the data stream is transmitted when therapy is on. Transmitted data includes:

- Therapy parameters (flow rate, temperature, FiO₂, etc.)
- Pulse oximetry data (if unit is OAM capable)
- System state information

No patient identifiable information is included in the data stream.

NOTE: To maintain electrical safety, use only Vapotherm provided interface cables.

Integration with an EMR system is the responsibility of the customer and is often done by third-party integrators. While operation of HVT 2.0 is not affected by the configuration of devices connected to this port, safe and effective use of transmitted data is the responsibility of the integrator and responsible organization. Bernoulli Systems (formally Nuvon) and Capsule are the Vapotherm-supported third party integrators.



Figure B1: HVT 2.0 EMR Link Cable placed in EMR data port



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Appendix C – Technical Specifications

Physical Characteristics

Dimensions:	Height 9.1" (23.114 cm), width 11.1" (28.194 cm), depth 12.1" (30.734 cm)	
Weight:	16 lbs.	
Mounting:	Tabletop or roll stand; fits IV poles up to 1.5" (38mm) diameter	
Gas Connections:	: Standard DISS non-interchangeable fittings for oxygen	
Fuses: Type T4AH250V		
Contact with Patient:	Indirectly via the delivery tube and cannula	
Context of Use:	General care floors, emergency departments, intensive care units, long-term acute care facilities, skilled nursing facilities.	

System Requirements

AC Power:	100 to 240VAC, 50 to 60Hz
Backup Power/SafetyThe reserve power will last for a minimum of 15 minutes atBattery:45 L/min flow rate	
Oxygen Supply:	Oxygen or oxygen concentration at inlet pressures between 4 and 87 psi (600 kPa)
	For respiratory distress: Oxygen at inlet pressures between 40 and 87 psi (600 kPa)
Water Supply:	Sterile water in pre-filled sealed bags or bottles.

Environment

Operation:	Ambient Temperature: 18 to 30°C
	Ambient Relative Humidity: 15 to 90% non-condensing
	Ambient Pressure: Standard atmospheric (not to be used in hyperbaric cond.)
Storage and Shipping:	Ambient Temperature: -10 to +50°C
	Ambient Relative Humidity: 10 to 90% RH non-condensing
Altitude:	0 to 3000m (0 to 9843 ft): full flow range available.

Performance

	Range: 33°C to 39°C at exit from the delivery tube; adjustable		
Temperature:	Resolution: 1°C		
	Accuracy: ± 2°C		
Warm-up Time:	± 2°C of 33°C set point < 5 minutes (at ambient 23°C)		
	Range: 21% to 100% O ₂		
Oxygen Percentage:	Accuracy: ± 2% Wall or Tank Source		
	±4% Oxygen Concentrator		
	Resolution: 1%		
Flow Rate:	5 to 45 L/min BTPS		
Flow Rate Accuracy	Greater of 0.5 L/min or 10% of setting		
Expected Service Life of Device:	5 years		

NOTE: Temperature, O_2 %, and Flow Rate accuracies have been established inclusive of all test equipment measurement uncertainties.

NOTE: Absolute Humidity > 16 mg/L at all combinations of flow rates, temperatures, and oxygen concentrations.

NOTE: Absolute Humidity \ge 33 mg/L is maintained at flow rates of \ge 8 L/min, temperatures \ge 37°C, and at all available FiO₂ settings.

NOTE: When using an oxygen concentrator, 45 L/min flow often has a maximum FiO₂ range of 36%.

Useful Life

Part	Useful Life
HVT 2.0 device	5 years
Disposable Patient Circuit (DPC)	30 days
ProSoft Nasal Cannulas	30 days
Tubing Adapter	30 days
Power Supply Cord	5 years
Patient Air Intake Filter	6 months, or as recommended per institution policy
Internal Safety Battery	5 years
Internal Blower	5 years
Roll Stand	5 years
Transfer Upgrade Kit	5 years



Use with Oxygen Concentrators

WARNING: If used with an oxygen concentrator, the maximum oxygen percentage will be limited depending on the type of concentrator used and total set flow.

The table below shows the maximum O_2 delivered for the set HVT 2.0 flow rate and concentrator flow limit of 5 L/min and 10 L/min. The HVT 2.0 is compatible with the Philips EverFlo Concentrator, Inogen Home Concentrator and Respironics Millennium M10 Concentrator.

	HVT 2.0 FiO ₂ Output					
		OC output of 5 L/min OC output of 10 L/m				
Ð	5	90%	90%			
HVT 2.0 Flow rate	10	56%	90%			
No	15	44%	67%			
.0 F	20	38%	56%			
/Т 2	25	35%	49%			
Ŧ	30	33%	44%			
	35	31%	41%			
	40	30%	38%			
	45	29%	36%			

Inputs

Airway Gas:	Oxygen DISS Connector or Oxygen ¼" barb nipple	
External Device Comm.: USB and Ethernet		

Outputs

(Only compatible with other IEC60601-1 approved devices)

Nurse Call:	1/4" modular jack
-------------	-------------------

Bidirectional Connectivity

USB:	USB 2.0, Type A Connector. Up to 64 MB capability.	
		WARNING : Do not connect any device, system, or accessory that has not been approved by Vapotherm.
Ethernet:	RJ45 mo	dular jack
EMR Connectivity:	RS-232 serial connection via an interface cable	
WiFi:	2.4 GHz,	, 802.11 b/g/n



Minimum Alarm Sound Pressure Ranges

High Priority Alarm	> 80 dBA
Medium Priority Alarm:	> 50 dBA
Low Priority Alarm:	> 45 dBA

Alarm Type and Test Conditions	Calculated Average A-weighted sound pressure level (dBA)	
	Max	Min
Medium priority alarm	75.65	51.46
Low priority alarm	72.27	45.00

Available Parameter Settings and Factory Defaults

Parameter	Available Settings	Factory Defaults
TEMP SetPoint	33°C to 39°C	37°C
Gas Flow	5 to 45 L/min	n/a
Percentage Oxygen	21% to 100%	n/a
Care Area (Service Setting)	Acute Care (hospital or sub-acute)	Acute Care (hospital or sub-acute)

NOTE: When using an oxygen concentrator, 45 L/min flow often has a maximum FiO_2 range of 36%.

Cybersecurity

Vapotherm has designed the HVT 2.0 system to be resistant to cyberattacks in order to maintain the integrity and availability of the device. These features do not require any user configuration or action.

- Device does not collect or maintain any protected health information (PHI).
- Real time therapy control is isolated through a secure proprietary link to hardware with no available network ports.
- Device is configured securely to prevent unauthorized access either wirelessly or through the communication ports intended for future use.
- Software updates are performed via USB drive, rather than over the Internet.
- Software updates are verified by digital signature.

Standards/References

Reference	Description
ANSI/AAMI ES60601-1: 2005 / 12012 CAN/CSA-C22.2 No. 60601-1:14 IEC 60601-1: 2005 + A1: 2012, Third Edition-	Medical Electrical Equipment - Part 1: General Requirements for Basic Safety and Essential Performance
IEC 60601-1-2 ed 4.0 (2014-02)	Medical Electrical Equipment - General requirements for safety - Collateral Standard Electromagnetic disturbances
IEC 60601-1-6:2010, AMD1:2013	Medical Electrical Equipment Part 1 – 6 General Requirements for Safety – Collateral Standard: Usability
IEC 60601-1- 8: 2006 (Second Edition) + Am.1: 2012	General requirements for basic safety and essential performance - Collateral Standard: general requirements, tests, and guidance for alarm systems in medical electrical equipment and medical electrical systems
IEC 60601-1-9:2007, AMD1:2013	General requirements for basic safety and essential performance – Collateral Standard: Requirements for environmentally conscious design
IEC 62366-1: 2015	Medical Devices - Application of Usability Engineering to Medical Devices
ISO 80601-2-74:2017	Respiratory tract humidifiers for medical use — Particular requirements for respiratory humidification systems

Guidance and Manufacturer's Declaration

Electromagnetic Emissions

The HVT 2.0 is intended for use in the electromagnetic environment specified below. The user of the device should ensure that it is used in such an environment.

Emissions Test	Compliance	Electromagnetic environment- guidance
RF emissions CISPR 11	Group 1	The device 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 equipment.
RF emissions CISPR 11	Class B	The device is suitable for use in all establishments
Harmonic emissions IEC 61000-3-2	Class A	and may be used in domestic establishments and those directly connected to the public low-voltage power supply network that supplies buildings used
Voltage fluctuations/ flicker emissions IEC 61000-3-3	Complies	for domestic purposes.

Manufacturer's Declaration – Electromagnetic Immunity

IEC 60601-1-2:2014		
Sub Test	Passed Parameters	
Electro-Static Discharge* IEC 61000-4-2 ed2.0 (2008-12)	±8kV Contact discharge ±15kV Air discharge	
Radiated RF Susceptibility IEC 61000-4-3:2006, +A1:2007, +A2:2010	80- 2700MHz @ 20 V/m, 80% AM @2Hz	
Electrical Fast Transients IEC 61000-4-4 ed3.0 (2012-04)	±2kV AC mains	
Surges IEC 61000-4-5 ed2.0 (2005)	±0.5,1kV Line to line	
Line Conducted RF Susceptibility IEC 61000-4-6 ed4.0 (2013)	0.15-80MHz @ 3Vrms (6Vrms in ISM and Amateur Radio Bands) 1kHz AM 80% modulation	
Power Frequency Magnetics IEC 61000-4-8 ed2.0 (2009-09)	30A/m @ 50/60Hz	
Voltage Dips and Dropouts IEC 61000-4-11 ed2.0 (2004-03)	Per Standard	

*ESD mitigation measures include maintaining adequate relative humidity and touching a large metal object that is away from the HVT 2.0 and the patient before touching the device. Note that a "**Water Out**" message could be displayed in response to electrostatic discharge to the device.

Test Specifications for Enclosure Port Immunity to RF wireless communications equipment

Test Frequency (MHz)	Immunity Test Level (V/m)
385	27
450	28
710	
745	9
780	
810	
870	
930	
1720	28
1845	
1970	
2450	
5240	
5500	9
5785	



Appendix D: Software Update Process

The software update process allows the user to press one button on the Software Update screen to update both the Controller software and GUI application.

IMPORTANT NOTES on the Software Update Process:

- Therapy must be stopped before a software update can be started.
- The device will update the Controller software first and then the GUI.
- The device will reboot after each software application update.
- Dialog boxes will be displayed throughout the update process to keep the user informed on the progress of the update.
- A software update will not be allowed while in therapy. A dialog box will be displayed telling the user they must stop therapy to perform a software update.
- The [**Update software**] button will only be enabled when a USB (which contains the complete update package) is plugged into the back of the device.
- A dialog box will display if a failure condition occurs .However, the device will be in a recoverable state, i.e., it will automatically revert to the previous state before the failed software update attempt.
- A final dialog box will be displayed when the software update is successful.

Steps to update Controller and GUI software:

 Place the USB stick containing the update into the back of the device.

> When the USB is in place, the [**Update software**] button will be enabled on the device screen.

2. Press the [Update software] button to begin.



Igure DI: [**Update software**] button enabled when USB plugged into back of device



3. A confirmation popup will display.



Figure D2: Confirmation pop-up screen

4. Press the [Yes, start] button to begin the process.

The Software Update process will begin with the Controller software update and then will reboot the system.

5. After the Controller software update and reboot, the GUI update occurs.

The device reboots after updating the GUI software.

6. When the process is complete, a success dialog box will display on the screen.

Software update succeeded.

7. Press the [OK] button to acknowledge that the software update succeeded.

NOTE: If the update failed, a dialog box will display informing the user to contract customer service.



Appendix E: Symbol & Icon Key

Device Screen Icons

	Silence Alarm button
	Internal safety battery charge status
C	Dim display
Settings	Settings Menu
Unlock	Touch screen Lock/Unlock selector (System Settings Menu)
♦ Start therapy	Start Therapy / Stop Therapy buttons
Ċ	Power button
\sim	AC Power connected
===	Not in use
دب]	Internal safety battery charging (when AC power is connected)
۲	Internal safety battery is active
	Transfer battery - 4 green bars plus border (full)
	Transfer battery - 4 green bars (almost full)
(🦻	Transfer battery - 3 green bars (~ 75% charge)



(7	Transfer battery - 2 yellow bars (~ 50% charge)
(1	Transfer battery - 1 red bar (~ 25% charge)
(/	Transfer battery - empty battery (empty)
Temp	Temperature
Flow	Flow rate
FiO ₂	FiO ₂ percentage

Labeling Symbols

	Caution: U.S. Federal Law restricts this device to sale by or on the order of a physician.
Note: this is not an internationally recognized symbol.	Medical Device Symbol. (Note: This is not an internationally recognized symbol.)
RoHS 2011/65/EU	Product complies with requirements of the RoHS Directive 2011/65/EU and must bear the CE marking.
	For Indoor Use Only.
IP22	IP22 Drip Proof and prevents adult finger ingress
~	Alternating Current
2	Single Patient Use; Do Not Re-Use
X	Do Not Cover



	CLASS II Equipment
Ť	Shock Protection: Type BF
X	This symbol indicates that the waste of electrical and electronic equipment must not be disposed of as unsorted municipal waste and must be collected separately. Please contact an authorized representative of the manufacturer for information concerning the decommissioning of your equipment.
18°C 30	Ambient Temperature Operating Range
	Consult Instructions for Use
Ĩ	e-IFU: Consult Instructions for Use, indicates the web address where the e-IFU is located, e.g., <u>https://vapotherm.com/international-documents/</u>
MR	MR Unsafe – keep away from magnetic resonance imaging (MRI) equipment
€ 0297	Mandatory marking for devices entering the European market to indicate conformity with the essential health and safety requirements set out in European Directives. Accompanied by the 4-digit ID number of the notified body.
	Manufacturer, adjacent to name and address of the manufacturer
	Date of Manufacture, YYYY-MM-DD
LOT	Lot Number, accompanied by the manufacturer's batch code.



REF	Reference Number
SN	Manufacturer's Serial Number
	Use By Expiration Date, YYYY-MM-DD
NON STERILE	Non-Sterile – Device has not been sterilized
	Caution, consult accompanying documents or Attention, see Instructions for Use
EC REP	Authorized Representative in the European Community, accompanied by the name and address of the authorized representative in the European Union.
CH REP	Authorized Representative in Switzerland, accompanied by the name and address of the authorized representative in Switzerland.
	External DC Power Input
٢	Internal Battery: Lithium-ion 14.4V, 6900mA hour. Replaceable by service technician only. See the <u>"Service"</u> section in <u>"Maintenance & Disinfection"</u> .
	Fuse: replace with indicated fuse only
O 2	Oxygen Connection Port. See Oxygen Supply specifications in <u>"System</u> <u>Requirements"</u> of <u>"Appendix C – Technical Specifications"</u> .



+	Nurse Call Connection. See <u>"Appendix A – Nurse Call System Installations".</u>
	Electronic Medical Record Connection.
aux	Auxiliary Connection. For Factory Use Only.

Vapotherm Inc. has declared that this product conforms with the European Council Directive 93/42/EEC Medical Device Directive when it is used in accordance with the instructions provided in the Instructions for Use.

Trademark citations:

Masimo SET™, X-Cal™, LNCS®, and RD SET™ are registered trademarks of Masimo Corporation

Appendix F: Glossary

Terms

- Disposable Patient Circuit (DPC) The DPC (or "Circuit") enables delivery of high velocity therapy for pediatric and adult patients for the full flow range from 5 to 45 L/min. The DPC is a component of the HVT 2.0 system and is detachable and disposable. Note: The DPC is provided fully assembled and ready to use.
- Electronic Medical Records (EMR) System Electronic Medical Records system. The HVT 2.0 system can be interfaced with the hospital's Electronic Medical Records (EMR) system.
- FiO_2 Fraction of inspired oxygen. It is the concentration or percentage of oxygen that a person inhales.
- **HVT 2.0 device** The HVT 2.0 system is made up of the controller device and the disposable patient circuit (DPC), which together enable the delivery of high velocity therapy to the patient.
- **Nasal Cannula** Device used to deliver supplemental oxygen or increased airflow to a patient in need of respiratory help. It consists of a lightweight tube that, on one end, splits into two prongs that are placed in the nostrils and from which a mixture of air and oxygen flows.
- Nurse Call System A system in the health care facility that allows patients to call or contact their nurse or nurse's station. HVT 2.0 allows connectivity between the device and the hospital's Nurse Call system.
- **Transfer Upgrade Kit** This kit is used with the roll stand and allows moving the patient from one location to another within the hospital while the patient continues to be connected and receiving therapy. The kit includes a battery, oxygen manifold, oxygen hoses, and adjustable oxygen tank holder.
- **Tubing Adapter** This adapter allows the DPC to be connected to a trach collar or t-piece in place of a nasal cannula.

Appendix G: Transfer and Ambulation Tank Table

The HVT 2.0 allows the mobile delivery of optimally humidified high velocity therapy within a hospital environment. The length of available mobile therapy depends on 2 factors:

- Cylinder gas supply:
 - o From 14 minutes to indefinite, based on the FiO2 and Flow setting.
- Transfer Battery run time:
 - o Up to 75 minutes from a full charge @ 25 LPM
 - o At least 35 minutes from a full charge at 45 LPM
 - o Transfer battery can be hot-swapped

Flow	% Oxygen								
LPM	21%	30%	40%	50%	60%	70%	80%	90%	100%
5	ω	1076	510	334	248	198	164	140	123
10	ω	538	255	167	124	99	82	70	61
15	ω	359	170	111	83	66	55	47	41
20	ω	269	127	84	62	49	41	35	31
25	ω	215	102	67	50	40	33	28	25
30	ω	179	85	56	41	33	27	23	20
35	ω	154	73	48	35	28	23	20	18
40	ω	135	64	42	31	25	21	18	15
45	œ	120	57	37	28	22	18	16	14

- Runtime in minutes above accounting for ca. 20% safety surplus
- Calculation based on E-Type cylinder @2000PSI (680I Oxygen)
- Actual runtimes may vary based on the amount of gas in cylinder

Warranty

Vapotherm expressly warrants, for a period of one (1) year from the date of shipment by Vapotherm to the initial purchaser of the HVT 2.0 device ("Customer"), that the HVT 2.0 device shall meet the specifications set forth in the applicable official operating Instructions for Use provided with each HVT 2.0 device (the "Instructions"). The sole remedy for this warranty is that Vapotherm shall, at its sole option, either refund, repair or replace any or all of any HVT 2.0 device that is defective at no cost to the Customer. Vapotherm shall pay any shipping charges required in repairing or replacing any part, or all of an HVT 2.0 device during the warranty period. Thereafter, shipping charges shall be paid by the Customer. The Customer shall also be responsible for the cost of labor for repairs. This warranty does not apply to any disposable component to the HVT 2.0 device, including without limitation the disposable patient circuits and hoses supplied with the HVT 2.0 device.

The warranty set forth herein shall become null and void if: (1) the HVT 2.0 device is not used or serviced in accordance with the applicable Instructions or any related preventative maintenance instructions provided with the HVT 2.0 device; or (2) the HVT 2.0 device is opened or tampered with, or if repairs or service are performed or attempted on the HVT 2.0 device by anyone other than Vapotherm or a Vapotherm-certified service center.

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